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DEDICATION

To Michael, Mary, Paul, and Margaret
The purpose of A Manual of Laboratory and Diagnostic Tests, in this Seventh edition, is to promote the delivery of safe, effective, and informed care for patients undergoing diagnostic tests and procedures and also to provide the clinician and student with a unique resource. This comprehensive manual provides a foundation for understanding the relatively simple to the most highly complex diagnostic tests that are delivered to varied populations in varied settings. It describes the clinician's role in providing effective diagnostic services in depth, through affording the necessary information for quality care planning, individualized patient assessment, analysis of patient needs, appropriate interventions, patient education, patient follow-up, and timely outcome evaluation.

Potential risks and complications of diagnostic testing mandate that proper test protocols, interfering factors, follow-up testing, and collaboration among those involved in the testing process be a significant part of the information included in this text.

This book is organized into 16 chapters and 12 appendices. Chapter 1 outlines the clinician's role in diagnostic testing and includes interventions for safe, effective, informed pre-, intra-, and posttest care. This chapter includes a Patient's Bill of Rights and Responsibilities, a model for the role of the clinical team in providing diagnostic care and services, test environments, reimbursement for diagnostic services, and the importance of communication as key to desired outcomes. The intratest section is expanded to include information about collaborative approaches facilitating family presence during invasive procedures, risk management, the collection, handling, and transport of specimens, infection control, controlling pain, comfort measures, administration of drugs and solutions, monitoring fluid intake and loss, using required equipment kits and supplies, properly positioning the patient for the procedure, managing the environment, and patient monitoring. The reader is referred back to Chapter 1, Diagnostic Testing, throughout the text for information about the clinician's role and diagnostic services. Chapter 2, Chapter 3, Chapter 4, Chapter 5, Chapter 6, Chapter 7, Chapter 8, Chapter 9, Chapter 10, Chapter 11, Chapter 12, Chapter 13, Chapter 14, Chapter 15 and Chapter 16 focus upon specific categories that include:

- Chapter 2: Blood Studies
- Chapter 3: Urine Studies
- Chapter 4: Stool Studies
- Chapter 5: Cerebrospinal Fluid Studies
- Chapter 6: Chemistry Studies
- Chapter 7: Microbiologic Studies
- Chapter 8: Immunodiagnostic Studies
- Chapter 9: Nuclear Medicine Studies
- Chapter 10: X-ray Studies
- Chapter 11: Cytology, Histology, and Genetic Studies
- Chapter 12: Endoscopic Studies
- Chapter 13: Ultrasound Studies
- Chapter 14: Pulmonary Function and Blood Gas Studies
- Chapter 15: Prenatal Diagnosis and Tests of Fetal Well-Being
- Chapter 16: Special Systems, Organ Functions, and Postmortem Studies

CHAPTER CONTENT AND FEATURES

- Background rationale
- Test purpose
- Interfering factors
- Description of the procedure protocol and time frames and test completion
- Reference ranges and normal values, expectations
- Patient involvement (eg, history of signs and symptoms, body position, breathing instructions, electrode placement, compliance issues, patient right to refuse testing)
- Method of specimen collection (biohazard guidelines), handling, and transportation
- Clinical implications with interpretation of abnormal findings, unexpected outcomes, and disease patterns
- Interventions for pretest patient preparation (medications, fasting), explanation of benefits and risks, intratest patient care (appropriate monitoring, conscious sedation), and posttest patient aftercare (includes monitoring, explanation of further testing and treatment modalities)

Special features integrated into the format include:

- The clinician’s role in providing diagnostic services.
- Clinical Alerts and Education Alerts that signal special cautions.
- Specific guidelines listed for each test phase.
- Expected outcomes with evidence-based patient expectations and reference ranges as defined by the specialty.
- A user-friendly format of the text to support easy information retrieval.
- Both conventional and SI units are listed and, where possible, age-related reference values are also listed as a
component of normal reference values.

- Numerous examples of test values and clinical considerations for newborn, infant, child, adolescent, and older adult groups where appropriate.
- A bibliography at the end of each chapter representing a composite of selected references from various disciplines and directs the clinician to information available beyond the scope of this book.
- Extensive appendices providing the clinician with additional data for everyday practice.
- Current, complete, and accurate content, which has been compiled from various multidisciplinary sources, then carefully scrutinized and continually reevaluated.

NEW INFORMATION IN THE SEVENTH EDITION

The addition of many new tests and methodologies includes:

- Newborn screening for inherited disease
- Updated Pap smears and protocols for further testing
- Cytokines
- Metabolic autopsies
- Tissue (histology) biopsies and predictive markers for treatment response
- Tests for bone disease
- Tests for heart disease, congestive and acute MI disease
- Microbiological testing, bioterrorism agents, detecting food poisoning, anthrax, plague, and hemorrhagic fever
- Breast diagnostic and prognostic markers
- Fetal predictive tests of abnormal development
- Breath tests for ulcers, alcohol, lactose, etc.
- Fertility tests
- Expanded scope of magnetic resonance (MRI) scans
- Expanded scope of sleep/sleepiness studies in newborns, children, older adults
- New nuclear tumor and infection scans
- PET scans combined with CT spiral imaging and ultrasound
- Ductal lavage for determining Gail Index for breast cancer risk
- New sentinel node localization
- LEEP GYN procedure
- Eye tests for retinal disorders, macular degeneration, visual acuity, and glaucoma
- Expanded content on keeping records of diagnostic tests, use of proper forms, and standardized patient reports
- Panels of multiple tests (e.g., metabolic syndrome, syndrome X) within Chapter 6 Chemistry Tests

The appendices are completely revised and contain many additions. For example, Appendix D offers information regarding collection of saliva, breath, nail, sputum, and hair specimens. Appendix H provides examples of commonly used forms and infrequently used forms (videotaping, refusal). Appendix L deals with guidelines for collecting evidentiary specimens.

Revised chapters include changes in the clinician's role and reflect current laboratory and diagnostic practice standards.

Throughout the text, a greater emphasis is placed upon communication skills and collaboration between patients, their significant others, and health professionals from diverse disciplines. When clinicians see patients in the context of what the patient and loved ones are experiencing (i.e., situational needs, expectations, previous experiences, and the environment in which they live), only then can they offer meaningful support and care. When patients believe the clinician is on their side, they have an increased sense of control. Identifying with the patient's point of view leads to a more profound level of communication.

CURRENT DEVELOPMENTS IN LABORATORY AND DIAGNOSTIC TESTING

New technologies foster new scientific modalities for patient assessment and clinical interventions. Thus, the clinician is provided a greater understanding of the long chain of events from diagnosis through treatment and outcomes. In a brief span of years, new technologies have introduced greatly improved developments in total body and brain x-ray scanners; digital and enhanced imaging; magnetic resonance (MR); positron emission tomography (PET) scanners, combination scans such as PET and CT to diagnose cancer and infections; greatly enhanced ultrasound and nuclear medicine procedures; genetic mutation studies; new tests for cancer; new cancer markers for diagnosis and prognosis; sleep disorders tests; technology for fetal testing before birth, and postmortem testing after death. Many new technologies are faster, more patient-friendly, more comfortable, and provide an equivalent or higher degree of accuracy (i.e., HIV or hepatitis detection, monitoring for drug abuse or managing therapeutic drug levels). Saliva and breath testing is gaining ground as a mirror of body function and emotional, hormonal, immune, and neurologic status, as well as providing clues about faulty metabolism. Noninvasive and minimally invasive testing, (i.e., need only one drop of blood, nail and hair clippings), is better suited for testing in environments such as the workplace, private home, and other nontraditional health care settings such as churches, is made possible by better collection methods and standardized collection techniques. Newest diagnostic lab technologies include hand-held nucleic acid detectors for specific bacteria and viruses, hand-held miniaturized chip-based DNA analyzers, reagentless diagnostics that introduce the sample (hand, finger, ear lobe, etc.) to magnetic fields, and magnetic resonance spectroscopy (MRS). Non-invasive and minimally invasive diagnostics include infrared light to estimate glucose, rapid oral screen for HIV, proteomics, functional and molecular technologies. Managed care and its drive for control of costs for diagnostic services exerts a tremendous effect on consumers' ability to access testing services care. This results in mixed access to services, depending upon approval or denial of coverage.
A resurgence in the use of traditional, trusted diagnostic modalities, such as electroencephalogram (EEG), is being seen in certain areas. Diseases such as HIV, antibiotic-resistant strains of pathological organisms, and Type 2 diabetes are becoming more prevalent. In the workplace, thorough diagnostic testing is more common as applications are made for disability benefits. Also, requirements for periodic monitoring of exposures to potentially hazardous workplace substances (chemicals, heavy metals), breathing and hearing tests, and TB and latex allergy testing requires skill in administering and procuring specimens. The number of forensic DNA tests being performed has increased tremendously. Concurrently, consumer perceptions have shifted from implicit faith in the health care system to concerns regarding less control over choices for health care and more distrust of the system in general.

These trends—combined with a shift in diagnostic care from acute care hospital settings to outpatient departments, physicians' offices, clinics, community-based centers, nursing homes, and sometimes even churches, stores and pharmacies—challenge clinicians to provide standards-based, safe, effective, and informed care. Because the health care system is becoming a community-based model, the clinician's role is also changing. Updated knowledge and skills, flexibility, and a heightened awareness of the testing environment (point of care testing) are needed to provide diagnostic services in these settings.

Clinicians must also adapt their practice to changes in other areas. This includes developing, coordinating, and following policies and standards set forth by institutions, governmental bodies, and regulatory agencies. Being informed regarding ethical and legal implications of such things as informed consent, privacy, patient safety, the right to refuse tests, end-of-life decisions, and trends in diagnostic research procedures add another dimension to the clinician's accountability and responsibility. The consequences of certain types of testing (ie, HIV and genetic) and the implications of confidential versus anonymous testing must also be kept in mind. For example, anonymous tests do not require the individual to give his or her name, whereas confidential tests do require the name. This difference has implications in the requirements and process of agency reporting all patients as well as for select groups of infectious diseases such as HIV.

Responding to these trends, the Seventh edition of *A Manual of Laboratory and Diagnostic Tests* is a comprehensive, up-to-date diagnostic reference source that includes information about newer technologies, together with the time-honored classic tests that continue to be an important component of diagnostic work. It meets the needs of clinicians, educators, researches, students, and others whose work and study requires this type of resource or reference manual.

Frances Talaska Fischbach
It is with sincere gratitude and pleasure that I acknowledge the collaboration of Dr. Marshall B. Dunning for his diligence, extra effort, and graciousness in accomplishing the task of renewal and enhancement for the revision of this text, for the 7th edition, all in a timely manner.

I want to give special praise and recognition to my husband, Jack Fischbach, the best researcher I have ever had; to Corrinne Strandell, Mary Pat Schmidt, Bernice DeBoer, Pat Pomohac, and Jean Schultz for their dedication, kindness, support, and generous help in manuscript preparation; to Kathie Gordon, Kathleen Dunning, Deanne Shmitz, and Margaret Fischbach, for carefully arranging, organizing, and typing the manuscript.

I would also like to acknowledge and thank all the reviewers, researchers, and consultants who provided ideas for manuscript revision and whose comments to me have helped make the book better. This work would not have been complete without the help and information provided by the librarians and staff of the Todd Wehr Library of the Medical College of Wisconsin, the Marquette University Library, and St. Joseph's Hospital Library; with thanks to Dynacare Laboratories and Medical Science Laboratories, especially for referencing their Laboratory Handbooks, and to the Infection Control Staff, Neuroscience Center, Transplant Services, Transfusion Services, Eye Institute, at Froedtert Memorial Hospital of Milwaukee, Wisconsin.

Appreciation and recognition are also due these persons who helped with this and previous editions: my daughters, Mary Fischbach Johnson, BS, MS Ed, and Margaret Fischbach, BA, JD; my son-in-law, Richard Johnson, BA; my daughter-in-law, Ann Shafranski Fischbach, BSN; and the hard work on this edition and in the past of the entire staff at Lippincott Williams & Wilkins, especially Sharon Nowak, Marie Rim, Quincy McDonald, Debra Schiff, Kim Lilly, Kathie Barrie, and, as always, Jay Lippincott. Writing a book is truly a labor of love, and the process makes me humble and thankful to many, many individuals, named and unnamed, who have made it possible. Thanks for a job well done.

Frances Fischbach
OVERVIEW OF THE CLINICIAN'S ROLE: RESPONSIBILITIES, STANDARDS, AND REQUISITE KNOWLEDGE

In this era of high technology, health care delivery involves many different disciplines and specialties. Consequently, clinicians must have an understanding and working knowledge of modalities other than their own area of expertise. This includes diagnostic evaluation and diagnostic services. Laboratory and diagnostic tests are tools to gain additional information about the patient. By themselves, these tests are not therapeutic; however, when used in conjunction with a thorough history and physical examination, these tests may confirm a diagnosis or provide valuable information about a patient's status and response to therapy that may not be apparent from the history and physical examination alone. Generally, a tiered approach to selecting tests is used:

1. Basic screening (frequently used with wellness groups and case finding)
2. Establishing (initial) diagnoses
3. Differential diagnosis
4. Evaluating current medical case management and outcomes
5. Evaluating disease severity
6. Monitoring course of illness and response to treatment
7. Group and panel testing
8. Regularly scheduled screening tests as part of ongoing care
9. Testing related to specific events, certain signs and symptoms, or other exceptional situations (e.g., infection and inflammation [bladder infection or cellulitis], sexual assault, drug screening, pheochromocytoma, postmortem tests, to name a few) (Table 1.1)

Table 1.1 Examples of Selecting Tests

<table>
<thead>
<tr>
<th>Diagnostic Test</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stool occult blood</td>
<td>Yearly screening after 45 years of age</td>
</tr>
<tr>
<td>Serum potassium</td>
<td>Yearly in patients on diuretic agents or potassium supplements; in cases of some cardiac arrhythmias</td>
</tr>
<tr>
<td>Liver enzyme levels</td>
<td>Monitoring patient on hepatotoxic drugs; establish baseline values</td>
</tr>
<tr>
<td>Serum amylase</td>
<td>In the presence of abdominal pain, suspect pancreatitis</td>
</tr>
<tr>
<td>Thyroid-stimulating hormone (TSH) test</td>
<td>Suspicion of hypothyroidism, hyperthyroidism, or thyroid dysfunction, 50 years of age and older</td>
</tr>
<tr>
<td>Chlamydia and gonorrhea</td>
<td>In sexually active persons with multiple partners to monitor for pelvic inflammatory disease</td>
</tr>
<tr>
<td>Hematocrit and hemoglobin</td>
<td>Baseline study; abnormal bleeding; detection of anemia (use CBC results if they are recent)</td>
</tr>
<tr>
<td>Papanicolaou cervical smear (Pap)</td>
<td>Yearly for all women = 18 years of age; more often with high-risk factors (e.g., dysplasia, human immunodeficiency virus [HIV], herpes simplex) now checks for human papillomavirus (HPV), chlamydia, and gonorrhea, using DNA</td>
</tr>
<tr>
<td>Urine culture</td>
<td>Pyuria</td>
</tr>
<tr>
<td>Syphilis serum fluorescent treponemal antibody (FTA) test</td>
<td>Positive rapid plasma reagin (RPR) test result</td>
</tr>
<tr>
<td>Tuberculosis (TB) skin test</td>
<td>Easiest test to use for TB screening of individuals &lt; 35 years of age or those with history of negative TB skin tests, for persons in resident homes</td>
</tr>
<tr>
<td>Fasting blood glucose (FBG)</td>
<td>Every 3 years starting at 45 years of age; monitor diabetes control</td>
</tr>
<tr>
<td>Urinalysis (UA)</td>
<td>Signs or history of recurrent urinary tract disease; pregnant women; men with prostatic hypertrophy</td>
</tr>
<tr>
<td>Prothrombin time (PT) (INR)</td>
<td>Monitoring anticoagulant treatment</td>
</tr>
<tr>
<td>Prostate-specific antigen (PSA) and digital rectal examination</td>
<td>Screen men = 50 years of age for prostate cancer yearly</td>
</tr>
<tr>
<td>Chest x-ray</td>
<td>Monitor for lung lesions and infiltrates; congestive heart failure; anatomic deformities, posttrauma, before surgery, follow-up for positive TB skin test and monitor treatment</td>
</tr>
<tr>
<td>Mammogram</td>
<td>Screen by 40 years of age in women, then every 12–18 months between 40 and 49 years of age, annually = 50 years of age; follow-up for history and treatment of breast cancer; routine screening when strong family history of breast carcinoma</td>
</tr>
<tr>
<td>Colon x-rays and proctosigmoidoscopy</td>
<td>Screen adults for colon cancer beginning at age 45; follow up for presence of hemoglobin- or guaiac-positive stools, polyps, diverticulosis</td>
</tr>
<tr>
<td>Computed tomography (CT) scans</td>
<td>Before and after treatment for certain cancers, injuries, illness (e.g., suspected transient ischemic attack, cerebro-vascular accident; diagnostic evaluation of certain signs/symptoms)</td>
</tr>
<tr>
<td>DNA testing of hair, blood, skin tissue, or semen samples</td>
<td>To gather postmortem evidence, in certain criminal cases; to establish identity and parentage</td>
</tr>
</tbody>
</table>

Some tests are mandated by government agencies or clinical practice guidelines of professional societies; others are deemed part of necessary care based on the individual practitioner's judgment and expertise or a group practitioner consensus. There is not a consensus as to the frequency of testing (e.g., annually or after a certain age).

Test selections are based on subjective clinical judgment. Often diagnostic tests or procedures are used as predictors of surgical risk and/or morbidity and mortality rates (e.g., maximum oxygen consumption determination to assess risk before esophageal cancer surgery) as the risk may outweigh the benefit. Use of evidence-based guidelines for scheduling, selecting, retaining, or eliminating certain diagnostic tests may help in more effective case management and cost containment. These guidelines use a system that grades the quality of scientific evidence based on published reports of clinical trials, expert consensus, or clinical expertise. Levels of evidence are A to C and E, with A being the best evidence and E referring to expert opinion or consensus (Chart 1.1).

Education Alert

Not all information on the Internet is reliable.
Chart 1.1 Grading Guidelines for Scientific Evidence

<table>
<thead>
<tr>
<th>Grading</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Clear evidence from all appropriately conducted trials</td>
<td>A. Measure plasma glucose through an accredited lab to diagnose or screen for diabetes</td>
</tr>
<tr>
<td>B. Supportive evidence from well-conducted studies or registries</td>
<td>B. Draw fasting blood plasma specimens for glucose analysis</td>
</tr>
<tr>
<td>C. No published evidence; or only case, observational, or historical evidence</td>
<td>C. Self-monitoring of blood glucose may help to achieve better control</td>
</tr>
<tr>
<td>D. Expert consensus or clinical experience or Internet polls</td>
<td>D. Measure ketones in urine or blood to monitor and diagnose diabetic ketoacidosis (DKA) (in home or clinic)</td>
</tr>
</tbody>
</table>

As an integral part of their practice, clinicians have long supported patients and their significant others in meeting the demands and challenges incumbent in the simplest to the most complex diagnostic testing. This testing begins before birth and frequently continues after death. The clinician who provides diagnostic services must have basic requisite knowledge to plan patient care and an understanding of psychoneuroimmunology (effects of stress on health status), must make careful judgments, and must gather vital information about the patient and the testing process, to diagnose appropriately within the parameters of the clinician’s professional standards (Table 1.2; Chart 1.2).

Table 1.2 Examples of Inappropriate Tests and Replacement Tests

<table>
<thead>
<tr>
<th>Inappropriate</th>
<th>Replacement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prostatic acid phosphatase</td>
<td>PSA or free PSA</td>
</tr>
<tr>
<td>Ammonia</td>
<td>AST, GGT</td>
</tr>
<tr>
<td>Crossmatch (needed if blood is actually to be given)</td>
<td>Type and screen</td>
</tr>
<tr>
<td>Calcium</td>
<td>Ionized calcium</td>
</tr>
<tr>
<td>CBC</td>
<td>Hemogram</td>
</tr>
<tr>
<td>HCV antibody</td>
<td>HCV RNA by PCR</td>
</tr>
<tr>
<td>Iron</td>
<td>Ferritin</td>
</tr>
<tr>
<td>Lupus cell</td>
<td>ANA</td>
</tr>
<tr>
<td>Creatinine</td>
<td>Urea</td>
</tr>
<tr>
<td>CRP</td>
<td>ESR</td>
</tr>
</tbody>
</table>

PSA, prostate-specific antigen; AST, aspartate transaminase; GGT, gamma-glutamyltransferase; CBC, complete blood count; HCV, hepatitis C virus; PCP, polymerase chain reaction; ANA, antinuclear antibody; CRP, C-reactive protein; ESR, erythrocyte sedimentation rate.

Chart 1.2 Basics of Informed Care

- Manage testing environment using collaborative approach
- Communicate effectively and clearly
- Prepare the patient properly
- Follow standards
- Consider culture, gender, and age diversity
- Measure and evaluate outcomes; modify treatment as necessary
- Manage effective diagnostic services using team approach
- Interpret, treat, monitor, and counsel about abnormal test outcomes
- Maintain proper test records

The diagnostic testing model incorporates three phases: pretest, intratest, and posttest (Fig. 1.1). The clinical team actively interacts with the patient and his or her significant others throughout each phase. The following components are included with each laboratory or diagnostic test in this text:
FIGURE 1.1 Model for the role of the clinical team in diagnostic care and services.

<table>
<thead>
<tr>
<th>Pretest Interventions</th>
<th>Intratest Interventions</th>
<th>Posttest Interventions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Test background information</td>
<td>1. Actual description of procedures</td>
<td>1. Patient aftercare</td>
</tr>
<tr>
<td>2. Normal (reference values)</td>
<td>2. Specimen collection and transport results</td>
<td>2. Clinical, education, and procedure alerts</td>
</tr>
<tr>
<td>4. Indications for testing</td>
<td>4. Interfering factors</td>
<td>4. Interpretation of test results</td>
</tr>
</tbody>
</table>

Each phase of testing requires that a specific set of guidelines and standards be followed for accurate, optimal test results. Patient care standards and standards of professional practice are key points in developing a collaborative approach to patient care during diagnostic evaluation. Standards of care provide clinical guidelines and set minimum requirements for professional practice and patient care. They protect the public against less-than-quality care (Table 1.3).

Table 1.3 Standards for Diagnostic Evaluation

<table>
<thead>
<tr>
<th>Source of Standards for Diagnostic Service</th>
<th>Standards for Diagnostic Testing</th>
<th>Examples of Applied Standards for Diagnostic Testing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Professional practice parameters of American Nurses Association (ANA), American Medical Association (AMA), American Society of Clinical Pathologists (ASCP), American College of Radiology, Centers for Disease Control and Prevention (CDC), JCAHO health care practice requirements</td>
<td>Use a model as a framework for choosing the proper test or procedure and in the interpretation of test results. Use laboratory and diagnostic procedures for screening, differential diagnoses, follow-up, and case management.</td>
<td>Test strategies include single tests or combinations/panels of tests. Panels can be performed in parallel, series, or both.</td>
</tr>
<tr>
<td>The guidelines of the major agencies, such as American Heart Association, Cancer Society, and American Diabetes Association</td>
<td>Order the correct test, appropriately collect and transport specimens. Properly perform tests in an accredited laboratory or diagnostic facility. Accurately report test results. Communicate and interpret test findings. Treat or monitor the disease and the course of therapy. Provide diagnosis as well as prognosis.</td>
<td>Patients receive diagnostic services based on a documented assessment of need for diagnostic evaluation. Patients have the right to necessary information, benefits, or rights, to enable them to make choices and decisions that reflect their need or wish for diagnostic care.</td>
</tr>
<tr>
<td>Individual agency and institution policies and procedures and quality-control criteria for specimen collection, procedure statement for monitoring the patient after an invasive procedure, and policy for universal witnessed consent situations. Statements on quality improvement standards. Use standards of professional practice and standards of patient care. Use policy for obtaining informed consent/witnessed consent. Use policies for unusual situations.</td>
<td>Observe standard precautions (formerly known as universal precautions). Use latex allergy protocols and required methodology of specimen collection. Use standards and statements for monitoring patients who receive conscious sedation and analgesia. Vital signs are monitored and recorded at specific times before and after the procedure. Patients are monitored for bleeding and respiratory or neurovascular changes. Record data regarding outcomes when defined care criteria are implemented and practiced. Protocols to obtain appropriate consents are employed, and deviations from basic consent policies are documented and reported to the proper individual.</td>
<td>The clinician wears protective eyewear and gloves when handling all body fluids and employs proper handwashing before and after handling specimens and between patient contacts. Labeled biohazard bags are used for specimen transport. Vital signs are monitored and recorded at specific times before and after the procedure. Patients are monitored for bleeding and respiratory or neurovascular changes. Record data regarding outcomes when defined care criteria are implemented and practiced. Protocols to obtain appropriate consents are employed, and deviations from basic consent policies are documented and reported to the proper individual.</td>
</tr>
</tbody>
</table>
State and federal government communicable disease reporting regulations; Centers for Disease Control and Prevention (CDC), U.S. Department of Health and Human Services, Agency for Health Care Policy and Research (AHCPR), and Clinical Laboratory Improvement Act (CLIA)

Clinical laboratory personnel and other health care providers follow regulations to control the spread of communicable diseases by reporting certain disease conditions, outbreaks, and unusual manifestations, morbidity, and mortality data. Findings from research studies provide health care policy makers with evidence-based guidelines for appropriate selection of tests and procedures.

U.S. Department of Transportation

Alcohol testing is done in emergency rooms in special situations (eg, following a motor vehicle accident, homicide, or suicide, or an unconscious individual).

Occupational Safety and Health Administration (OSHA)

Workplace testing

The clinician reports laboratory evidence of certain disease classes (eg, sexually transmitted diseases, diphtheria, Lyme disease, symptomatic HIV infection; see list of reportable diseases). Personnel with hepatitis A may not handle food or care for patients, young children, or the elderly for a specific period of time. Federal government regulates shipment of diagnostic specimens. MR and CT are used to evaluate persistent low back pain according to AHCPR guidelines. Properly trained personnel perform blood, saliva, and breath alcohol testing and use required kits as referenced by federal law.

JCAHO, Joint Commission on Accreditation of Healthcare Organizations; HIV, human immunodeficiency virus; MR, magnetic resonance; CT, computed tomography.

If test results are inconclusive or negative and no definitive medical diagnosis can be established, other tests and procedures may be ordered. Thus, testing can become an involved and lengthy process (see Fig. 1.1).

Understanding the basics of safe, effective, and informed care is important. These basics include assessing risk factors and modifying care accordingly, using a collaborative approach, following proper guidelines for procedures and specimen collection, and delivering appropriate care throughout the process. Providing reassurance and support to the patient and his or her significant others, intervening appropriately, and clearly documenting patient teaching, observations, and outcomes during the entire process are important (see Fig. 1.1).

A risk assessment before testing identifies risk-prone patients and helps to prevent complications. The following factors increase a patient's risk for complications and may affect test outcomes:

1. Age > 70 years
2. History of falls
3. History of serious chronic illnesses
4. History of allergies (eg, latex, contrast iodine, radiopharmaceuticals, and other medications)
5. Infection or increased risk for infection (eg, human immunodeficiency virus [HIV], organ transplantation, chemotherapy, radiation therapy)
6. Aggressive or antisocial behavior
7. Seizure disorders
8. Uncontrolled pain
9. Gastric motility dysfunction
10. Use of assistive devices for activities of daily living (ADLs)
11. Unsteady gait, balance problems
12. Neuromuscular conditions
13. Weakness, fatigability
14. Paresthesias
15. Impaired judgment or illogical thinking
16. Severe visual problems
17. Hearing impairment
18. Use of diuretics, sedatives, analgesics, or other prescription or over-the-counter (OTC) drugs
19. Alcohol or illegal drug use or addiction

The environments in which diagnostic services are provided, the degree of cultural diversity present in the community, and the physical, emotional, social, and spiritual state of the patient all influence the patient's response to the procedure. Including the patient's significant others is a vital component of the entire process and must not be taken lightly or casually dismissed.

Testing environments vary. Certain tests (eg, cholesterol screening, blood glucose, electrocardiogram [ECG], lipid profiles, tuberculosis [TB] skin tests) can be done “in the field,” meaning that the service is brought to the patient's environment. Other tests (eg, x-rays using contrast media and those that require special patient preparation, invasive procedures, nuclear medicine procedures, hormone levels, and 24-hour urine testing panels) must be done in a physician's office, clinic, or hospital setting. Magnetic resonance (MR) imaging and ultrasound procedures (eg, echocardiograms) are commonly performed in freestanding or specialty diagnostic centers. Complex tests such as endoscopic retrograde cholangiopancreatography (ERCP), cardiac catheterization, or bronchoscopy may require hospital admission or at least outpatient status. As testing equipment becomes more technologically sophisticated and risks associated with testing are reduced, the environment in which diagnostic procedures take place will also shift. Insurance reimbursement for testing also influences trends. Managed care and case management, together with collaboration among the diverse health care disciplines and the patient, are key factors in determining how and to what degree optimal
diagnostic services are used. Clear, timely, accurate communication among all patients and professionals is key to minimizing problems and frustrations.

As societies become more culturally blended, the need to appreciate and work within the realm of cultural diversity becomes imperative. Interacting with patients and directing them through diagnostic testing can present certain challenges if one is not familiar and sensitive to the health care belief system of the patient and his or her significant others. Something as basic as attempting to communicate in the face of language differences may necessitate arrangements for a relative or translator to be present during all phases of the process. Special attention and communication skills are necessary for these situations as well as when caring for children and for comatose, confused, or frail patients. Consideration of these issues will significantly influence compliance, outcomes, and positive responses to the procedure. To be most effective, professional care providers must be open to a holistic perspective and attitude that affects their care giving, communication, and patient-empowering behaviors. Clinicians who understand the patient’s basic needs and expectations and strive to accommodate those as much as possible are truly acting as patient advocates.

Preparing patients for diagnostic or therapeutic procedures, collecting specimens, carrying out and assisting with procedures, and providing follow-up care have long been requisite activities of professional practice. This care may continue even after the patient’s death. Diagnostic postmortem services include death reporting, possible postmortem investigations, and sensitive communication with grieving families and significant others regarding autopsies, unexplained death, other postmortem testing, and organ donation (see Chap. 16).

Professionals need to work as a team to meet diverse patient needs, to facilitate certain decisions, to develop comprehensive plans of care, and to help patients modify their daily activities to meet test requirements in all three phases. It is a given that institutional protocols are followed.

**PRETEST PHASE: ELEMENTS OF SAFE, EFFECTIVE, INFORMED CARE**

The emphasis of pretest care is on appropriate test selection, obtaining proper consent, proper patient preparation, individualized patient education, emotional support, and effective communication. These interventions are key to achieving the desired outcomes and preventing misunderstandings and errors.

**Basic Knowledge and Necessary Skills**

Know the test terminology, purpose, process, procedure, and normal test reference values or results. The names of diseases are a convenient way of briefly stating the endpoint of a diagnostic process that begins with assessment of symptoms and signs and ends with knowledge of causation and detection of underlying disorders of structure and function.

The clinical value of a test is related to its sensitivity, specificity, and the incidence of the disease in the population tested. Sensitivity and specificity do not change with different populations of ill and healthy patients. The *predictive value* of the same test can vary significantly with age, gender, and geographic location.

**Specificity** refers to the ability of a test to identify correctly those individuals who do not have the disease. The division formula for specificity is as follows:

\[
\% \text{ of specificity} = \frac{\text{persons without disease who test negative}}{\text{divided by the total number of persons without the disease}} \times 100
\]

**Sensitivity** refers to the ability of a test to correctly identify those individuals who truly have the disease. The division formula for sensitivity is as follows:

\[
\% \text{ of sensitivity} = \frac{\text{persons with disease who test positive}}{\text{divided by the total number of persons with the disease}} \times 100
\]

**Incidence** refers to the prevalence of a disease in a population or community. The predictive value of the same test can be very different when applied to people of differing ages, genders, geographic locations, and cultures.

**Predicted Values** refer to the ability of a screening test result to correctly identify the disease state. *True-positive results* correctly identify individuals who actually have the disease, and *true-negative results* correctly identify individuals who do not actually have the disease. *Positive predictive value* equals the percentage of positive tests with true-positive results (ie, the individual does have the disease). *Negative predictive value* refers to the percentage of negative tests with true-negative results (ie, the individual does not have the disease).

See Table 1.4 for an example that demonstrates the specificity, sensitivity, and predictive values for a new screening test to identify the cystic fibrosis gene.

<table>
<thead>
<tr>
<th>Test Result Have Gene for Cystic Fibrosis Do Not Have Gene for Cystic Fibrosis Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive</td>
</tr>
<tr>
<td>Negative</td>
</tr>
<tr>
<td>TOTAL</td>
</tr>
</tbody>
</table>
Thus, this new screening test will give a false-negative result about 20% of the time (e.g., the person does have the cystic fibrosis gene but his or her test results are negative).

\[
\text{positive predictive value} = \frac{62}{67} \times 100 = 92.5\% 
\]

Thus, there is about an 8% chance that the person will test positive for the cystic fibrosis gene but does not have it.

\[
\text{negative predictive value} = \frac{341}{356} \times 100 = 95.7\% 
\]

Thus, there is about a 5% chance that the person will test negative for the cystic fibrosis gene but actually does have it.

Look at both current and previous test results and review the most recent laboratory data first, then work sequentially backward to evaluate trends or changes from previous data. The patient’s plan of care may need to be modified because of test results and changes in medical management.

Testing Environments

Diagnostic testing occurs in many different environments. Many test sites have shifted into community settings and away from hospitals and clinics.

Point-of-Care Testing refers to tests done in the primary care setting. In acute care settings (e.g., critical care units, ambulances), state-of-the-art testing can produce rapid reporting of test results.

Testing in the home care environment requires skill in procedures such as drawing blood samples, collecting samples from retention catheters, proper specimen labeling, documentation, specimen handling, and specimen transporting. Moreover, teaching the patient and his or her significant others how to collect specimens is an important part of the process.

In occupational health environments, testing may be done to reduce or prevent known workplace hazards (e.g., exposure to lead) and to monitor identified health problems. This can include preemployment baseline screening, periodic monitoring of exposure to potentially hazardous workplace substances, and drug screening. Skill in drawing blood samples, performing breathing tests, monitoring chain of custody (see page 226 in Chap. 3), and obtaining properly signed and witnessed consent forms for drug testing is required.

More pretest, posttest, and follow-up testing occurs in nursing homes because patients are more frequently taken or transferred to hospitals for more complex procedures (e.g., computed tomography [CT] scans, endoscopies), whereas this is not the case with routine testing. Increasing numbers of “full code” (i.e., resuscitation) orders leads to greater numbers and varieties of tests. Additionally, confused, combative, or uncooperative behaviors are seen more frequently in these settings. An attitude adopted by nursing home patients of “not wanting to be bothered” or engaging in outright refusal to undergo prescribed tests can make testing difficult. Consequently, understanding patient behaviors and using appropriate communication strategies and interventions for this population are necessary skills for practicing in this arena.

For those who practice in the realm of public health, diagnostic test responsibilities focus on wellness screenings, preventive services, disease control, counseling, and treatment of individuals with problems. Case finding frequently occurs at health fairs, outreach centers, homeless shelters, neighborhood nurse offices, mobile health vans, and church settings. Responsibilities vary according to setting and may include providing test information, procuring specimens, and providing referrals to appropriate caregivers. These responsibilities may even extend to transporting and preparing specimens for analysis or actually performing specimen analysis (e.g., stool tests for occult blood, TB skin testing, and procuring blood or saliva samples for HIV/acquired immunodeficiency syndrome [AIDS] testing).

History and Assessment

Obtain a relevant, current health history; perform a physical assessment if indicated. Identify conditions that could influence the actual testing process or test outcomes (e.g., pregnancy, diabetes, cultural diversity, language barrier, physical impairment, altered mental state).

1. Perform a risk assessment for potential injury or noncompliance.
2. Identify contraindications to testing such as allergies (e.g., iodine, latex, medications, contrast media). Records of previous diagnostic procedures may provide clues.
3. Assess for coping styles and knowledge or teaching needs.
4. Assess fears and phobias (eg, claustrophobia, “panic attacks,” fear of needles and blood). Ascertain what strategies the patient uses to deal with these reactions and try to accommodate these.

5. Observe standard/universal precautions with every patient (see Appendix A). A patient may choose not to disclose drug or alcohol use or HIV and hepatitis risks.

6. Document relevant data. Address patient concerns and questions. This information adds to the database for collaborative problem-solving activities among the medical, laboratory/diagnostic, and nursing disciplines.

**Reimbursement for Diagnostic Services**

Differences in both diagnostic care services and reimbursement may vary between private and government insurance. Nonetheless, quality of care should not be compromised in favor of cost reduction. Advocate for patients regarding insurance coverage for diagnostic services. Inform the patient and his or her family or significant others that it may be necessary to check with their insurance company before laboratory and diagnostic testing to make certain that costs are covered.

Many insurance companies employ case managers as gatekeepers for monitoring costs, diagnostic tests ordered, and other care. As a result, the insurance company or third-party payer may reimburse only for certain tests or procedures or may not cover tests considered by them to be preventive care. So that reimbursement completely covers diagnostic services provided, be sure to include proper documentation and proper Common Practice Terminology (CPT) codes. Note date laboratory service is performed and date specimen is collected (must use). Based on 1999 data, Chart 1.3 lists laboratory tests that are covered by most insurance carriers, both private and government.

**Chart 1.3 Tests Covered by Most Insurance Carriers**

<table>
<thead>
<tr>
<th>Test</th>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alpha-fetoprotein</td>
<td>Human chorionic gonadotropin</td>
</tr>
<tr>
<td>Blood counts</td>
<td>Lipids</td>
</tr>
<tr>
<td>Blood glucose testing</td>
<td>Partial thromboplastin time</td>
</tr>
<tr>
<td>Carcinoembryonic antigen</td>
<td>Prostate-specific antigen</td>
</tr>
<tr>
<td>Collagen crosslinks, any method (urine osteoporosis)</td>
<td>Prothrombin time</td>
</tr>
<tr>
<td>Digoxin therapeutic drug assay</td>
<td>Serum iron studies</td>
</tr>
<tr>
<td>Fecal occult blood</td>
<td>Thyroid testing</td>
</tr>
<tr>
<td>Gamma-glutamyltransferase</td>
<td>Tumor antigen by immunoassay—CA125</td>
</tr>
<tr>
<td>Glycated hemoglobin/glycated protein</td>
<td>Tumor antigen by immunoassay—CA15-3/CA27</td>
</tr>
<tr>
<td>Hepatitis panel</td>
<td>Tumor antigen by immunoassay—CA19-9</td>
</tr>
<tr>
<td>HIV testing (diagnosis)</td>
<td>Urine culture</td>
</tr>
<tr>
<td>HIV testing (prognosis including monitoring)</td>
<td></td>
</tr>
</tbody>
</table>

**Methodology of Testing**

Follow testing procedures accurately. Verify orders and document them with complete, accurate, and legible information. Document all drugs the patient is taking because these may influence test outcomes (see Appendix J).

1. Ensure that specimens are correctly obtained, preserved, handled, labeled, and delivered to the appropriate department. For example, it is not generally acceptable to draw blood samples when an intravenous line is infusing proximal to the intended puncture site.
2. Observe precautions for patients in isolation. Use standard/universal precautions.
3. As much as possible, coordinate patient activities with testing schedules to avoid conflicts with meal times and administration of medications, treatments, or other diagnostic tests and travel time.
   a. Maintain NPO (ie, nothing by mouth) status when necessary.
   b. Administer the proper medications in a timely manner. Schedule tests requiring contrast substances in the proper sequence so as not to invalidate succeeding tests.

**Interfering Factors**

Minimize test outcome deviations by following proper test protocols. Make certain the patient and his or her significant others know what is expected of them. Written instructions are very helpful.
Reasons for deviations may include the following:

1. Incorrect specimen collection, handling, storage, or labeling
2. Wrong preservative or lack of preservative
3. Delayed specimen delivery
4. Incorrect or incomplete patient preparation
5. Hemolyzed blood samples
6. Incomplete sample collection, especially of timed samples
7. Old or deteriorating specimens

Patient factors that can alter test results may include the following:

1. Incorrect pretest diet
2. Current drug therapy
3. Type of illness
4. Dehydration
5. Position or activity at time of specimen collection
6. Postprandial status (ie, time patient last ate)
7. Time of day
8. Pregnancy
9. Level of patient knowledge and understanding of testing process
10. Stress
11. Nonadherence or noncompliance with instructions and pretest preparation
12. Undisclosed drug or alcohol use
13. Age and gender

Avoiding Errors

To avoid costly mistakes, know what equipment and supplies are needed and how the test is performed. Communication errors account for more incorrect results than do technical errors. Properly identify and label every specimen as soon as it is obtained. Determine the type of sample needed and the collection method to be used. Is the test invasive or noninvasive? Are contrast media injected or swallowed? Is there a need to fast? Are fluids restricted or forced? Are medications administered or withheld? What is the approximate length of the procedure? Are consent forms and conscious sedation, oxygen, analgesia, or anesthesia required? Report test results as soon as possible. “Critical” or “panic” values must be reported to the proper persons immediately (STAT).

Instruct patients and their significant others regarding their responsibilities. Accurately outline the steps of the testing process and any restrictions that may apply. Conscientious, clear, timely communication among health care departments can reduce errors and inconvenience to both staff and patients.

Proper Preparation

Prepare the patient correctly. This preparation begins at the time of scheduling.

1. Provide information about testing site and give directions for locating the facility; allow time to enter the facility and find the specific testing laboratory. If a copy of the written test order was given to the patient to bring to the laboratory, interpret the test order. For example, an order for a renal sonogram means that an ultrasound of the kidney will be done to “rule out” (RO) evidence or presence of abnormality or suspected problem. The terms “ultrasound” and “sonogram” are used interchangeably.
2. Plan to be at the department 15 minutes before testing if the test is scheduled for a specific time. Review all pretest instructions and be certain they are explained clearly (eg, “fasting” directions for test, tell patient what fasting actually means).
3. Be aware of special needs of those with conditions such as physical limitations or disabilities, ostomies, or diabetes; children; elderly patients; and culturally diverse patients.
4. Give simple, accurate, precise instructions according to the patient’s level of understanding. For example, the patient needs to know when and what to eat and drink or how long to fast.
5. Encourage dialogue about fears and apprehensions. “Walking” a patient through the procedure using imagery and relaxation techniques may help them to cope with anxieties. Never underestimate the value of a caring presence.
6. Assess for the patient’s ability to read and understand instructions. Poor eyesight or hearing difficulties may impair understanding and compliance. Speak slowly and clearly. Do not bombard the patient with information. Instruct the patient to use assistive devices such as eyeglasses and hearing aids if necessary. Clear, written instructions can reinforce verbal instructions and should be used whenever possible. In some cases, a translator or “signer,” or legal representative may be necessary.
7. Assess for language and cultural barriers. Patients behave according to personal values, perceptions, beliefs, traditions, and cultural and ethnic influences. Take these into consideration and value the patient’s uniqueness to the highest degree possible.
8. Document accurately in all testing phases.

Patient Education

Educate the patient and family regarding the testing process and what will be expected of them. Record the date, time, type of teaching, information given, and to whom the information was given.

1. Giving sensory and objective information that relates to what the patient will likely physically feel and the equipment
that will be used is important so that patients can “see” a realistic representation of what will occur. Avoid technical and medical jargon and adapt information to the patient's level of understanding. Slang terms may be necessary to get a point across.

2. Encourage questions and verbalization of feelings, fears, and concerns. Do not dismiss, minimize, or invalidate the patient's anxiety through trivial remarks such as “Don't worry.” Develop “listening ears and eyes” skills. Be aware of nonverbal signals (ie, body language) because these frequently provide a more accurate picture of what the patient really feels than what he or she says. Above all, be nonjudgmental.

3. Emphasize that there is usually a waiting period (ie, “turn-around time”) before test results are relayed back to the clinicians and nursing unit. The patient may have to wait several days for results. Offer listening, presence, and support during this time of great concern and anxiety.

4. Record test result information. Include the patient's response. Just because something is taught does not necessarily mean that it is learned or accepted. The possibility that a diagnosis will require a patient to make significant lifestyle changes (eg, diabetes) requires intense support, understanding, education, and motivation. Document specific names of audiovisual and reading materials to be used for audit, reimbursement, and accreditation purposes.

**Testing Protocols**

Develop consistent protocols for teaching and testing that encompass comprehensive pretest, intratest, and posttest care modalities.

Prepare patients for those aspects of the procedure experienced by the majority of patients. Clinicians can collaborate to collect data and to develop a list of common patient experiences, responses, and reactions.

**Patient Independence**

Allow the patient to maintain as much control as possible during the diagnostic phases to reduce stress and anxiety. Include the patient and his or her significant others in decision making. Because of factors such as anxiety, language barriers, and physical or emotional impairments, the patient may not fully understand and assimilate instructions and explanations. To validate the patient's understanding of what is presented, ask the patient to repeat instructions given to evaluate assimilation and understanding of presented information.

Include and reinforce information about the diagnostic plan, the procedure, time frames, and the patient's role in the testing process.

**Test Results**

Know normal or reference values.

1. Normal ranges can vary to some degree from laboratory to laboratory. Frequently, this is because of the particular type of equipment used. Theoretically, “normal” can refer to the ideal health state, to average reference values, or to types of statistical distribution. Normal values are those that fall within 2 standard deviations (ie, random variation) of the mean value for the normal population.

2. The reported reference range for a test can vary according to the laboratory used, the method employed, the population tested, and methods of specimen collection and preservation.

3. The majority of normal blood test values are determined by measuring “fasting” specimens.

4. Be aware of specific influences on test results. For example, patient posture is important when plasma volume is measured because this value is 12% to 15% greater in a person who has been supine for several hours. Changing from a supine to a standing position can alter values as follows: increased hemoglobin (Hb), red blood cell (RBC) count, hematocrit (Hct), calcium (Ca), potassium (K), phosphorus (P), aspartate aminotransferase (AST), phosphatases, total protein, albumin, cholesterol, and triglycerides. Going from an upright to a supine position results in increased hematocrit, calcium, total protein, and cholesterol. A tourniquet applied for > 1 minute produces laboratory value increases in protein (5%), iron (6.7%), AST (9.3%), and cholesterol (5%) and decreases in K⁺ (6%) and creatinine (2%–3%).

Laboratories must specify their own normal ranges. Many factors affect laboratory test values and influence ranges. Thus, values may be normal under one set of prevailing conditions but may exhibit different limits in other circumstances. Age, gender, race, environment, posture, diurnal and other cyclic variations, foods, beverages, fasting or postprandial state, drugs, and exercise can affect derived values. Interpretation of laboratory results must always be in the context of the patient's state of being. Circumstances such as hydration, nutrition, fasting state, mental status, or compliance with testing protocols are only a few of the situations that can influence test outcomes.

**Laboratory Reports**

Scientific publications and many professional organizations are changing clinical laboratory data values from conventional units to Système International (SI) units. Currently, many data are reported in both ways. The SI system uses seven dimensionally independent units of measurement to provide logical and consistent measurements. For example, SI concentrations are written as amount per volume (grams, milligrams, or milliequivalents per deciliter, 100 milliliters, or liter). Numerical values may differ between systems or may be the same. For example, chloride is the same in both systems: 95 to 105 mEq/L (conventional) and 95 to 105 mmol/L (SI) (see Appendix D).
Margins of Error

Recognize margins of error. For example, if a patient has a battery of chemistry tests, the possibility exists that some tests will be abnormal owing purely to chance. This occurs because a significant margin of error arises from the arbitrary setting of limits. Moreover, if a laboratory test is considered normal up to the 95th percentile, then 5 times out of 100, the test will show an abnormality even though a patient is not ill. A second test performed on the same sample will probably yield the following: 0.95 \times 0.95, or 90.25%. This means that 9.75 times out of 100, a test will show an abnormality even though the person has no underlying health disorder. Each successive testing will produce a higher percentage of abnormal results. If the patient has a group of tests performed on one blood sample, the possibility that some of the tests will “read abnormal” due purely to chance is not uncommon.

Ethics and the Law

Consider legal and ethical implications. These include the patient’s right to information, properly signed and witnessed consent forms, and explanations and instructions regarding chain-of-custody requirements and risks as well as benefits of tests.

1. Chain of custody is a legal term descriptive of a procedure to ensure specimen integrity from collection to transport to receipt to analysis and specimen storage. A special form is used to provide a written record. The right to informed consent before certain tests and procedures pertains to patient autonomy, the ethical right of self-determination, the legal right to be free of procedures to which one does not consent, and to determine what will be done to one’s own person. Risks, benefits, and alternatives are explained and written consent obtained well in advance of the procedure.

2. The patient must demonstrate appropriate cognitive and reasoning faculties to sign a legally valid consent. Conversely, a patient may not legally give consent while under the immediate influence of sedation, anesthetic agents, or certain classes of analgesics and tranquillizers. If the patient cannot validly and legally sign a consent form, an appropriately qualified individual may give consent for the patient.

3. Guidelines and wishes set forth in advance directives or “living will”–type documents must be honored, especially in life-threatening situations. Such directives may prevent more sophisticated invasive procedures from being performed. Some states have legislated that patients can procure do-not-resuscitate (DNR) orders and medical DNR bracelets that indicate their wishes. A copy of a patient’s advance directives in the health care record can be very helpful in unpredictable situations.

4. A collaborative team approach is essential for responsible, lawful, and ethical patient-focused care. The clinician who orders the test has a responsibility to inform the patient about risks and test results and to discuss alternatives for follow-up care. Other caregivers can provide additional information and clarification and can support the patient and family in achieving the best possible outcomes. The duty to maintain confidentiality, to provide freedom of choice, and to report infectious diseases may result in ethical dilemmas.

Respect for the dignity of the individual reflects basic ethical considerations. Patients and family have a right to consent, to question, to request other opinions, and to refuse diagnostic tests. Conversely, caregivers have the right to know the diagnoses of the patients they care for so that they can minimize the risks to themselves.

Patient’s Bill of Rights and Patient Responsibilities

Patients have a right to expect that an agency’s or institution’s policies and procedures will ensure certain rights and responsibilities for them. At all times, the patient has the right:

1. To considerate, honest, respectful care, with consideration given to privacy and maintenance of personal dignity, cultural and personal values and beliefs, and physical and developmental needs, regardless of the setting.

2. To be involved in decision making and to participate actively, if so desired, in the testing process, assuming the patient is competent to make these choices.

3. To participate in the informed consent process before testing and to be told of the benefits, risks, and reasonable alternative approaches to tests ordered.

4. To be informed regarding test costs and reimbursement responsibility.

5. To refuse diagnostic testing.

6. To expect to have the support of family or significant others, if so desired and appropriate during the testing process.

7. To expect that standards of care will be followed by all personnel involved in the testing process.

8. To expect safe, skilled, quality care provided by trained personnel with expertise in their field.

9. To expect patient and family education and instructions regarding all phases of the testing process and procedure, including the nature and purpose of the test, pretest preparation, actual testing, posttest care benefits, risks, side effects, and complications. Information should be provided in a sensitive and objective manner.

10. To expect to be informed in a timely manner of test results and implications, treatment, and future testing if necessary.

11. To expect to be counseled appropriately regarding abnormal test outcomes as well as alternative options and available treatments.

12. To expect to have acceptable pain control and comfort measures provided throughout the testing process.

13. To expect that all verbal, written, and electronic communication, medical records, and medical record transfers will be accurate and confidential. Exception: when reporting of situation is required by law (eg, certain infectious diseases, child abuse).

The patient has the following responsibilities:

1. To comply with test requirements (eg, fasting, special preparations, medications, enemas) and to inform the
To report active or chronic disease conditions that may alter test outcomes, be adversely affected by the testing process, or pose a risk to health care providers (eg, HIV, hepatitis).

3. To keep appointments for diagnostic procedures and follow-up testing.

4. To disclose drug and alcohol use as well as use of supplements and herbal products despite being informed that these products could affect test outcomes (eg, erroneous test results).

5. To disclose allergies and past history of complications or adverse reactions to tests. Example: Reaction to contrast materials.

6. To report any adverse effects attributed to tests and procedures after being advised regarding signs and symptoms of such.

7. To supply specimens that are their own.

8. To report visual or hearing impairments or inability to read, write, or understand English.

Cultural Sensitivity

Preserving the cultural well-being of any individual or group promotes compliance with testing and easier recovery from routine as well as more invasive and complex procedures. Sensitive questioning and observation may provide information about certain cultural traditions, concerns, and practices related to health. For example, the Hmong people believe the soul resides in the head and that no one should touch an adult's head without permission. Patting a Hmong child on the head may violate this belief. Health care personnel should make an effort to understand the cultural differences of populations they serve without passing judgment. Most people of other cultures are willing to share this information if they feel it will be respected. Sometimes, a translator is necessary for accurate communication.

Many cultures have diverse beliefs about diagnostic testing that requires blood sampling. For example, alarm about having blood specimens drawn or concerns regarding the disposal of body fluids or tissue may require health care workers to demonstrate the utmost patience, sensitivity, and tact when communicating information about blood tests.

INTRATEST PHASE: ELEMENTS OF SAFE, EFFECTIVE, INFORMED CARE

Basic Knowledge and Required Skills

Intratest care focuses on specimen or tissue collection, monitoring the testing environment tissue collection, performing and/or assisting with procedures, providing emotional and physical comfort and reassurance, administering analgesics and sedatives, and monitoring vital signs and other parameters during testing. The clinician must have basic knowledge about the procedure and test and should have the required skills to perform testing or to assist in the process. Safe practices, proper collection of specimens, minimizing delays, providing support to the patient, preparing or administering analgesia and sedatives, monitoring various parameters as necessary, and being alert to potential side effects or complications are integral activities of the intratest phase. Invasive procedures place patients at greater risk for complications and require ongoing vigilance and observation. Monitoring fluid intake and loss, body temperature, and respiratory and cardiovascular systems and treating problems in these domains require critical thinking and quick responses.

Infection Control

Institute accepted infection control protocols. Observe special measures and sterile techniques as appropriate. Identify patients at risk for infection. Institute strict respiratory and contact isolation as necessary. Quality assurance requires proper collection, transport, and receipt of specimens and use of properly cleaned and prepared instruments and equipment. Appendix A offers more information on standard precautions for safe practice and infection control and isolation. The term standard precautions refers to a system of disease control that presupposes each direct contact with body fluids or tissues is potentially infectious and that every person exposed to these must protect himself or herself. Consequently, health care workers must be both informed and conscientious about adhering to standard precautions and strict infection control guidelines. It goes without saying that health care workers must be scrupulous about proper hand hygiene (see Appendix A). Proper protective clothing and other devices must be worn as necessary.

Procurement and disposal of specimens according to U.S. Occupational Safety and Health Administration (OSHA) standards must be adhered to. Moreover, institutions may have procedures and policies of their own to ensure compliance (eg, specimens are to be placed directly into biohazard bags).

NOTE

Standard precautions (formerly known as universal precautions) prevail in all situations in which risk for exposure to blood, tissue, and other body fluids is even remotely possible. The terms standard precautions and universal precautions are often used interchangeably.

Collaborative Approaches

A collaborative team approach is necessary for most procedures. Clinicians must assist and understand each other's role in the procedure. Invasive procedures (such as lumbar punctures or cystoscopy) place patients at greater risk for complications and usually require closer monitoring during the test. Frequently, administration of intravenous (IV) sedation and other drugs is part of the procedure. Astute ongoing observation of the patient and critical thinking and quick decision-making skills during intense situations is a requisite for clinicians in these settings.
Risk Management

Assess for and provide a safe environment for the patient at all times. Identify patients at risk and environments that may pose a risk. Previous falls, cerebrovascular accident (CVA), neuromuscular disorders, loss of balance, or use of ambulatory and other assistive devices are contributory risk factors. Prevention of complications and management of risk factors are an important part of the intratest phase. As part of risk management, observe standard precautions and infection control protocols as necessary (see Appendix A, Appendix B, and Appendix C).

Use special care during procedures that include iodine and barium contrasts, radiopharmaceuticals, latex products, conscious sedation, and analgesia (see Chap. 9, Chap. 10, and Chap. 15 for precautions for imaging procedures.) Certain risk factors contribute to a higher incidence of adverse reactions when contrast agents and radiopharmaceuticals are used (Table 1.5).

Table 1.5 Classification of Risk Factors

<table>
<thead>
<tr>
<th>Preexisting Disorders</th>
<th>Contributing Elements</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asthma</td>
<td>Allergy</td>
</tr>
<tr>
<td>Diabetes</td>
<td>Age-related (newborn and older adults)</td>
</tr>
<tr>
<td>Liver insufficiency</td>
<td>Dehydration</td>
</tr>
<tr>
<td>Multiple myeloma</td>
<td>Frequent use of contrast agents</td>
</tr>
<tr>
<td>Pheochromocytoma</td>
<td>High dosage of contrast and radiopharmaceuticals</td>
</tr>
<tr>
<td>Renal failure</td>
<td>Previous reaction to contrast agents</td>
</tr>
<tr>
<td>Seizure history</td>
<td></td>
</tr>
</tbody>
</table>

Remove jewelry, false teeth, and other prosthetic devices as necessary. Check for NPO or fasting status if appropriate.

Specimens and Procedures

Assist with and/or conduct certain diagnostic procedures. Examples of the types of assisted procedures include endoscopy, lumbar puncture, and cardiac catheterization. Diagnostic procedures often performed independently of other medical personnel include Papanicolaou (Pap) smears, centrifugation of blood samples, ECGs, breathing tests, and pulse oximetry. For example, the pulse oximeter is used to monitor noninvasively the oxygen saturation (SpO₂); S

\[
O_2 \text{ refers to pulse oximetry, whereas } S
\]

\[
O_2 \text{ refers to arterial saturation measured on an arterial blood sample. Sensors may be applied on the index, middle, or ring finger; on the nose, earlobe, toe, or foot; and on the forehead. Be aware of factors that interfere with accurate results, such as patient movement, ambient light, electronic interference, artificial nails and polish, anemia, edema, or poor circulation to an area. Chapter 14 provides more information on pulse oximetry.}
\]

Collecting specimens and conducting procedures are the main interventions in the diagnostic pretest and intratest phases. Procure, process, transport, and store specimens properly. The community environment and health care setting in which testing takes place dictate protocols for doing this. Everyone involved in the process must have a thorough understanding of testing principles and protocols and must adhere to them to ensure accurate results.

Determine specimen type needed and method of sample procurement. Special equipment and supplies may be necessary (eg, sterile containers, special kits).

Collection by the patient requires patient cooperation, understanding, and instruction. It does not always require direct supervision. Conversely, supervised collection requires supervision of the patient by trained personnel during specimen collection. Examples of these two types of collection include a routine urine sample collected by the patient privately versus a urine sample procured in a supervised setting for drug screening.

A third method of collection requires that the clinician perform the entire collection. An example of this type of collection is aspirating a urine sample from an indwelling catheter.

Time of collection is also important. For example, results from a fasting blood glucose test versus results from a 2-hour-postprandial blood glucose test are significantly different as diagnostic parameters.

Specimens can be rejected for analysis because of factors related to the specimen itself or to the collection process (Table 1.6).
### Table 1.6 Errors in Collection

<table>
<thead>
<tr>
<th>Specimen Errors</th>
<th>Collector Errors</th>
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<tbody>
<tr>
<td>Insufficient volume</td>
<td>Transport delay</td>
</tr>
<tr>
<td>Improper type</td>
<td>Improper collection method</td>
</tr>
<tr>
<td>Insufficient number of samples</td>
<td>Wrong specimen container</td>
</tr>
<tr>
<td>Wrong transport medium or wrong or absent preservative</td>
<td>Wrong time</td>
</tr>
<tr>
<td>Air bubbles in tube</td>
<td>Incorrect storage</td>
</tr>
<tr>
<td>Storage at incorrect temperature</td>
<td>Unlabeled or mislabeled specimen and/or wrong patient identification info</td>
</tr>
<tr>
<td>Incorrect order of draw</td>
<td>Improperly completed forms or computer data entry</td>
</tr>
<tr>
<td>Do not cut test tapes in half</td>
<td>Discrepancies between test ordered and specimen collected</td>
</tr>
<tr>
<td>Improper centrifugation time</td>
<td>Failure to properly transcribe and process orders</td>
</tr>
</tbody>
</table>

*Note: Observing institutional protocols can prevent mishaps.

Blood collection is normally done by trained persons. (An exception is the self-test for blood glucose using equipment designed specifically for that purpose.) The time of collection is an important factor (eg, a sequence of samples for a cardiac panel). For example, a “peak” drug-level blood specimen is collected when highest drug concentration in the blood is expected. This type of test is used for therapeutic drug monitoring and dosing. Conversely, a “trough” sample is collected when lowest drug concentration is expected. These types of tests are used for therapeutic drug monitoring, and specimens are collected and results reported before the next scheduled dose of medication.

Legal and forensic specimens are collected as evidence (see Appendix L) in legal proceedings, criminal investigations, and after death. Examples include DNA samples and drug and alcohol levels. Factors such as chain-of-custody situations and witnessed collections may be involved.

The following list addresses some general comments about specimen collections:

1. Stool and urine collection requires clean, dry containers and kits.
2. Timed urine collection requires refrigeration and/or containers with special additives.
3. Sterile, dry containers and special kits are needed for midstream clean-catch urine specimens.
4. Oral, saliva, and sputum specimens require specific techniques and kits and, sometimes, special preservatives.
5. Blood collection equipment includes gloves, needles, collection tubes, syringes, tourniquets, needle disposal containers, lancets for skin puncture, cleansing agents or antimicrobial skin preparations, and adhesive bandages.
6. Color-coded stoppers and tubes indicate the type of additive present in the collection tube (Table 1.7).

### Table 1.7 Blood Specimen Collections

<table>
<thead>
<tr>
<th>Collection Tube Color and Additives</th>
<th>Use and Precautions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yellow-topped tube: sodium polyethylene sulfonate (SPS)</td>
<td>For collection of blood cultures; aseptic technique for blood draw; invert tube 7–10 times to prevent clot formation</td>
</tr>
<tr>
<td>Red or gold serum separator tubes (SST): no anticoagulant</td>
<td>For collecting serum samples such as chemistry analysis. SST tubes should be gently inverted (completely, end over end) 5 times after collection to ensure mixing of clot activator with blood and clotting within 30 minutes. After the 30-minute period, centrifuge promptly at designated relative centrifugal force (rcf) for 15 ± 5 minutes to separate serum from cells. Serum can be stored in gel separator tubes after centrifugation for up to 48 hours. Do not freeze SST tubes. If frozen specimen is needed, separate serum into a labeled plastic transfer vial. Serum separation tubes must not be used to obtain therapeutic drug levels because the gel may lower the values.</td>
</tr>
<tr>
<td>Red-topped (plain) tube: no anticoagulant, no additive</td>
<td>For serum chemistry, serology, blood bank, collection of clotted blood specimens</td>
</tr>
<tr>
<td>Royal blue–topped tube: without ethylenediaminetetraacetic acid (EDTA) or sodium heparin (no anticoagulant—blood will clot)</td>
<td>For aluminum, arsenic, chromium, copper, nickel, and zinc levels; tube free of trace elements</td>
</tr>
<tr>
<td>Light blue–topped tube: with sodium citrate as anticoagulant (removes calcium to prevent clotting)</td>
<td>For plasma-coagulation studies (eg, prothrombin times [PT]; PT/partial thromboplastin time [PTT] and factor assays). The tube must be allowed to fill to its capacity or an improper blood/anticoagulant ratio will invalidate coagulation test results. Invert tube 7–10 times to prevent clotting. For serum, used for most chemistry tests; these tubes should be gently inverted 5 times after collection to ensure mixing of clot activator with blood and clotting within 30 minutes. After 3-minute period, centrifuge promptly at designated rcf for 15 ± 5 minutes to separate serum from cells. Serum can be stored in gel separator tubes after centrifugation for up to 48 hours. Do not freeze SST tubes. If frozen specimen is needed, separate serum into a labeled plastic transfer vial. Serum separation tubes must not be used for therapeutic drug levels. The gel may lower values. Not for blood bank use</td>
</tr>
<tr>
<td>Gold or red marbled–topped tube: serum gel separator tube (SST)</td>
<td></td>
</tr>
</tbody>
</table>
Additives preserve the specimen, prevent deterioration and coagulation, and/or block action of certain enzymes in
Tubes with anticoagulants should be gently and completely inverted (end over end) 7 to 10 times after collection.
Do not use if you notice a defect (eg, moisture, pinholes, tears). In cases of sexual assault, special rape kits are
Store specimens properly after collecting or transport them to the laboratory immediately for processing and
Use barrier drapes as directed. For example, arthroscopy drapes are positioned with the fluid control pouch at the
Taking photographs of injuries in suspected abuse situations is another example.
Unacceptable specimens lead to increased costs and time wasted in getting results to the clinician, patient,
Operating special equipment such as video monitors for endoscopic procedures may be required in some
Maintain aseptic technique during certain procedures (eg, cystoscopy, bone marrow biopsy).

<table>
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<tr>
<th>Tube Color/Treatment</th>
<th>Use Case</th>
</tr>
</thead>
<tbody>
<tr>
<td>Light green marbled–topped tube: gel separator/lithium, heparin as anticoagulant</td>
<td>For potassium determination</td>
</tr>
<tr>
<td>Tan/brown-topped tube: with heparin as anticoagulant</td>
<td>For heparinized plasma specimens for testing lead levels (ie, lead-free tube). Invert tube 7–10 times.</td>
</tr>
<tr>
<td>Lavender-topped tube: with EDTA; removes calcium to prevent clotting</td>
<td>For whole blood and plasma, for hematology and complete blood counts (CBCs); prevents the filled tube from clotting. If the tube is less than half-filled, the proportion of anticoagulant to blood may be sufficiently altered to produce unreliable laboratory test results. Invert tube 6–8 times.</td>
</tr>
<tr>
<td>Royal blue–topped tube: no additive with EDTA or sodium heparin anticoagulant</td>
<td>For toxicology, cadmium and mercury; tube free of trace elements. Invert tube 7–10 times.</td>
</tr>
<tr>
<td>Gray-topped tube: with potassium oxalate and sodium fluoride</td>
<td>For glucose levels, glucose tolerance levels, and alcohol levels.</td>
</tr>
<tr>
<td>Plain pink tube: no additive or anticoagulant</td>
<td>For blood bank</td>
</tr>
<tr>
<td>Black tube: with sodium citrate (binds calcium)</td>
<td>For Westergren sedimentation rate</td>
</tr>
<tr>
<td>Green-topped tube: with anticoagulant heparin (sodium, lithium, and ammonium heparin)</td>
<td>For heparinized plasma specimens, plasma chemistries, arterial blood gases, and special tests such as ammonia levels, hormones, and electrolytes. Invert 7–10 times to prevent clot formation.</td>
</tr>
</tbody>
</table>

*List is arranged in sequence of draw according to NCCLS guidelines.

7. Additives preserve the specimen, prevent deterioration and coagulation, and/or block action of certain enzymes in blood cells.
8. Tubes with anticoagulants should be gently and completely inverted (end over end) 7 to 10 times after collection. This process ensures complete mixing of anticoagulants with the blood sample and prevents clot formation.
9. Store specimens properly after collecting or transport them to the laboratory immediately for processing and analysis if possible. Failure to do so may result in specimen deterioration. STAT-ordered tests should always be hand-delivered to the laboratory and then processed as STAT.
10. Unacceptable specimens lead to increased costs and time wasted in getting results to the clinician, patient, institution, and third-party payer. Exposure to sunlight, air, or other substances and warming or cooling are examples of things that can alter specimen integrity (see Appendix E). Check with the laboratory for proper storage (eg, ice, ice water, separate from ice), transport, and time limits.
11. As environments for specimen collection become more variable, modified procedures and protocols require the clinician to keep abreast of the latest information related to these factors (see Appendix E).

Equipment and Supplies
1. Use required kits, equipment, and supplies. Special kits are used for obtaining heel sticks and finger sticks, blood alcohol samples, saliva or oral fluid specimen, and urine specimens.
2. Do not use if you notice a defect (eg, moisture, pinholes, tears). In cases of sexual assault, special rape kits are required and a strict procedure, consisting of several steps, is followed.
3. Operating special equipment such as video monitors for endoscopic procedures may be required in some instances. Familiarity with current audiovisual technology is necessary.
4. Taking photographs of injuries in suspected abuse situations is another example.
5. Use barrier drapes as directed. For example, arthroscopy drapes are positioned with the fluid control pouch at the knee.
6. Maintain aseptic technique during certain procedures (eg, cystoscopy, bone marrow biopsy).

Family Presence
Involving family members in the diagnostic care process has helped families by making them active participants. Facilitating family presence may provide the opportunity to calm the patient, offer additional comfort, and reduce anxiety and fear. However, some families may find the option of observing procedures to be distressing or uncomfortable. Other patients may not want family members present. Nurses acting as patient advocates recognize the importance of supporting the patient's need for reassurance and the family's need and right to be present during diagnostic procedures. The goal is to achieve an acceptable balance between all parties.

Positioning for Procedures
Proper body positioning and alignment involves placing the patient in the best possible position for the procedure and aligning the body correctly for optimal respiratory and circulatory function. Positions include jackknife, prone, lithotomy, sitting, supine, and Trendelenburg. Using positioning devices, arranging padding, and repositioning are important interventions to prevent skin pressure and skin breakdown. The potential adverse effects of various positions, especially during lengthy procedures, include skin breakdown, venous compression, sciatic nerve injury, muscle injury, and low back strain. Necessary positioning skills include ensuring that the patient's airway, IV lines, skin integrity, and monitoring devices are not compromised and identifying those persons at potential risk for injury (eg, elderly, thin, frail, unconscious patients) before positioning. If wounds, skin breakdown, abrasions, or bruises are present before the procedure, accurately document their presence and location.

Administration of Drugs and Solutions
All drugs and solutions administered during diagnostic procedures are given according to accepted protocols. Drugs are given by mouth, by intubation, parenterally (intramuscularly, intravenously, or subcutaneously), and by local or topical skin applications. IV fluids and endoscopic irrigating fluids are commonly administered.

Be aware of the potential for adverse reactions to drugs. Before procedure begins, confirm previous drug problems with the patient before the procedure. Risks for injury are related to hypersensitivity, allergic or toxic reactions, impaired drug tolerance due to liver or kidney malfunction, extravasation of intravenous fluids, and absorption of irrigating fluids into the systemic circulation. Required skills include managing airways and breathing patterns; monitoring fluid intake and loss; monitoring body, skin, and core temperature; and observing the effects of sedation and analgesia (Appendix C) (eg, vital signs, rashes, edema). Use tape with caution, especially when skin integrity can be easily compromised, as in frail elderly patients.

Management of Environment

The main goal of environmental control is safe practice to ensure that the patient is free from injury related to environmental hazards and is free from discomfort. Be attentive to temperature and air quality; the patient's temperature; exposure to noise, radiation, latex, and noxious odors; sanitation; and cleanliness.

1. Eliminate or modify sensory stimuli (eg, noise, odors, sounds).
2. Post a PATIENT AWAKE sign if the patient is awake during a procedure or PATIENT ASLEEP for sleep studies.
3. Be sensitive to conversation among team members in the presence of the patient. At best, it can be annoying to the patient; at worst, it may be misinterpreted and have far-reaching negative effects and consequences.

Pain Control, Comfort Measures, and Patient Monitoring

Provide proper information, reassurance, and support throughout the entire procedure to allay anxiety and fear. Administer sedatives, pain medication, or antiemetics as ordered. Uphold the dignity of each patient, provide privacy, and minimize any situation that might cause embarrassment or stress. Continue monitoring throughout procedures as well as after completion, if indicated.

1. Do not permit the patient to remain disrobed any longer than necessary. Allow personal clothing and other accessories such as rings or religious medals provided they do not pose a risk or interfere with the procedure. Ensure a reasonable degree of privacy.
2. Control pain and provide comfort measures. IV conscious sedation and drugs given to reverse the effects of test medications are part of this scenario. Allow the patient to maintain as much control as possible during all testing phases without compromising safety, the process and procedure, and test integrity. If possible, plan ahead to accommodate persons with special needs such as learning disabilities, visual or hearing impairment, ostomy, or diabetes management.
3. Monitor and document vital signs and other relevant parameters (eg, pulse oximetry, ECG) throughout the procedure. Observe for problems and abnormal reactions and take appropriate measures to correct such situations. Make sure emergency equipment is readily available and functional.
4. Document the patient's response to the procedure during all phases. Also document significant events or situations that occur during testing. Record disposition of specimens.

POSTTEST PHASE: ELEMENTS OF SAFE, EFFECTIVE, INFORMED CARE

Basic Knowledge and Necessary Skills

The focus of the posttest phase is on patient aftercare and the follow-up activities, observations, and monitoring necessary to prevent or minimize complications. Evaluation of outcomes and effectiveness of care, follow-up counseling, discharge planning, and appropriate posttest referrals are the major components of this phase.

Abnormal Test Results

Report and interpret test outcomes correctly. Abnormal test patterns or trends can sometimes provide more useful information than single test outcome deviations. Conversely, single test results can be normal in patients with a proven disease or illness.

1. Recognize abnormal test results and consider the implications for the patient in both the acute and the chronic stages of the disease as well as during screening.
2. The greater the degree of test abnormality, the more likely the outcome will be more serious.
3. Consider the role of drugs when tests are abnormal. Use of OTC drugs, vitamins, iron, and other minerals may produce false-positive or false-negative test results. Patients often do not disclose all medications they use, either unintentionally or deliberately. Commonly prescribed drugs that most often affect laboratory test outcomes include anticoagulants, anticonvulsants, antibiotic or antiviral agents, oral hypoglycemics, hormones, and psychotropic drugs. Consult a pharmacist or Physicians Desk Reference (PDR) source about drugs the patient is taking (eg, current literature search, computerized data, or manufacturer's drug insert sheet) (Appendix J). Be aware that patients who are addicted to drugs or alcohol may not provide accurate, reliable information about their use of these agents. In the same vein, sometimes athletes may not disclose their use of performance-enhancing drugs.
4. Consider biocultural variations when interpreting test results. See Table 1.8 for examples of some common variations.
Biocultural Variation

Support the patient and his or her significant others in understanding and coping with positive or negative test outcomes. Recognize that “panic values” may pose an immediate threat to the patient’s health status. Report these findings when providing genetic counseling, the clinician needs to be sensitive to the implications of genetic or metabolic disorders. Informing the patient or family about the genetic defect requires special training in genetic science, family counseling skills, and an understanding of legal and ethical issues. Confidentiality and privacy of information are vital.

Devastating physical, psychological, and social consequences can result from being misdiagnosed with a serious disease because of false-positive or false-negative test results. Major alterations in lifestyles and relationships without just cause can be a consequence of these clinical aberrations (eg, misdiagnosis of HIV or syphilis).

Follow-Up Counseling

1. Counsel the patient regarding test outcomes and their implications for further testing, treatment, and possible lifestyle changes. Provide time for the patient to ask questions and voice concerns about the entire testing process.
2. Test outcome interpretation involves reassessment of interfering factors and patient compliance if the results significantly deviate from normal and previous results.
3. No test is perfect; however, the greater the degree of abnormality indicated by the test result, the more likely it is that this outcome deviation is significant or represents a real disorder.
4. Notify the patient about test results after consultation with the clinician. Treatment may be delayed if test results are misplaced or not communicated in a timely manner.
5. Help patients interpret the results of community-based testing.
6. Identify differences in the patient's view of the situation, the clinician's views about tests and disease, and the health care team’s perceptions.
7. When providing genetic counseling, the clinician needs to be sensitive to the implications of genetic or metabolic disorders. Informing the patient or family about the genetic defect requires special training in genetic science, family coping skills, and an understanding of legal and ethical issues. Confidentiality and privacy of information are vital.
8. Be familiar with crisis intervention skills for patients who experience difficulty dealing with the posttest phase.
9. Encourage the patient to take as much control of the situation as possible.
10. Recognize that the different stages of behavioral responses may last several weeks.

Table 1.8 Biocultural Considerations

<table>
<thead>
<tr>
<th>Diagnostic Test</th>
<th>Biocultural Variation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orthopedic x-rays</td>
<td>Body proportions and tendencies: African American people exhibit longer arms and legs and shorter trunks than Caucasians. African American women tend to be wider shouldered and more narrow hipped, but with more abdominal adipose tissue than Caucasian women. Caucasian men tend to exhibit more abdominal adipose tissue than do African American men. Native Americans and Asian Americans have longer trunks and shorter limbs than do African American and Caucasian people. Asian American people tend to be wider hipped and more narrow shouldered than do other peoples.</td>
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<tr>
<td>Bone density measurements</td>
<td>African American men have the densest bones, followed by African American women and Caucasian men, who have similar bone densities. Caucasian women have the least dense bones. Chinese, Japanese, and Inuit bone density is less than that of Caucasian Americans. Additionally, bone density decreases with age.</td>
</tr>
<tr>
<td>Test for glucose-6-phosphate dehydrogenase (G6PD) deficiency</td>
<td>G6PD deficiency may be the cause of hemolytic disease of newborns in Asian Americans and those of Mediterranean descent. Three G6PD variants occur frequently: type A is common in African Americans (10% of males); the Mediterranean type is common in Iraqis, Kurds, Lebanese, and Sephardic Jews; and the Mahedial type is common in Southeast Asians (22% of males).</td>
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<td>Cholesterol levels</td>
<td>African American and Caucasian ethnic groups have similar cholesterol levels at birth. During childhood, African American people develop higher levels than do Caucasian people; however, African American adults have lower cholesterol levels than do Caucasian adults.</td>
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<tr>
<td>Hemoglobin/hematocrit levels</td>
<td>The normal hemoglobin level for African American people is 1 g lower than that for other groups. Given similar socioeconomic conditions, Asian Americans and Mexican Americans have hemoglobin/hematocrit levels higher than those of Caucasian people.</td>
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<tr>
<td>Sickle cell anemia</td>
<td>Sickle cell anemia affects millions of people throughout the world. It is particularly common among people whose ancestors come from sub-Saharan Africa; Spanish-speaking regions (South America, Cuba, Central America), Saudi Arabia, India, and Mediterranean countries, such as Turkey, Greece, and Italy. In the United States, it affects approximately 72,000 people, most of whose ancestors come from Africa. The disease occurs in approximately 1 in every 1,000 to 1,400 Hispanic American births. Approximately 2 million Americans, or 1 in 12 African Americans, carry the sickle cell trait.</td>
</tr>
</tbody>
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Clinical Alert

1. Correct test interpretation also requires knowledge of all medications the patient is taking.
2. Support the patient and his or her significant others in understanding and coping with positive or negative test outcomes.
3. Recognize that “panic values” may pose an immediate threat to the patient’s health status. Report these findings to the attending physician or other designated person immediately. Carefully document results and actions taken as soon as possible.
4. Nearly all tests have limitations. Some tests cannot predict future outcomes or events. For example, an ECG cannot predict a future myocardial infarction; it can merely tell what has already occurred. No test is absolute.
5. African American and Caucasian ethnic groups have similar cholesterol levels at birth. During childhood, African American people develop higher levels than do Caucasian people; however, African American adults have lower cholesterol levels than do Caucasian adults.

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4. Nearly all tests have limitations. Some tests cannot predict future outcomes or events. For example, an ECG cannot predict a future myocardial infarction; it can merely tell what has already occurred. No test is absolute.
5. Devastating physical, psychological, and social consequences can result from being misdiagnosed with a serious disease because of false-positive or false-negative test results. Major alterations in lifestyles and relationships without just cause can be a consequence of these clinical aberrations (eg, misdiagnosis of HIV or syphilis).
Monitoring for Complications

Observe for complications or other risks, and take appropriate measures to prevent or deal with them in a safe patient environment.

1. The most common complications after invasive procedures are bleeding, infection (frequently a later complication), respiratory difficulties, perforation of organs, and adverse effects of conscious sedation and local anesthesia. Watch for related signs and symptoms such as redness, swelling, skin irritation, pain or tenderness, dyspnea, abnormal breath sounds, cyanosis, decreased or increased pulse rate, blood pressure deviations (eg, hypertension, hypotension), laryngospasm, agitation or combative behavior, pallor, and complaints of dizziness. If adverse reactions or events occur, contact the physician immediately and initiate treatment as soon as possible.

2. Posttest assessments include evaluation of patient behaviors, complaints, activities, and compliance within the emotional, physical, psychosocial, and spiritual dimensions. Alterations in any of these domains may indicate a need for interventions appropriate to the dimension affected.

3. Older patients and children may require closer, more lengthy monitoring and observation. For example, invasive procedure sites should be observed and assessed for potential bleeding and circulatory problems in the immediate postprocedure phase and for infection as a later event (possibly several days later).

4. Patients who receive sedation, drugs, contrast media (eg, iodine, barium), or radioactive substances must be evaluated and treated according to established protocols (see Appendix C and Chapter 9 and Chapter 10).

5. Infection control measures with standard precautions and aseptic techniques must be observed.

Test Result Availability

Collaborate with other disciplines to ensure that test results are made available to the clinician, patient, and staff as soon as possible. Time-critical information is of limited value if it is delayed or not received. Even though computerized communication technologies contribute to faster information delivery, clinicians are often left waiting for crucial clinical data. Using facsimile (fax) machines, computers, and wireless networks properly can expedite the reporting of vital patient data to the health care provider so that treatment can begin without delay.

Clinical Alert

The issue of confidentiality demands that access to records and information should be on a strict need-to-know basis with secure and protected access available to select individuals.

Referral and Treatment

Referrals for further testing and beginning treatment are a part of the collaborative process. For example, the clinician refers patients with abnormal Pap smear results to the specialist for colposcopy, loop electrocautery excision procedure (LEEP), or cervical or endometrial biopsy. The clinician refers the patient for genetic counseling and dietary therapy for genetic disorders such as phenylketonuria (PKU) cholesterol in the newborn.

Follow-Up Care

Follow-up care should be consistent and should provide clearly understood discharge instructions. Emphasize the importance of and protocols for follow-up visits if these are ordered. Schedule ordered follow-up visits as appropriate. Follow established protocols for discharge to home after testing is completed. For complex procedures that are invasive or require sedation, be certain that a responsible individual escorts the patient home. Provide specific instructions regarding infection control, barium elimination, iodine sensitivity, and resuming pretest activities. Have the patient repeat this information back to the person providing the information to ensure that it has been understood. Plan time for listening, support, discussion, and problem solving according to the patient’s needs and requests. Follow-up by phone may be done after discharge if indicated.

Documentation, Record Keeping, and Reporting

Record information about all phases of the diagnostic testing process in the patient’s health care record. Accurately document diagnostic activities and procedures during the pretest, intratest, and posttest phases because of legal, budgetary, reimbursement, and diagnostic-related grouping (DRG) and common practice terminology (CPT) code implications and constraints.

The patient’s health care record is the only way to validate the need for diagnostic care, the quality and type of care given, and the patient’s response to the care and to ensure that current standards of medical and nursing care and diagnostic testing are being met. The medical record may also be the basis for reimbursement for diagnostic tests by Medicare (government), or private insurance programs. Accuracy, completeness, objectivity, and legibility are of utmost importance in the documentation process. Documentation for laboratory and diagnostic testing includes recording all pretest, intratest, and posttest care:

1. Document that the purpose, side effects, risks, and expected results and benefits, as well as alternative methods, have been explained to the patient, and note who gave the explanation. Include information about medications, IV conscious sedation, start and end times, and patient responses. Describe allergic or adverse reactions (see Appendix B). Record data regarding disposition of specimens as well as information about follow-up care and discharge instructions.

2. Document the patient’s reasons for refusing a test along with any other pertinent information about the situation and who was given this report.
3. Maintain records of laboratory and diagnostic test data. Frequently, these records are transferred onto compact record storage systems such as microfilm or computer disks. For example, when an individual tests positive for HIV, it is necessary to review donor records at blood donor centers to determine whether the individual ever donated blood. If the infected person donated blood, the recipients of those blood components must be contacted and informed of the situation. This process is called “look back.” Because many years may pass between donation and transfusion and the time the donor tests HIV positive, medical history records of blood donors must be stored indefinitely.

4. Indicate the time, day, month, and year of entries. This information can assume great importance in the office or clinic setting when charts become very lengthy. Enter appropriate assessment data and note the patient’s concerns and questions that help to define the nursing diagnosis and focus for care planning. Document specific teaching and preparation of the patient before the procedure. Avoid generalizations.

5. When an interpreter is present, document the name and relationship to the patient. Record that patient consent to give confidential test information through an interpreter was obtained before revealing the information. Record any deviations from basic witnessed consent policies (eg, illiteracy, non–English-speaking client, sedation immediately before the request for consent signature, consent per telephone); include nurse measures employed to obtain appropriate consent for the procedure.

6. Record that the preparation, side effects, expected results, and interfering factors have been explained. Document the information given and the patient’s response to that information. Keep a record of all printed and written instructions. Record medications, treatments, food and fluids, intake status, beginning and end of specimen collection and procedure times, outcomes, and the patient’s condition during all phases of diagnostic care. If the patient does not appear for testing, document this fact; include any follow-up discussion with the patient. Completely and clearly describe side effects, symptoms, adverse reactions, and complications along with follow-up care and instructions for posttest care and monitoring.

7. Record the patient’s refusal to undergo diagnostic tests. Note the reasons, using the patient’s own words if possible. Document significant noncompliant behaviors such as refusal or inability to fast or to restrict or increase fluid or food intake, incomplete timed specimens, inadequate or improperly self-collected specimens, and missed or canceled test appointments. Place copies of letters sent in the patient’s chart.

8. Reporting includes patient notification regarding test outcomes in a timely fashion and documentation that the patient or family has been notified regarding test results. Document follow-up patient education and counseling.

9. Report results to designated professionals. Report critical (“panic”) values immediately, and document to whom results were reported, orders received, and urgent treatments initiated.

10. Report all communicable diseases to appropriate agencies.

11. Report and document situations that are mandatory by state statute (eg, suspected elder abuse, child abuse as evidenced by x-rays).

Reporting infectious diseases and outbreaks to state and federal governments is part of record keeping. Chart 1.4 and Chart 1.5 are examples of one state’s (Maryland) required reporting. Check with your individual state or province for specific guidelines.
<table>
<thead>
<tr>
<th>Disease and Condition</th>
<th>Other Information</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acquired immunodeficiency syndrome (AIDS)</td>
<td>Legionellosis</td>
</tr>
<tr>
<td>Amebiasis</td>
<td>Leptospirosis</td>
</tr>
<tr>
<td>Animal bites *</td>
<td>Lyme disease</td>
</tr>
<tr>
<td>Anthrax *</td>
<td>Malaria</td>
</tr>
<tr>
<td>Botulism *</td>
<td>Measles (rubeola) *</td>
</tr>
<tr>
<td>Brucellosis</td>
<td>Meningitis (viral, bacterial, parasitic, and fungal)</td>
</tr>
<tr>
<td>Chancroid</td>
<td>Meningococcal disease</td>
</tr>
<tr>
<td>Cholera *</td>
<td>Mumps (infectious parotitis)</td>
</tr>
<tr>
<td>Diphtheria *</td>
<td>Mycobacteriosis other than tuberculosis and leprosy</td>
</tr>
<tr>
<td>Encephalitis</td>
<td>Pertussis *</td>
</tr>
<tr>
<td>Gonococcal infection</td>
<td>Pertussis vaccine adverse reactions</td>
</tr>
<tr>
<td>Haemophilus influenzae type b invasive disease *</td>
<td>Plague *</td>
</tr>
<tr>
<td>Hepatitis, viral (A, B, C, all other types and undetermined)</td>
<td>Plague *</td>
</tr>
<tr>
<td>Kawasaki syndrome</td>
<td>*Reportable immediately by telephone.</td>
</tr>
</tbody>
</table>

*Reportable immediately by telephone.

From State of Maryland Department of Health and Mental Hygiene. Epidemiology and Disease Control Program. (Reviewed: December 2002.)
<table>
<thead>
<tr>
<th>Diseases</th>
<th>Conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amebiasis</td>
<td>Microsporidiosis</td>
</tr>
<tr>
<td>Anthrax</td>
<td>Mumps</td>
</tr>
<tr>
<td>Bacteremia in newborns</td>
<td>Pertussis</td>
</tr>
<tr>
<td>Botulism</td>
<td>Plague</td>
</tr>
<tr>
<td>Brucellosis</td>
<td>Poliomyelitis</td>
</tr>
<tr>
<td>Campylobacter infection</td>
<td>Psittacosis</td>
</tr>
<tr>
<td>CD4+ count, if &lt;200/mm³</td>
<td>Q fever</td>
</tr>
<tr>
<td>Chlamydia infection</td>
<td>Rabies</td>
</tr>
<tr>
<td>Cholera</td>
<td>Ricin toxin</td>
</tr>
<tr>
<td>Coccidiodomycosis</td>
<td>Rocky Mountain spotted fever</td>
</tr>
<tr>
<td>Cryptosporidiosis</td>
<td>Rubella and congenital rubella syndrome</td>
</tr>
<tr>
<td>Cyclosporiasis</td>
<td>Salmonellosis (nontyphoid fever types)</td>
</tr>
<tr>
<td>Dengue fever</td>
<td>Shiga-like toxin production</td>
</tr>
<tr>
<td>Diphtheria</td>
<td>Shigellosis</td>
</tr>
<tr>
<td>Ehrlichiosis</td>
<td>Smallpox and other orthopox viruses</td>
</tr>
<tr>
<td>Encephalitis, infectious</td>
<td>Staphylococcal enterotoxin</td>
</tr>
<tr>
<td>E. coli O157:H7 infection</td>
<td>Streptococal invasive disease, group A</td>
</tr>
<tr>
<td>Giardiasis</td>
<td>Streptococal invasive disease, group B</td>
</tr>
<tr>
<td>Gonorrhea</td>
<td>Streptococcus pneumonia, invasive disease</td>
</tr>
<tr>
<td>Haemophilus influenza, invasive</td>
<td>Syphilis</td>
</tr>
<tr>
<td>Hansen's disease (leprosy)</td>
<td>Trichinosis</td>
</tr>
<tr>
<td>Hantavirus infection</td>
<td>Tularemia</td>
</tr>
<tr>
<td>Hepatitis, viral, types A, B, C</td>
<td>Typhoid fever</td>
</tr>
<tr>
<td>HIV infection</td>
<td>Varicella (chickenpox), fatal cases only</td>
</tr>
<tr>
<td>Isosporiasis</td>
<td>Vibriosis, noncholera</td>
</tr>
<tr>
<td>Legionellosis</td>
<td>Viral hemorrhagic fever (all types)</td>
</tr>
<tr>
<td>Leptospirosis</td>
<td>Yellow fever</td>
</tr>
<tr>
<td>Lyme disease</td>
<td>Yersiniosis</td>
</tr>
<tr>
<td>Malaria</td>
<td></td>
</tr>
<tr>
<td>Measles</td>
<td></td>
</tr>
<tr>
<td>Meningococcal invasive disease</td>
<td>2 invasive disease means a disease in which an organism is detected in a specimen taken from a normally sterile body site;</td>
</tr>
<tr>
<td>Meningitis, infectious</td>
<td></td>
</tr>
</tbody>
</table>

1 Reportable by unique patient identifying number;

2 invasive disease means a disease in which an organism is detected in a specimen taken from a normally sterile body site;
Guidelines for Disclosure

Follow agency guidelines for disclosure. Ethical standards may be a source of conflict and anxiety when the professional clinician is acting in the role of patient advocate. Recommended guidelines for telling a patient about test results can alleviate some of this frustration. Under normal circumstances, the patient has the right to be informed of test results. Although the clinician who orders the test is responsible for providing initial test result information, other designated individuals may need to facilitate and support the patient's right to know information about their health status.

In cases in which the patient brings family and significant others together to inform them about test results, communication becomes open and shared. This prevents the so-called conspiracy of silence, in which individuals in the scenario withhold information because they feel they are protecting the patient or family or because they do not know how to deal with the situation.

Patient Responses to Expected or Unexpected Outcomes

Develop crisis intervention skills to use when communicating with the patient who experiences difficulty dealing with abnormal test results or confirmation of disease or illness.

1. Encourage the patient to take as much control of the situation as possible.
2. Recognize that the different stages of behavioral responses to negative results may last several weeks or longer.
3. Monitor changes in patient affect, mood, behaviors, and motivation. Do not assume that persons who initially have a negative perception of their health (eg, denial of diabetes) will not be able to integrate better health behaviors into daily life once they accept the diagnosis.
4. Use the following strategies to lessen the impact of a threatening situation:
   a. Offer appropriate comfort measures.
   b. Allow patients to work through feelings of anxiety and depression. At the appropriate time, reassure them that these feelings and emotions are normal initially. Be more of a therapeutic listener than a talker.
   c. Assist the patient and family in making necessary lifestyle and self-concept adjustments through education, support groups, and other means. Emphasize that risk factors associated with certain diseases can be reduced through lifestyle changes. Be realistic.

It is better to introduce change slowly rather than trying to promote adjustments on a grand scale in a short period of time (Table 1.9).

<table>
<thead>
<tr>
<th>Immediate Response</th>
<th>Secondary Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute emotional turmoil, shock, disbelief about diagnosis, denial</td>
<td>Insomnia, anorexia, difficulty concentrating, depression, difficulty in performing work-related responsibilities and tasks</td>
</tr>
<tr>
<td>Anxiety will usually last several days until the person assimilates the information.</td>
<td>Depression may last several weeks as the person begins to incorporate the information and to participate realistically in a treatment plan and lifestyle adaptation.</td>
</tr>
</tbody>
</table>

Expected and Unexpected Outcomes

Evaluate outcomes using the following steps:

1. Learn the normal or reference values and expected outcomes of the test. The patient or his or her significant others should be able to describe the purpose of the test and the testing process and should properly perform expected activities associated with testing. Offer assistance if necessary. If test outcomes are abnormal, the patient should be encouraged to comply with repeat testing and to introduce appropriate lifestyle changes realistically. Deal with anxiety and fears in a timely manner. Refer the patient to appropriate counseling resources if indicated. Above all, do not dismiss the patient's feelings and concerns casually.
2. Compare normal values with abnormal results and apply these comparisons to the patient's situation. Sometimes, desired outcomes cannot be achieved. For example, the patient cannot, for various reasons, fully participate in the teaching/learning process or the actual testing itself. Recommendations for follow-up care and lifestyle changes may not be able to be followed. Verbal and nonverbal cues can sometimes provide reasons (eg, Alzheimer's disease, physical limitations) for this inability. In another instance, the patient might be noncompliant with pretest preparations and posttest activities. Denial of the situation is frequently a reason, although there are many other causes for noncompliance. Patients may refuse diagnostic testing because they feel the results may confirm their
Numerous and varied responses can be related to lack of appropriate problem-solving behaviors, inappropriate behaviors, fears or denial, concern about potential complications, inability to cope with or take control of the situation, depression or abnormal emotional patterns of response, and lack of support from significant others and family. Perceptions of having experienced uncaring acts can lead to frustration, despair, and hopelessness on the patient's part.

Adverse events (eg, perforation, anaphylaxis, death) and health hazards may occur as a result of diagnostic procedures or problems with a medical device or product (eg, reactions to latex gloves or other latex-containing medical devices). Health professionals are asked to monitor and voluntarily report faulty medical devices to the U.S. Food and Drug Administration (FDA) so that action can be taken to protect the public. Reporting does not necessarily constitute an admission that medical personnel or the product caused or contributed to the adverse event.

Prompt action is necessary when results are abnormally high or low and are indicative of a serious situation (eg, positive blood culture, abnormally elevated potassium level).

Modify, report, and collaborate with other clinicians when unexpected or abnormal values occur and when changes in medical care may be necessary as a result of test outcomes.

Examples of expected and unexpected test outcomes follow in Table 1.10.

<table>
<thead>
<tr>
<th>Expected Outcomes</th>
<th>Unexpected Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Some anticipated outcomes may not be achieved, possibly because of specific patient behaviors that interfere with care interventions (eg, patient does not appear for testing appointment, patient did not fast or withhold medication when directed before testing).</td>
<td></td>
</tr>
<tr>
<td>Inability to fully participate in teaching or learning process is evidenced by verbal and nonverbal cues. Patient cannot perform expected activities. Misinterpretation and misinformation of diagnostic process results in panic, avoidance behaviors, and refusal to have tests done.</td>
<td></td>
</tr>
<tr>
<td>Patient does not comply with test preparation guidelines and posttest recommended lifestyle changes; hides test results; and minimizes or exaggerates meaning of test outcomes.</td>
<td></td>
</tr>
<tr>
<td>Patient exhibits untoward signs and symptoms (eg, allergic response, shock, bleeding, nausea, vomiting, retention of barium).</td>
<td></td>
</tr>
<tr>
<td>Complications are not fully resolved, health state is compromised, and more extensive testing and care are needed.</td>
<td></td>
</tr>
</tbody>
</table>

**Table 1.10 Test Outcomes**

**IMPORTANCE OF COMMUNICATION**

At the heart of informed care is the ability to communicate effectively. Frequently, communication must take place within a compressed time frame because of time constraints. Thus, the importance of communicating effectively cannot be emphasized enough. Effective communication is the key to achieving desired outcomes, preventing misunderstanding and errors, and helping patients feel secure and connected to the diagnostic process. One must always keep in mind that the human person is an integration of body, mind, and spirit and that these three entities are intimately bound together to make each person unique. Skillful assessment of physical, emotional, psychosocial, and spiritual dimensions provides a sound database from which to plan communication and teaching/instruction strategies.

Individuals have different needs and changing capacities for learning as they progress from child to adult to older adult. It is important for the clinician to know the different developmental levels and stages and the ways in which clear communication can be achieved at any level.

For the pediatric patient, teaching tools might include tours of the diagnostic area, play therapy, films or videos, models of equipment that the child can touch or manipulate, and written materials and pictures appropriate to the child's developmental stage. Shorter attention spans and the unpredictable nature of children can make teaching a challenge in this population. Mentally retarded or mentally ill patients may need significant others close by who can guide communication between caretaker and patient. Gentle, simple, nurturing behaviors usually work well with children and developmentally challenged individuals.
Adolescents may be at the stage of developing their own unique identity as they move toward adulthood. Teaching may be more effective without parents present; however, it is important to include parents at some point. Drawings, illustrations, or videos are helpful. Because body image is very important at this stage, honest, supportive behaviors are necessary, especially if some alteration in physical appearance will be necessary (e.g., removal of jewelry, no makeup allowed).

The opportunity to participate actively and to ask questions is important for adults. They bring to the communication process their lifetime of perceptions and experiences. This can be a proverbial "double-edged sword." Listening well to verbal cues as well as paying attention to nonverbal messages cannot be overemphasized. For example, interacting with patients who have Alzheimer's disease can present special challenges. The presence of a significant other who has experience communicating with this patient can be the key to performing a successful procedure.

Provide an environment that is quiet, private, and free of distractions to promote dialogue and communication. Ask by what name or title the patient wishes to be addressed. Referring to a patient as a room number, a procedure, or a disease is demeaning and inexcusable; it reduces the patient to the level of an object rather than a person.

Nonverbal communication behaviors such as proper eye contact, firm handshake, sense of respect, and appropriate humor can reduce anxiety. Do not dismiss the power of touch, the sense of "making time" for the patient, and the use of appropriate and positive verbal cues. The greater part of communication (>70%) is perceived through body language. If words don't match body language and behaviors, patients will react to the body language they observe as their primary frame of reference. Negative communication by caregivers often is experienced by patients as an uncaring attitude and results in a sense of discouragement.

Every person engaged in the entire process of testing is a link in the ongoing communication continuum. This continuum is only as effective as the weakest link that joins all activities and all communication together.

CONCLUSION

As professionals, we need to remember that patients are people just like us. These individuals present with their perceptions, worries, and anxieties regarding the diagnostic process and what their illness means to them and their loved ones, what strategies they use for coping, what resources are available for their use, and what other knowledge they have about themselves. As clinicians and patient advocates, we must be willing to "take on the mind" of another—that is, to identify with the patient's point of view as much as possible and to show empathy. Once we reach that point, we can then begin to understand and communicate with each other at the deeper levels necessary for a therapeutic relationship to occur.

BIBLIOGRAPHY

Auxter SF: Identifying inappropriate laboratory testing strategies. What testing procedures should be eliminated or updated? Clinical Laboratory News, Vol. 27, No. 6, June 2001


Doak CC, Doak LG, Root J: Teaching Patients with Low Literacy Skills, 2nd ed. Philadelphia, Lippincott Williams & Wilkins, 1995


Fischbach FT: Quick Reference to Common Laboratory and Diagnostic Tests, 3rd ed. Philadelphia, Lippincott Williams & Wilkins, 2002

Fischbach FT: Documenting Care—Communication; Nursing Process; Documentation Standards. Philadelphia, FA Davis, 1991


Giger J, Davidhizar R: Transcultural Nursing. St. Louis, Mosby, 1999


Haeldorsdotter S: The effects of uncaring. Reflections, 4th Quarter, 28–30, 1999


OVERVIEW OF BASIC BLOOD HEMATOLOGY AND COAGULATION TESTS
Composition of Blood

Blood Tests

BLOOD SPECIMEN COLLECTION PROCEDURES
Capillary Puncture (Skin Puncture)
Venipuncture

NOTE
NOTE
Bone Marrow Aspiration

BASIC BLOOD TESTS

Complete Blood Count (CBC)

TESTS OF WHITE BLOOD CELLS
White Blood Cell Count (WBC; Leukocyte Count)
Differential White Blood Cell Count (Diff; Differential Leukocyte Count)

TESTS OF RED BLOOD CELLS
Red Blood Cell Count (RBC; Erythrocyte Count)
Hematocrit (Hct); Packed Cell Volume (PCV)
Hemoglobin (Hb)

NOTE
Reticulocyte Count
Sedimentation Rate (Sed Rate); Erythrocyte Sedimentation Rate (ESR)

TESTS FOR HEMOGLOBIN DISORDERS
Hemoglobin Electrophoresis

Fetal Hemoglobin (Hemoglobin F; Alkali-Resistant Hemoglobin)
Hemoglobin A2 (Hb A2)
Hemoglobin S (Sickle Cell Test; Sickledex)
Methemoglobin (Hemoglobin M)
Sulfhemoglobin

Other Blood Tests for Anemia
Vitamin B12 (VB12)
Folic Acid (Folate)
Erythropoietin (Ep)

TESTS FOR HEMOSTASIS AND COAGULATION

Iron (Fe), Total Iron-Binding Capacity (TIBC), and Transferrin Tests
Ferritin
Iron Stain (Stainable Iron in Bone Marrow; Prussian Blue Stain)

Clinical Alert
Fetal Hemoglobin (Hemoglobin F; Alkali-Resistant Hemoglobin)
Hemoglobin A2 (Hb A2)
Hemoglobin B (Sickle Cell Test; Sickledex)
Methemoglobin (Hemoglobin M)
Sulfhemoglobin
Carboxyhemoglobin; Carbon Monoxide (CO)
Myoglobin (Mb)
Haptoglobin (Hp)
Bot's Hemoglobin
Paroxysmal Nocturnal Hemoglobinuria (PNH) Test; Acid Hemolysis Test; Ham's Test

Other Blood Tests for Anemia
Vitamin B12 (VB12)
Folic Acid (Folate)
Erythropoietin (Ep)
Tests for Disseminated Intravascular Coagulation (DIC)

- Bleeding Time (Ivy Method; Template Bleeding Time)
- Platelet Count; Mean Platelet Volume (MPV)
- Platelet Aggregation
- Thrombin Time (TT); Thrombin Clotting Time (TCT)
- Partial Thromboplastin Time (PTT); Activated Partial Thromboplastin Time (APTT)

**NOTE**
- Activated Coagulation Time (ACT)
- Prothrombin Time (Pro Time; PT)
- Coagulant Factors (Factor Assay)
- Plasminogen (Plasmin; Fibrinolysin)
- Fibrinolysis (Euglobulin Lysis Time; Diluted Whole Blood Clot Lysis)
- Fibrin Split Products (FSPs); Fibrin Degradation Products (FDPs)
- D-Dimer
- Fibrinopeptide A (FPA)
- Prothrombin Fragment (F1 + 2)
- Fibrin Monomers (Protamine Sulfate Test; Fibrin Split Products)
- Fibrinogen
- Protein C (PC Antigen)

**NOTE**
- Protein S
- Antithrombin III (AT-III; Heparin Cofactor Activity)

**BIBLIOGRAPHY**

**OVERVIEW OF BASIC BLOOD HEMATOLOGY AND COAGULATION TESTS**

**Composition of Blood**

The average person circulates about 5 L of blood (1/13 of body weight), of which 3 L is plasma and 2 L is cells. Plasma fluid derives from the intestines and lymphatic systems and provides a vehicle for cell movement. The cells are produced primarily by bone marrow and account for blood "solids." Blood cells are classified as white cells (leukocytes), red cells (erythrocytes), and platelets (thrombocytes). White cells are further categorized as granulocytes, lymphocytes, monocytes, eosinophils, and basophils.

Before birth, hematopoiesis occurs in the liver. In midfetal life, the spleen and lymph nodes play a minor role in cell production. Shortly after birth, hematopoiesis in the liver ceases, and the bone marrow is the only site of production of erythrocytes, granulocytes, and platelets. B lymphocytes are produced in the marrow and in the secondary lymphoid organs; T lymphocytes are produced in the thymus.

**Blood Tests**

Tests in this chapter are basic screening tests that address disorders of hemoglobin (Hb) and cell production (hematopoiesis), synthesis, and function. Blood and bone marrow examinations constitute the major means of determining certain blood disorders (anemias, leukemia and porphyrias disorders, abnormal bleeding and clotting), inflammation, infection and inherited disorders of red blood cells, white blood cells, and platelets. Specimens are obtained through capillary skin punctures (finger, toe, heel), dried blood samples, arterial or venous sampling, or bone marrow aspiration. Specimens may be tested by automated or manual hematology instrumentation and evaluation.

**BLOOD SPECIMEN COLLECTION PROCEDURES**

Proper specimen collection presumes correct technique and accurate timing when necessary. Most hematology tests use liquid ethylenediaminetetraacetic acid (EDTA) as an anticoagulant. Tubes with anticoagulants should be gently but completely inverted end over end 7 to 10 times after collection. This action ensures complete mixing of anticoagulants with blood to prevent clot formation. Even slightly clotted blood invalidates the test, and the sample must be redrawn.

For plasma coagulator studies, such as prothrombin time (PT) and partial thromboplastin time (PTT), the tube must be allowed to fill to its capacity or an improper blood-to-anticoagulant ratio will invalidate coagulator results. Invert 7 to 10 times to prevent clotting.

**Capillary Puncture (Skin Puncture)**

Capillary blood is preferred for a peripheral blood smear and can also be used for other hematology studies.

**Procedure**

1. Observe standard precautions (see Appendix A). Check for latex allergy. If allergy is present, do not use latex-containing products (see Appendix B).
2. Obtain capillary blood from fingertips or earlobes (adults) or from the great toe or heel (infants).
3. Disinfect puncture site, dry the site, and puncture skin with sterile disposable lancet no deeper than 2 mm. If povidone-iodine is used, allow to dry thoroughly.
4. Wipe away the initial drop of blood. Collect subsequent drops in a microtube or prepare a smear directly from a
drop of blood.

**Clinical Alert**

1. Do not squeeze the site to obtain blood because this alters blood composition and invalidates test values.
2. Warming the extremity or placing it in a dependent position may facilitate specimen collection.

**Interventions**

**Pretest Patient Care**

1. Instruct patient about purpose and procedure of test.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Apply small dressing or adhesive strip to site.
2. Evaluate puncture site for bleeding or oozing.
3. Apply compression or pressure to the site if it continues to bleed.
4. Evaluate patient's medication history for anticoagulation or acetylsalicylic acid (ASA)-type drug ingestion.
5. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Clinical Alert**

In this method, a lancet is used, and the resulting droplets of blood are collected by blotting them with filter paper directly. Check the stability of equipment and integrity of supplies when doing a finger stick. If provided, check the humidity indicator patch on the filter paper card. If the humidity circle is pink, do not use this filter paper card. The humidity indicator must be blue to ensure specimen integrity.

After wiping the first drop of blood on the gauze pad, fill and saturate each of the circles in numerical order by blotting the blood droplet with the filter paper. Do not touch the patient's skin to the filter paper; only the blood droplet should come in contact with the filter paper. If an adult has a cold hand, run warm water over it for approximately 3 minutes. The best flow occurs when the arm is held downward, with the hand below heart level, making effective use of gravity. If there is a problem with proper blood flow, milk the finger with gentle pressure to stimulate blood flow or attempt a second finger stick; do not attempt more than two. When the blood circles penetrate through to the other side of the filter paper, the circles are fully saturated.

**Venipuncture**

**Venipuncture** allows procurement of larger quantities of blood for testing. Usually, the antecubital veins are the veins of choice because of ease of access. Blood values remain constant no matter which venipuncture site is selected, so long as it is venous and not arterial blood.

1. Observe standard precautions (see Appendix A). If latex allergy is suspected, use latex-free supplies and equipment (see Appendix B).
2. Position and tighten a tourniquet on the upper arm to produce venous congestion.
3. Ask the patient to close the fist in the designated arm. Select an accessible vein.
4. Cleanse the puncture site and dry it properly with sterile gauze. Povidone-iodine must dry thoroughly.
5. Puncture the vein according to accepted technique. Usually, for an adult, anything smaller than a 21-gauge needle might make blood withdrawal more difficult. A Vacutainer system syringe or butterfly system may be used.
6. Once the vein has been entered by the collecting needle, blood will fill the attached vacuum tubes automatically because of negative pressure within the collection tube.
7. Remove the tourniquet before removing the needle from the puncture site or bruising will occur.
8. Remove needle. Apply pressure and sterile dressing strip to site.
9. The preservative or anticoagulant added to the collection tube depends on the test ordered. In general, most hematology tests use EDTA anticoagulant. Even slightly clotted blood invalidates the test, and the sample must be redrawn.
10. Take action to prevent these venipuncture errors:
    a. Pretest errors
       1. Improper patient identification
       2. Failure to check patient compliance with dietary restrictions
       3. Failure to calm patient before blood collection
       4. Use of wrong equipment and supplies
       5. Inappropriate method of blood collection
    b. Procedure errors
       1. Failure to dry site completely after cleansing with alcohol
       2. Inserting needle with bevel side down
       3. Using too small a needle, causing hemolysis of specimen
       4. Venipuncture in unacceptable area (eg, above an intravenous [IV] line)
       5. Prolonged tourniquet application
       6. Wrong order of tube draw
       7. Failure to mix blood immediately that is collected in additive-containing tubes
       8. Pulling back on syringe plunger too forcefully
9. Failure to release tourniquet before needle withdrawal

c. Posttest errors
   1. Failure to apply pressure immediately to venipuncture site
   2. Vigorous shaking of anticoagulated blood specimens
   3. Forcing blood through a syringe needle into tube
   4. Mislabeling of tubes
   5. Failure to label specimens with infectious disease precautions as required
   6. Failure to put date, time, and initials on requisition
   7. Slow transport of specimens to laboratory

<table>
<thead>
<tr>
<th>NOTE</th>
</tr>
</thead>
<tbody>
<tr>
<td>A blood pressure cuff inflated to a point between systolic and diastolic pressure values can be used.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>NOTE</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Vacutainer system consists of vacuum tubes (Vacutainer tubes), a tube holder, and a disposable multisample collecting needle.</td>
</tr>
</tbody>
</table>

**Interventions**

**Pretest Patient Care**

1. Instruct patient regarding sampling procedure. Assess for circulation or bleeding problems and allergy to latex.
2. Reassure patient that mild discomfort may be felt when the needle is inserted.
3. Place the arm in a fully extended position with palmar surface facing upward (for antecubital access).
4. If withdrawal of the sample is difficult, warm the extremity with warm towels or blankets. Allow the extremity to remain in a dependent position for several minutes before venipuncture.

**Clinical Alert**

In patients with leukemia, agranulocytosis, or lowered resistance, finger-stick and earlobe punctures are more likely to cause infection and bleeding than venipunctures. Should a capillary sample be necessary, the cleansing agent should remain in contact with the skin for at least 5 to 10 minutes. Povidone-iodine is the cleansing agent of choice. It should be allowed to dry. It may then be wiped off with alcohol and the site dried with sterile gauze before puncture.

**Posttest Patient Aftercare**

1. If oozing or bleeding from the puncture site continues for an unusually long time, elevate the area and apply a pressure dressing. Observe the patient closely. Check for anticoagulant or ASA-type ingestion.
2. Be aware that the patient occasionally becomes dizzy, faint, or nauseated during the venipuncture. The phlebotomist must be constantly aware of the patient's condition. If a patient feels faint, immediately remove the tourniquet and terminate the procedure. If the patient is sitting, lower the head between the legs and instruct the patient to breathe deeply. A cool, wet towel may be applied to the forehead and back of the neck, and, if necessary, ammonia inhalant may be applied briefly. If the patient remains unconscious, notify a physician immediately.
3. Prevent hematomas by using proper technique (not sticking the needle through the vein), releasing the tourniquet before the needle is withdrawn, applying sufficient pressure over the puncture site, and maintaining an extended extremity until bleeding stops.

**Clinical Alert**

1. Never draw blood from the same extremity being used for IV medications, fluids, or transfusions. If no other site is available, make sure the venipuncture site is below the site. Avoid areas that are edematous, are paralyzed, are on the same side as a mastectomy, or have infections or skin conditions present. Venipuncture may cause infection, circulatory impairment, or retarded healing.

**Bone Marrow Aspiration**

Bone marrow is located within cancellous bone and long bone cavities. It consists of a pattern of vessels and nerves, differentiated and undifferentiated hematopoietic cells, reticuloendothelial cells, and fatty tissue. All of these are encased by endosteum, the membrane lining the bone marrow cavity. After proliferation and maturation have occurred in the marrow, blood cells gain entrance to the blood through or between the endothelial cells of the sinus wall.

A bone marrow specimen is obtained through aspiration or biopsy or needle biopsy aspiration. A bone marrow examination is important in the evaluation of a number of hematologic disorders and infectious diseases. The presence or suspicion of a blood disorder is not always an indication for bone marrow studies. A decision to employ this procedure is made on an individual basis.

Sometimes, the aspirate does not contain hematopoietic cells. This “dry tap” occurs when hematopoietic activity is so sparse that there are no cells to be withdrawn or when the marrow contains so many tightly packed cells that they cannot be suctioned out of the marrow. In such cases, a bone marrow biopsy would be advantageous. Before the bone marrow procedure is started, a peripheral blood smear should be obtained from the patient and a differential leukocyte count done.
Reference Values

Normal See Table 2.1 for normal values.

Table 2.1 Normal Values for Bone Marrow *

<table>
<thead>
<tr>
<th>Formed Cell Elements</th>
<th>Normal Mean (%)</th>
<th>Range (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Undifferentiated cells</td>
<td>0.0</td>
<td>0.0–1.0</td>
</tr>
<tr>
<td>Reticulum cells</td>
<td>0.4</td>
<td>0.0–1.3</td>
</tr>
<tr>
<td>Myeloblasts</td>
<td>2.0</td>
<td>0.3–5.0</td>
</tr>
<tr>
<td>Promyelocytes</td>
<td>5.0</td>
<td>1.0–8.0</td>
</tr>
<tr>
<td>Myelocytes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neutrophilic</td>
<td>12.0</td>
<td>5.0–19.0</td>
</tr>
<tr>
<td>Eosinophilic</td>
<td>1.5</td>
<td>0.5–3.0</td>
</tr>
<tr>
<td>Basophilic</td>
<td>0.3</td>
<td>0.0–0.5</td>
</tr>
<tr>
<td>Metamyelocytes</td>
<td>25.6</td>
<td>17.5–33.7</td>
</tr>
<tr>
<td>Neutrophilic</td>
<td>0.4</td>
<td>0.0–1.0</td>
</tr>
<tr>
<td>Eosinophilic</td>
<td>0.0</td>
<td>0.0–0.2</td>
</tr>
<tr>
<td>Segmented granulocytes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neutrophilic</td>
<td>20.0</td>
<td>11.6–30.0</td>
</tr>
<tr>
<td>Eosinophilic</td>
<td>2.0</td>
<td>0.5–4.0</td>
</tr>
<tr>
<td>Basophilic</td>
<td>0.2</td>
<td>0.0–3.0</td>
</tr>
<tr>
<td>Monocytes</td>
<td>2.0</td>
<td>0–3</td>
</tr>
<tr>
<td>Lymphocytes</td>
<td>10.0</td>
<td>8–20</td>
</tr>
<tr>
<td>Megakaryocytes</td>
<td>0.4</td>
<td>0.0–3.0</td>
</tr>
<tr>
<td>Plasma cells</td>
<td>0.9</td>
<td>0.0–2.0</td>
</tr>
<tr>
<td>Erythroid series</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pronormoblasts</td>
<td>0.5</td>
<td>0.2–4.2</td>
</tr>
<tr>
<td>Basophilic normoblasts</td>
<td>1.6</td>
<td>0.24–4.8</td>
</tr>
<tr>
<td>Polychromatic normoblasts</td>
<td>10.4</td>
<td>3.5–20.5</td>
</tr>
<tr>
<td>Orthochromatic normoblasts</td>
<td>6.4</td>
<td>3.0–25</td>
</tr>
<tr>
<td>Promegakoblasts</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Basophilic megaloblasts</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Polychromatic megaloblasts</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Orthochromatic megaloblasts</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Myeloid: erythroid ratio (ratio of WBC to nucleated RBC) 2.1–4:1 (Slightly higher in infants)

*These values are only for adults, and should be used as a guideline. (Each laboratory should establish its own reference range.)

Procedure

1. Follow standard precautions. Check for latex allergy; if allergy is present, do not use latex-containing products. Position the patient on the back or side according to site selected. The posterior iliac crest is the preferred site in all patients older than 12 to 18 months. Alternate sites include the anterior iliac crest, sternum, spinous vertebral processes T10 through L4, the ribs, and the tibia in children. The sternum is not generally used in children because the bone cavity is too shallow, the risk for mediastinal and cardiac perforation is too great, and the child may be uncooperative.

2. Shave, cleanse, and drape the site as for any minor surgical procedure.

3. Inject a local anesthetic (procaine or lidocaine). This may cause a burning sensation. At this time, a skin incision of 3 mm is often made.

4. Remember that the physician introduces a short, rigid, sharp-pointed needle with stylet through the periosteum into the marrow cavity.

5. Pass the needle-stylet combination through the incision, subcutaneous tissue, and bone cortex. The stylet is removed, and 1 to 3 mL of marrow fluid is aspirated. Alert the patient that when the stylet needle enters the marrow, he or she may experience a feeling of pressure. The patient may also feel moderate discomfort as aspiration is done, especially in the iliac crest. Use the Jamshidi needle for biopsy, although you can also use the Westerman-Jansen modification of the Vim-Silverman needle.

6. Rotate or “rock” the needle in several directions several times after adequate penetration of the base (3 cm) has been achieved. This “frees up” the specimen. Slowly withdraw the needle once this is done.

7. Push the biopsy specimen out backward from the needle. Use it to make touch preparations or immediately place in fixative. Make slide smears at the bedside.

8. Apply pressure to the puncture site until bleeding ceases. Dress the site.

9. Place specimens in biohazard bags, label properly, and route to the appropriate department.

Clinical Implications

1. A specific and diagnostic bone marrow picture provides clues to many diseases. The presence, absence, and ratio of cells are characteristic of the suspected disease.

2. Bone marrow examination may reveal the following abnormal cell patterns:
   a. Multiple myeloma, plasma cell myeloma, macroglobulinemia
   b. Chronic or acute leukemias
c. Anemia, including megaloblastic, macrocytic, and normocytic anemias
d. Toxic states that produce bone marrow depression or destruction
e. Neoplastic diseases in which the marrow is invaded by tumor cells (metastatic carcinoma, myeloproliferative and lymphoproliferative diseases); assists in diagnosis and staging
f. Agranulocytosis (a decrease in the production of white cells). This occurs when bone marrow activity is severely depressed, usually as a result of radiation therapy or chemotherapeutic drugs. Implications for the patient focus on the risk for death from overwhelming infection.
g. Platelet dysfunction
h. Some types of infectious diseases, especially histoplasmosis and tuberculosis
i. Deficiency of body iron stores, microcytic anemia
j. Lipid or glycogen storage disease

Interventions

Pretest Patient Care

1. Instruct the patient about the test procedure, purpose, benefits, and risks.
2. Ensure that a legal consent form is properly signed and witnessed. Bone marrow aspiration is usually contraindicated in the presence of hemophilia and other bleeding dyscrasias. However, risk versus benefit may dictate the choice made.
3. Reassure the patient that analgesics will be available if needed.
4. Be aware that bone marrow biopsies or aspirations can be uncomfortable. Squeezing a pillow may be helpful as a distraction technique.
5. Observe standard precautions.

Clinical Alert

1. Complications can include bleeding and sternal fractures. Osteomyelitis or injury to heart or great vessels is rare but can occur if the sternal site is used.
2. Manual and pressure dressings over the puncture site usually control excessive bleeding. Remove dressing in 24 hours. Redress site if necessary.
3. Fever, headache, unusual pain, or redness or pus at biopsy site may indicate infection (later event). Instruct patient to report unusual symptoms to physician immediately.

Posttest Patient Aftercare

1. Monitor vital signs until stable and assess site for excess drainage or bleeding.
2. Recommend bed rest for 30 minutes; then normal activities can be resumed.
3. Administer analgesics for sedatives as necessary. Soreness over the puncture site for 3 to 4 days after the procedure is normal. Continued pain may indicate fracture.
4. Interpret test outcomes and monitor appropriately.
5. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

BASIC BLOOD TESTS

Hemogram

A hemogram consists of a white blood cell count (WBC), red blood cell count (RBC), hemoglobin (Hb), hematocrit (Hct), red blood cell indices, and a platelet count. A complete blood count (CBC) consists of a hemogram plus a differential WBC.

Complete Blood Count (CBC)

The CBC is a basic screening test and is one of the most frequently ordered laboratory procedures. The findings in the CBC give valuable diagnostic information about the hematologic and other body systems, prognosis, response to treatment, and recovery. The CBC consists of a series of tests that determine number, variety, percentage, concentrations, and quality of blood cells:

1. White blood cell count (WBC): leukocytes fight infection
2. Differential white blood cell count (Diff): specific patterns of WBC
3. Red blood cell count (RBC): red blood cells carry O\(_2\) from lungs to blood tissues and CO\(_2\) from tissue to lungs
4. Hematocrit (Hct): measures RBC mass
5. Hemoglobin (Hb): main component of RBCs and transports O\(_2\) and CO\(_2\)
6. Red blood cell indices: calculated values of size and Hb content of RBCs; important in anemia evaluations
7. Mean corpuscular volume (MCV)
8. Mean corpuscular hemoglobin concentration (MCHC)
9. Mean corpuscular hemoglobin (MCH)
10. Stained red cell examination (film or peripheral blood smear)
11. Platelet count (often included in CBC): thrombocytes are necessary for clotting and control of bleeding
12. Red blood cell distribution width (RDW): indicates degree variability and abnormal cell size.
13. Mean platelet volume (MPV): index of platelet production

These tests are described in detail in the following pages.
Normal Values for Hemogram

<table>
<thead>
<tr>
<th>Age</th>
<th>WBC (× 10^3/mm^3)</th>
<th>RBC (× 10^6/mm^3)</th>
<th>Hb (g/dL)</th>
<th>Hct (%)</th>
<th>MCV (fL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Birth–2 wk</td>
<td>9.0–30.0</td>
<td>4.1–6.1</td>
<td>14.5–24.5</td>
<td>44–64</td>
<td>98–112</td>
</tr>
<tr>
<td>2–8 wk</td>
<td>5.0–21.0</td>
<td>4.0–6.0</td>
<td>12.5–20.5</td>
<td>39–59</td>
<td>98–112</td>
</tr>
<tr>
<td>2–6 mo</td>
<td>5.0–19.0</td>
<td>3.8–5.6</td>
<td>10.7–17.3</td>
<td>35–49</td>
<td>83–97</td>
</tr>
<tr>
<td>6 mo–1 y</td>
<td>5.0–19.0</td>
<td>3.8–5.2</td>
<td>9.9–14.5</td>
<td>29–43</td>
<td>73–87</td>
</tr>
<tr>
<td>1–6 y</td>
<td>5.0–19.0</td>
<td>3.9–5.3</td>
<td>9.5–14.1</td>
<td>30–40</td>
<td>70–84</td>
</tr>
<tr>
<td>6–16 y</td>
<td>4.8–10.8</td>
<td>4.0–5.2</td>
<td>10.3–14.9</td>
<td>32–42</td>
<td>73–87</td>
</tr>
<tr>
<td>16–18 y</td>
<td>4.8–10.8</td>
<td>4.2–5.4</td>
<td>11.1–15.7</td>
<td>34–44</td>
<td>75–89</td>
</tr>
<tr>
<td>&gt;18 y (males)</td>
<td>5.0–10.0</td>
<td>4.5–5.5</td>
<td>14.0–17.4</td>
<td>42–52</td>
<td>84–96</td>
</tr>
<tr>
<td>&gt;18 y (females)</td>
<td>5.0–10.0</td>
<td>4.0–5.0</td>
<td>12.0–16.0</td>
<td>36–48</td>
<td>84–96</td>
</tr>
</tbody>
</table>

Age | MCH (pg/cell) | MCHC (g/dL) | Platelets (× 10^3/mm^3) | RDW (%) | MPV (fL) |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Birth–2 wk</td>
<td>34–40</td>
<td>33–37</td>
<td>150–450</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>2–8 wk</td>
<td>30–36</td>
<td>32–36</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>2–6 mo</td>
<td>27–33</td>
<td>31–35</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>6 mo–1 y</td>
<td>24–30</td>
<td>32–36</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>1–6 y</td>
<td>23–29</td>
<td>31–35</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>6–16 y</td>
<td>24–30</td>
<td>32–36</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>16–18 y</td>
<td>25–31</td>
<td>32–36</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>&gt;18 y</td>
<td>28–34</td>
<td>32–36</td>
<td>140–400</td>
<td>11.5–14.5</td>
<td>7.4–10.4</td>
</tr>
</tbody>
</table>

Interventions

Pretest Patient Care for Hemogram, CBC, and Differential (Diff) Count (All Components)

1. Explain test procedure. Explain that slight discomfort may be felt when skin is punctured. Refer to venipuncture procedure for additional information.
2. Avoid stress if possible because altered physiologic status influences and changes normal hemogram values.
3. Select hemogram components ordered at regular intervals (eg, daily, every other day). These should be drawn consistently at the same time of day for reasons of accurate comparison; natural body rhythms cause fluctuations in laboratory values at certain times of the day.
4. Dehydration or overhydration can dramatically alter values; for example, large volumes of IV fluids can “dilute” the blood, and values will appear as lower counts. The presence of either of these states should be communicated to the laboratory.
5. Fasting is not necessary. However, fat-laden meals may alter some test results as a result of lipidemia.

Posttest Patient Aftercare for Hemogram, CBC, and Differential (Diff) Count (All Components)

1. Apply manual pressure and dressings to the puncture site on removal of the needle.
2. Monitor the puncture site for oozing or hematoma formation. Maintain pressure dressings on the site if necessary. Notify physician of unusual problems with bleeding.
3. Resume normal activities and diet.
4. Bruising at the puncture site is not uncommon. Signs of inflammation are unusual and should be reported if the inflamed area appears larger, if red streaks develop, or if drainage occurs.

Clinical Alert

NEVER apply a total circumferential dressing and wrap because this may compromise circulation and nerve function if constriction, from whatever cause, occurs.

TESTS OF WHITE BLOOD CELLS

White Blood Cell Count (WBC; Leukocyte Count)

White blood cells (or leukocytes) are divided into two main groups: granulocytes and agranulocytes. The granulocytes receive their name from the distinctive granules that are present in the cytoplasm of neutrophils, basophils, and eosinophils. However, each of these cells also contains a multilobed nucleus, which accounts for their also being called polymorphonuclear leukocytes. In laboratory terminology, they are often called “polys” or PMNs. The nongranulocytes, which consist of the lymphocytes and monocytes, do not contain distinctive granules and have nonlobular nuclei that are not necessarily spherical. The term mononuclear leukocytes is applied to these cells.

The endocrine system is an important regulator of the number of leukocytes in the blood. Hormones affect the production of leukocytes in the blood-forming organs, their storage and release from the tissue, and their disintegration. A local inflammatory process exerts a definite chemical effect on the mobilization of leukocytes. The life span of leukocytes varies from 13 to 20 days, after which the cells are destroyed in the lymphatic system; many are excreted from the body in fecal matter.

Leukocytes fight infection and defend the body by a process called phagocytosis, in which the leukocytes actually encapsulate foreign organisms and destroy them. Leukocytes also produce, transport, and distribute antibodies as part of...
the immune response to a foreign substance (antigen).

The WBC serves as a useful guide to the severity of the disease process. Specific patterns of leukocyte response can be expected in various types of diseases as determined by the differential count (percentages of the different types of leukocytes). Leukocyte and differential counts, by themselves, are of little value as aids to diagnosis unless the results are related to the clinical condition of the patient; only then is a correct and useful interpretation possible.

**Reference Values**

| Normal Black adults: | 3.2–10.0 × 10³ cells/mm³ or × 10⁹/L or 3200–10,000 cells/mm³ Adults: 4.5–10.5 × 10³ cells/mm³ or × 10⁹/L or 4500–10,500 cells/mm³ Children: 0–2 weeks: 9.0–30.0 × 10³ cells/mm³ or × 10⁹/L or 9000–30,000 cells/mm³ 2–8 weeks: 5.0–21.0 × 10³ cells/mm³ or × 10⁹/L or 5000–21,000 cells/mm³ 2 months–6 years: 5.0–19.0 × 10³ cells/mm³ or × 10⁹/L or 5000–19,000 cells/mm³ 6–18 years: 4.8–10.8 × 10³ cells/mm³ or × 10⁹/L or 4800–10,800 cells/mm³ |

**NOTE**

Different labs have slightly different reference values.

**Procedure**

1. Obtain a venous anticoagulated EDTA blood sample of 5 mL or a finger-stick sample. Place a specimen in a biohazard bag.
2. Record the time when specimen was obtained (eg, 7:00 a.m.).
3. Blood is processed either manually or automatically, using an electronic counting instrument such as the Coulter counter or Abbott Cell-Dyne.

**Clinical Implications**

1. **Leukocytosis:** WBC >11,000/mm³ or >11.0 × 10⁹/mm³ (or >11 × 10⁹/L)
   a. It is usually caused by an increase of only one type of leukocyte, and it is given the name of the type of cell that shows the main increase:
      1. Neutrophilic leukocytosis or neutrophilia
      2. Lymphocytic leukocytosis or lymphocytosis
      3. Monocytic leukocytosis or monocytosis
      4. Basophilic leukocytosis or basophilia
      5. Eosinophilic leukocytosis or eosinophilia
   b. An increase in circulating leukocytes is rarely caused by a proportional increase in leukocytes of all types. When this does occur, it is usually a result of hemoconcentration.
   c. In certain diseases (eg, measles, pertussis, sepsis), the increase of leukocytes is so great that the blood picture suggests leukemia. *Leukocytosis of a temporary nature* (leukemoid reaction) must be distinguished from leukemia. In leukemia, the leukocytosis is permanent and progressive.
   d. Leukocytosis occurs in acute infections, in which the degree of increase of leukocytes depends on severity of the infection, patient's resistance, patient's age, and marrow efficiency and reserve.
   e. Other causes of leukocytosis include the following:
      1. Leukemia, myeloproliferative disorders
      2. Trauma or tissue injury (eg, surgery)
      3. Malignant neoplasms, especially bronchogenic carcinoma
      4. Toxins, uremia, coma, eclampsia, thyroid storm
      5. Drugs, especially ether, chloroform, quinine, epinephrine (Adrenalin), colony-stimulating factors
      6. Acute hemolysis
      7. Hemorrhage (acute)
      8. After splenectomy
      9. Polycythemia vera
      10. Tissue necrosis
   f. Occasionally, leukocytosis is found when there is no evidence of clinical disease. Such findings suggest the presence of:
      1. Sunlight, ultraviolet irradiation
      2. Physiologic leukocytosis resulting from excitement, stress, exercise, pain, cold or heat, anesthesia
      3. Nausea, vomiting, seizures
   g. Steroid therapy modifies the leukocyte response.  
      1. When corticotropin (adrenocorticotropic hormone, or ACTH) is given to a healthy person, leukocytosis occurs.
      2. When ACTH is given to a patient with severe infection, the infection can spread rapidly without producing the expected leukocytosis; therefore, what would normally be an important sign is obscured.

2. **Leukopenia:** WBC <4000/mm³ or <4.0 × 10⁹/mm³ or <4.0 cells × 10⁹/L occurs during and following:
   a. Viral infections, some bacterial infections, overwhelming bacterial infections
   b. Hypersplenism
   c. Bone marrow depression caused by heavy-metal intoxication, ionizing radiation, drugs:
      1. Antimetabolites
      2. Barbiturates
      3. Benzene
      4. Antibiotics
      5. Antihistamines
      6. Anticonvulsives
      7. Antithyroid drugs
8. Arsenicals
9. Cancer chemotherapy (causes a decrease in leukocytes; leukocyte count is used as a link to disease)
10. Cardiovascular drugs
11. Diuretics
12. Analgesics and antiinflammatory drugs
d. Primary bone marrow disorders:
   1. Leukemia (aleukemic)
   2. Pernicious anemia
   3. Aplastic anemia
   4. Myelodysplastic syndromes
   5. Congenital disorders
   6. Kostmann’s syndrome
   7. Reticular agenesis
   8. Cartilage-hair hypoplasia
   9. Shwachman-Diamond syndrome
   10. Chédiak-Higashi syndrome
e. Immune-associated neutropenia
f. Marrow-occupying diseases (fungal infection, metastatic tumor)
g. Pernicious anemia

Clinical Alert

1. WBC <500/mm$^3$ or $<0.5 \times 10^9$/mm$^3$ (or $\times 10^9$/L) represents a panic value.
2. WBC >30,000/mm$^3$ or $>30.0 \times 10^9$/mm$^3$ (or $\times 10^9$/L) is a panic value.

Interfering Factors

1. Hourly rhythm: there is an early-morning low level and late-afternoon high peak.
2. Age: in newborns and infants, the count is high (10,000/mm$^3$ to 20,000/mm$^3$ or $10 \times 10^9$/L to $20 \times 10^9$/L); the count gradually decreases in children until the adult values are reached between 18 and 21 years of age.
3. Any stressful situation that leads to an increase in endogenous epinephrine production and a rapid rise in the leukocyte count

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Refer to standard pretest care for hemogram, CBC, and differential count on page 47. Also, see Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately. Refer to standard posttest care for hemogram, CBC, and differential count on page 47. Also, follow Chapter 1 guidelines for safe, effective, informed posttest care.
2. In prolonged severe granulocytopenia or pancytopenia, give no fresh fruits or vegetables because the kitchen, especially in a hospital, may be a source of food contamination. When the WBC is low, a person can get a bacterial, pseudomonal, or fungal infection from fresh fruits and vegetables. Use a minimal-bacteria or commercially sterile diet. All food must be served from a new or single-serving package. Consider a leukemia diet. See dietary department for restrictions (eg, cooked food only) and careful food preparation. Do not give intramuscular injections. Do not take rectal temperature, give suppositories, or give enemas. Do not use razor blades. Do not give aspirin or nonsteroidal anti-inflammatory drugs (NSAIDs), which cause abnormal platelet dysfunction. Watch carefully for signs or symptoms of infection. Without leukocytes to produce inflammation, serious infections can have very subtle findings. Often, patients have only a fever.

Differential White Blood Cell Count (Diff; Differential Leukocyte Count)

The total count of circulating white blood cells is differentiated according to the five types of leukocytes, each of which performs a specific function.

Function of Circulating WBCs According to Leukocyte Type

<table>
<thead>
<tr>
<th>Cell</th>
<th>These Cells Function to Combat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neutrophils</td>
<td>Pyogenic infections (bacterial)</td>
</tr>
<tr>
<td>Eosinophils</td>
<td>Allergic disorders and parasitic infestations</td>
</tr>
<tr>
<td>Basophils</td>
<td>Parasitic infections, some allergic disorders</td>
</tr>
<tr>
<td>Lymphocytes</td>
<td>Viral infections (measles, rubella, chickenpox, infectious mononucleosis)</td>
</tr>
<tr>
<td>Monocytes</td>
<td>Severe infections, by phagocytosis</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Differential for Leukocyte Count</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
</tr>
<tr>
<td>---------</td>
</tr>
</tbody>
</table>

Interfering Factors

1. Hourly rhythm: there is an early-morning low level and late-afternoon high peak.
2. Age: in newborns and infants, the count is high (10,000/mm$^3$ to 20,000/mm$^3$ or $10 \times 10^9$/L to $20 \times 10^9$/L); the count gradually decreases in children until the adult values are reached between 18 and 21 years of age.
3. Any stressful situation that leads to an increase in endogenous epinephrine production and a rapid rise in the leukocyte count

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Refer to standard pretest care for hemogram, CBC, and differential count on page 47. Also, see Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately. Refer to standard posttest care for hemogram, CBC, and differential count on page 47. Also, follow Chapter 1 guidelines for safe, effective, informed posttest care.
2. In prolonged severe granulocytopenia or pancytopenia, give no fresh fruits or vegetables because the kitchen, especially in a hospital, may be a source of food contamination. When the WBC is low, a person can get a bacterial, pseudomonal, or fungal infection from fresh fruits and vegetables. Use a minimal-bacteria or commercially sterile diet. All food must be served from a new or single-serving package. Consider a leukemia diet. See dietary department for restrictions (eg, cooked food only) and careful food preparation. Do not give intramuscular injections. Do not take rectal temperature, give suppositories, or give enemas. Do not use razor blades. Do not give aspirin or nonsteroidal anti-inflammatory drugs (NSAIDs), which cause abnormal platelet dysfunction. Watch carefully for signs or symptoms of infection. Without leukocytes to produce inflammation, serious infections can have very subtle findings. Often, patients have only a fever.

Differential White Blood Cell Count (Diff; Differential Leukocyte Count)

The total count of circulating white blood cells is differentiated according to the five types of leukocytes, each of which performs a specific function.
Birth–1 wk 10–18 32–62 0–2 0–1 26–36 0–6 —
1–2 wk 8–16 19–49 0–4 0–0 38–46 0–9 —
2–4 wk 7–15 14–34 0–3 0–0 43–53 0–9 —
4–8 wk 7–13 15–35 0–3 0–1 41–71 0–7 —
2–6 mo 5–11 15–35 0–3 0–1 42–72 0–6 —
6 mo–1 y 6–12 13–33 0–3 0–0 46–76 0–5 —
1–6 y 5–11 13–33 0–3 0–0 46–76 0–5 —
6–16 y 5–11 32–54 0–3 0–1 27–57 0–5 —
16–18 y 5–11 34–64 0–3 0–1 25–45 0–5 —
>18 y 3–6 50–62 0–3 0–1 25–40 3–7 0–1

Bands or stab cells, immature forms of neutrophils; Segs, segmented neutrophils; Polys, polymorphonuclear neutrophils; Eos, eosinophils; Basos, basophils; Lymphs, lymphocytes; Monos, monocytes; Metas, metamyelocytes.

The differential count is expressed as a percentage of the total number of leukocytes (WBC). The distribution (number and type) of cells and the degree of increase or decrease are diagnostically significant. The percentages indicate the relative number of each type of leukocyte in the blood. The absolute count of each type of leukocyte is obtained mathematically by multiplying its relative percentage by the total leukocyte count. The formula is:

\[
\text{Relative value (\%)} \times \text{WBC (cells/mm}^3) = \text{Absolute value (cells/mm}^3) 
\]

NOTE

This is now the preferred way of reporting.

The differential count alone has limited value; it must always be interpreted in relation to the WBC. If the percentage of one type of cell is increased, it can be inferred that cells of that type are relatively more numerous than normal, but it is not known whether this reflects an actual increase in the (absolute) number of cells that are relatively increased or an absolute decrease in cells of another type. On the other hand, if the relative (percentage) values of the differential count and the total WBC are both known, it is possible to calculate absolute values that are not subject to misinterpretation.

Historically, the differential count has been done manually, but the newer hematology instruments can now do an automated differential count. The count is based on different chemical components of each cell type. However, not all samples can be evaluated by automated methods. When a leukocyte count is extremely low or high, a manual count may have to be done. Extremely abnormal leukocytes, such as those in leukemia, also have to be counted by hand. The automated instrument has built-in quality control that senses abnormal cells and flags the differential. A microscopic count must then be done.

**Segmented Neutrophils (Polymorphonuclear Neutrophils, PMNs, Segs, Polys)**

Neutrophils, the most numerous and important type of leukocytes in the body’s reaction to inflammation, constitute a primary defense against microbial invasion through the process of phagocytosis. These cells can also cause some body tissue damage by their release of enzymes and endogenous pyrogens. In their immature stage of development, neutrophils are referred to as “stab” or “band” cells. The term band stems from the appearance of the nucleus, which has not yet assumed the lobed shape of the mature cell.

This test determines the presence of neutrophilia or neutropenia. Neutrophilia is an increase in the absolute number of neutrophils in response to invading organisms and tumor cells. Neutropenia occurs when too few neutrophils are produced in the marrow, too many are stored in the blood vessel margin, or too many have been called to action and used up.

**Reference Values**

**Normal** Absolute count: 3000–7000/mm$^3$ or 3–7 × 10$^9$/L

**NOTE**

All references are using this SI unit for reporting.

Black adults: 1.2–6.6 × 10$^9$/L Differential: 50% of total WBC 0%–3% of total PMNs are stab or band cells

**Procedure**

1. Obtain a 5-mL blood sample in EDTA coagulant and place it in biohazard bag.
2. Count as part of the differential.

**Clinical Implications**

1. Neutrophilia (increased absolute number and relative percentage of neutrophils) >8.0 × 10$^9$/L or 8000/mm$^3$, for African Americans: >7.0 × 10$^9$/L or 7000/mm$^3$
   a. Acute, localized, and general bacterial infections. Also, fungal and spirochetal and some parasitic and rickettsial infections.
   b. Inflammation (eg, vasculitis, rheumatoid arthritis, pancreatitis, gout), and tissue necrosis (myocardial infarction, burns, tumors).
c. Metabolic intoxications (eg, diabetes mellitus, uremia, hepatic necrosis)
d. Chemicals and drugs causing tissue destruction (eg, lead, mercury, digitalis, venoms)
e. Acute hemorrhage, hemolytic anemia, hemolytic transfusion reaction
f. Myeloproliferative disease (eg, myeloid leukemia, polycythemia vera, myelofibrosis)
g. Malignant neoplasms—carcinoma
h. Some viral infections (noted in early stages) and some parasitic infections

2. Ratio of segmented neutrophils to band neutrophils: normally 1%–3% of PMNs are band forms (immature neutrophils).
a. Degenerative shift to left: in some overwhelming infections, there is an increase in band (immature) forms with no leukocytosis (poor prognosis).
b. Regenerative shift to left: there is an increase in band (immature) forms with leukocytosis (good prognosis) in bacterial infections.
c. Shift to the right: decreased band (immature) cells with increased segmented neutrophils can occur in liver disease, megaloblastic anemia, hemolysis, drugs, cancer, and allergies.
d. Hypersegmentation of neutrophils with no band (immature) cells is found in megaloblastic anemias (eg, pernicious anemia) and chronic morphine addiction.

e. Neutropenia (decreased neutrophils)
a. <1800/mm$^3$ or <1.8 × 10$^9$/L
b. African Americans: <1000/mm$^3$ or <40% of differential count
c. Causes associated with decreased or ineffective production:
   1. Inherited stem cell disorders and genetic disorders or cellular development
   2. Acute overwhelming bacterial infections (poor prognosis) and septicemia
   3. Viral infections (eg, mononucleosis, hepatitis, influenza, measles)
   4. Some rickettsial and parasitical (protozoan) diseases (malaria)
   5. Drugs, chemicals, ionizing radiation, venoms
   6. Hematopoietic diseases (eg, aplastic anemia, megaloblastic anemias, iron-deficiency anemia, aleukemic leukemia, myeloproliferative diseases)
d. Causes associated with decreased survival:
   1. Infections mainly in persons with little or no marrow reserves, elderly people, and infants
   2. Collagen vascular diseases with antineutrophil antibodies (eg, systemic lupus erythematous [SLE] and Felty’s syndrome)
   3. Autoimmune and isoimmune causes
   4. Drug hypersensitivity (There is an extensive list of drugs that continues to grow. Women are more likely than men to have a drug sensitivity. Removal of offending drug results in return to normal.)
   5. Spleenic sequestration
e. Neutropenia in neonates (<5000/mm$^3$ or <5.0 × 10$^9$/L or <1000/mm$^3$ or <1.0 × 10$^9$/L after first week of life)
   1. Maternal neutropenia, maternal drug ingestion, maternal isoimmunization to fetal leukocytes (maternal immunoglobulin G [IgG] antibodies to fetal neutrophils)
   2. Inborn errors of metabolism (eg, maple syrup urine disease)
   3. Immune deficits—acquired
   4. Deficits and disorders of myeloid stem cell (eg, Kostmann’s agranulocytosis, benign chronic granulocytopenia of childhood)
   5. Congenital neutropenia
f. Pregnancy—progressive decrease until labor

g. Some viral infections (noted in early stages) and some parasitic infections

3. Neutropenia (decreased neutrophils)
a. <1800/mm$^3$ or <1.8 × 10$^9$/L
b. African Americans: <1000/mm$^3$ or <40% of differential count
c. Causes associated with decreased or ineffective production:
   1. Inherited stem cell disorders and genetic disorders or cellular development
   2. Acute overwhelming bacterial infections (poor prognosis) and septicemia
   3. Viral infections (eg, mononucleosis, hepatitis, influenza, measles)
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   6. Hematopoietic diseases (eg, aplastic anemia, megaloblastic anemias, iron-deficiency anemia, aleukemic leukemia, myeloproliferative diseases)
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   5. Congenital neutropenia
f. Pregnancy—progressive decrease until labor
g. Some viral infections (noted in early stages) and some parasitic infections

4. Other leukocyte abnormalities and corresponding diseases are listed in Table 2.2.

<table>
<thead>
<tr>
<th>Abnormality</th>
<th>Description</th>
<th>Associated Diseases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toxic granulation</td>
<td>Coarse, black or purple, cytoplasmic granules</td>
<td>Infections or inflammatory diseases; acute reactive states</td>
</tr>
<tr>
<td>Döhle bodies</td>
<td>Small (1–2 µm), blue, cytoplasmic inclusions in neutrophils</td>
<td>Infections or inflammatory diseases, burns</td>
</tr>
<tr>
<td>Pelger-Huët anomalies</td>
<td>Neutrophil with bilobed nucleus or no segmentation of nucleus; chromatin is coarse, and cytoplasm is pink with normal granulation</td>
<td>Hereditary (congenital), myelogenous leukemia</td>
</tr>
<tr>
<td>May-Hegglin anomaly</td>
<td>Basophilic, cytoplasmic inclusions of leukocytes; similar to Döhle bodies</td>
<td>May-Hegglin syndrome (hereditary), includes thrombocytopenia and giant platelets</td>
</tr>
<tr>
<td>Alder-Reilly anomaly</td>
<td>Prominent azurophilic granulation in leukocytes; similar to toxic granulation; granulation is seen better with Giemsa stain</td>
<td>Hereditary, mucopolysaccharidosis</td>
</tr>
<tr>
<td>Chédiak-Higashi anomaly</td>
<td>Gray-green, large cytoplasmic inclusions that are fused giant lysosomes (phospholipids)</td>
<td>Chédiak-Higashi syndrome; few cases of acute myeloid leukemia</td>
</tr>
<tr>
<td>LE (lupus erythematous) cells</td>
<td>Neutrophilic leukocyte with a homogenous red-purple inclusion that distends the cell’s cytoplasm</td>
<td>Lupus erythematous and other collagen diseases, chronic hepatitis, drug reactions, serum sickness (not naturally occurring in the body—must be induced to form by mechanical trauma in vitro)</td>
</tr>
<tr>
<td>Tart cell</td>
<td>Neutrophilic leukocyte with a phagocytized nucleus of a granulocyte that retains some nuclear structure</td>
<td>Drug reactions (eg, penicillin, procainamide) or actual phagocytosis</td>
</tr>
</tbody>
</table>
Myeloid “shift to left”

Presence of bands, myelocytes, metamyelocytes, or promyelocytes

Infections, intoxications, tissue necrosis, myeloproliferative syndrome, leukemia (chronic myelocytic), leukemoid reaction, pernicious anemia, hyposplenism

Hypersegmented neutrophil

Mature neutrophil with more than five distinct lobes

Megaloblastic anemia, hereditary constitutional hypersegmentation of neutrophils; long-term chronic infection

Leukemic cells (eg, lymphoblasts, myeloblasts, myelomonoblasts, promyelocytes)

Presence of lymphoblasts, myeloblasts, monoblasts, myelomonoblasts, promyelocytes (none normally present in peripheral blood)

Leukemia (acute or chronic), leukemoid reaction, severe infectious or inflammatory diseases, myeloproliferative syndrome, intoxications, malignancies, recovery from bone marrow suppression

Auer bodies

Rod-like, 1–6 µm long, red-purple, refractile inclusions in neutrophils

Acute myelocytic leukemia or myelomonocytic leukemia

Smudge cell

Disintegrating nucleus of a ruptured leukocyte

Increased numbers in leukemic blood, particularly in acute lymphocytic leukemia or chronic lymphocytic leukemia when WBC count is greater than 10,000/mm³ or >10 × 10⁹/L

NOTE

An ethnic difference exists only in neutrophils.

Interfering Factors

1. Physiologic conditions such as stress, excitement, fear, vomiting, electric shock, anger, joy, and exercise temporarily cause increased neutrophils. Crying babies have neutrophilia.
3. Steroid administration: neutrophilia peaks in 4 to 6 hours and returns to normal by 24 hours (in severe infection, expected neutrophilia does not occur).
4. Exposure to extreme heat or cold.
5. Age
   a. Children respond to infection with a greater degree of neutrophilic leukocytosis than adults do.
   b. Some elderly patients respond weakly or not at all, even when infection is severe.
6. Resistance
   a. People of any age who are weak and debilitated may fail to respond with a significant neutrophilia.
   b. When an infection becomes overwhelming, the patient's resistance is exhausted and, as death approaches, the number of neutrophils decreases greatly.
7. Myelosuppressive chemotherapy
8. Many drugs cause increases or decreases in neutrophils.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Refer to standard pretest care for hemogram, CBC, and differential count on page 47. Also, see Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately for neutrophilia or neutropenia.
2. Refer to standard posttest care for hemogram, CBC, and differential count on page 47. Also, follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

Agranulocytosis (marked neutropenia and leukopenia) is extremely dangerous and is often fatal because the body is unprotected against invading agents. Patients with agranulocytosis must be protected from infection by means of reverse isolation techniques with strictest emphasis on handwashing technique.

Eosinophils

Eosinophils, capable of phagocytosis, ingest antigen-antibody complexes and become active in the later stages of inflammation. Eosinophils respond to allergic and parasitic diseases. Eosinophilic granules contain histamine (one third of all the histamine in the body).

This test is used to diagnose allergic infections, assess severity of infestations with worms and other large parasites, and monitor response to treatment.

Reference Values

Normal Absolute count: 0–0.7 × 10⁹/L Differential: 0%–3% of total WBC
Procedure

1. Obtain a 5-mL blood sample in EDTA anticoagulant. Place it in a biohazard bag.
2. Note the time the blood sample is obtained (eg, 3:00 p.m.).
3. Perform a total WBC, make a blood smear, count 100 cells, and report the percentage of eosinophils.
4. Be aware that an absolute eosinophil count is also available. It is done with a special eosinophil stain and manual counting on a hemacytometer. It must be done within 4 hours after collection or, if refrigerated, within 24 hours.

Clinical Implications

1. **Eosinophilia** (increased circulating eosinophils) >5% or >500 cells/mm \(^3\) or >0.5 \(\times\) 10 \(^9\)/L occurs in:
   a. Allergies, hay fever, asthma
   b. Parasitic disease and trichinosis tapeworm, especially with tissue invasion
   c. Some endocrine disorders, Addison's disease, hypopituitarism
   d. Hodgkin's disease and myeloproliferative disorders, chronic myeloid leukemia, polycythemia vera
   e. Chronic skin diseases (eg, pemphigus, eczema, dermatitis herpetiformis)
   f. Systemic eosinophilia associated with pulmonary infiltrates (PIE)
   g. Some infections (scarlet fever, chorea), convalescent stage of other infections
   h. Familial eosinophilia (rare), hypereosinophilic syndrome (HES)
   i. Polyarteritis nodosa, collagen vascular diseases (eg, SLE), connective tissue disorders
   j. Eosinophilic gastrointestinal diseases (eg, ulcerative colitis, Crohn's disease)
   k. Immunodeficiency disorders (Wiskott-Aldrich syndrome, immunoglobulin A deficiency)
   l. Aspirin sensitivity, allergic drug reactions
   m. Löeffler's syndrome (related to *Ascaris* species infestation), tropical eosinophilia (related to filariasis)
   n. Poisons (eg, black widow spider, phosphorus)
   o. Hypereosinophilic syndrome (>1.5 \(\times\) 10 \(^9\)/L), persistent extreme eosinophilia with eosinophilic infiltration of tissues causing tissue damage and organ dysfunction
      1. Eosinophilic leukemia
      2. Trichinosis invasion
      3. Dermatitis herpetiformis
      4. Idiopathic

2. **Eosinopenia** (decreased circulating eosinophils) is usually caused by an increased adrenal steroid production that accompanies most conditions of bodily stress and is associated with:
   a. Cushing's syndrome (acute adrenal failure): <50/mm \(^3\)
   b. Use of certain drugs such as ACTH, epinephrine, thyroxine, prostaglandins
   c. Acute bacterial infections with a marked shift to the left (increase in immature leukocytes)

3. **Eosinophilic myelocytes** are counted separately because they have a greater significance, being found only in leukemia or leukemoid blood pictures.

Interfering Factors

1. Daily rhythm: normal eosinophil count is lowest in the morning, then rises from noon until after midnight. For this reason, serial eosinophil counts should be repeated at the same time each day.
2. Stressful situations, such as burns, postoperative states, electroshock, and labor, cause a decreased count.
3. After administration of corticosteroids, eosinophils disappear.
4. See Appendix J for drugs that affect test outcomes.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Refer to standard patient care for hemogram, CBC, and differential count on page 47. Also, see Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately.
2. Use special precautions if patient is receiving steroid therapy, epinephrine, thyroxine, or prostaglandins.
   Eosinophilia can be masked by steroid use.
3. Refer to standard posttest care for hemogram, CBC, and differential count on page 47. Also, follow Chapter 1 guidelines for safe, effective, informed posttest care.

Basophils

Basophils, which constitute a small percentage of the total leukocyte count, are considered phagocytic. The basophilic granules contain heparin, histamines, and serotonin. Tissue basophils are called mast cells and are similar to blood basophils. Normally, mast cells are not found in peripheral blood and are rarely seen in healthy bone marrow.

Basophil counts are used to study chronic inflammation. There is a positive correlation between high basophil counts and high concentrations of blood histamines, although this correlation does not imply cause and effect. It is extremely difficult to diagnose basopenia because a 1000–10,000 count differential would have to be done to get an absolute count.

Reference Values

**Normal** Absolute count: 15–50/mm \(^3\) or 0.02–0.05 \(\times\) 10 \(^9\)/L Differential: 0%–1.0% of total WBC
Procedure

1. Obtain a 5-mL blood sample in EDTA and count as part of the differential.
2. Place the sample in a biohazard bag.

Clinical Implications

1. **Basophilia** (increased count) >50/mm³ or >0.05 × 10⁹/L is commonly associated with the following:
   a. Granulocytic (myelocytic) leukemia
   b. Acute basophilic leukemia
   c. Myeloid metaplasia, myeloproliferative disorders
   d. Hodgkin's disease
2. It is less commonly associated with the following:
   a. Inflammation, allergy, or sinusitis
   b. Polycythemia vera
   c. Chronic hemolytic anemia
   d. After splenectomy
   e. After ionizing radiation
   f. Hyperthyroidism
   g. Infections, including tuberculosis, smallpox, chickenpox, influenza
   h. Foreign protein injection
3. **Basopenia** (decreased count) <20/mm³ or <0.02 × 10⁹/L is associated with the following:
   a. Acute phase of infection
   b. Hyperthyroidism
   c. Stress reactions (eg, pregnancy, myocardial infarction)
   d. After prolonged steroid therapy, chemotherapy, radiation
   e. Hereditary absence of basophils
   f. Acute rheumatic fever in children
4. **Presence of numbers of tissue mast cells** (tissue basophils) is associated with:
   a. Rheumatoid arthritis
   b. Urticaria, asthma
   c. Anaphylactic shock
   d. Hypoadrenalism
   e. Lymphoma
   f. Macroglobulinemia
   g. Mast cell leukemia
   h. Lymphoma invading bone marrow
   i. Urticaria pigmentosa
   j. Asthma
   k. Chronic liver or renal disease
   l. Osteoporosis
   m. Systemic mastocytosis

Interfering Factors

See Appendix J for drugs that affect test outcomes.

Interventions

**Pretest Patient Care**

1. Explain test purpose and procedure
2. Refer to standard patient care for hemogram, CBC, and differential count on page 47. Also, see Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and monitor appropriately.
2. Use special precautions if patient is receiving steroid therapy, epinephrine, thyroxine, or prostaglandins.
   Eosinophilia can be masked by steroid use.
3. Refer to standard posttest care for hemogram, CBC, and differential count on page 47. Also, follow Chapter 1 guidelines for safe, effective, informed posttest care.

Monocytes (Monomorphonuclear Monocytes)

These agranulocytes, the largest cells of normal blood, are the body's second line of defense against infection. Histiocytes, which are large macrophagic phagocytes, are classified as monocytes in a differential leukocyte count. Histiocytes and monocytes are capable of reversible transformation from one to the other.

These phagocytic cells of varying size and mobility remove injured and dead cells, microorganisms, and insoluble particles from the circulating blood. Monocytes escaping from the upper and lower respiratory tracts and the gastrointestinal and genitourinary organs perform a scavenger function, clearing the body of debris. These phagocytic cells produce the antiviral agent called interferon.

This test counts monocytes, which circulate in certain specific conditions such as tuberculosis, subacute bacterial endocarditis, and the recovery phase of acute infections.

Reference Values
**Normal** Absolute count: 100–500/mm³ or 0.1–0.5 × 10⁹/L Differential: 3%–7% of total WBC or 0.03–0.07 of total WBC

**Procedure**

1. Obtain a 5-mL blood sample in EDTA and count as part of the differential.
2. Observe standard precautions.

**Clinical Implications**

1. In monocytosis: a monocyte increase of >500 cells/mm³ or >0.5 × 10⁹/L or >10%. The most common causes are bacterial infections, tuberculosis, subacute bacterial endocarditis, and syphilis.
2. Other causes of monocytosis:
   a. Monocytic leukemia and myeloproliferative disorders
   b. Carcinoma of stomach, breast, or ovary
   c. Hodgkin’s disease and other lymphomas
   d. Recovery state of neutropenia (favorable sign)
   e. Lipid storage diseases (e.g., Gaucher's disease)
   f. Some parasitic mycotic and rickettsial diseases
   g. Surgical trauma
   h. Chronic ulcerative colitis, enteritis, and sprue
   i. Collagen diseases and sarcoidosis
   j. Tetrachlorethane poisoning
3. Phagocytic monocytes (macrophages) may be found in small numbers in the blood in many conditions:
   a. Severe infections (sepsis)
   b. Lupus erythematosus
   c. Hemolytic anemias
4. Decreased monocyte count (<100 cells/mm³ or <0.1 × 10⁹/L) is not usually identified with specific diseases:
   a. Prednisone treatment
   b. Hair cell leukemia
   c. Overwhelming infection that also causes neutropenia
   d. Human immunodeficiency virus (HIV) infection
   e. Aplastic anemia (bone marrow injury)

**Interfering Factors** See Appendix J for drugs that affect test outcomes.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. Refer to standard pretest care for hemogram, CBC, and differential count on page 47. Also, see Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and monitor appropriately for leukemia and infection.
2. Refer to standard posttest care for hemogram, CBC, and differential count on page 47. Also, follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Lymphocytes (Monomorphonuclear Lymphocytes); CD4, CD8 Count; Plasma Cells**

Lymphocytes are small, mononuclear cells without specific granules. These agranulocytes are motile cells that migrate to areas of inflammation in both early and late stages of the process. These cells are the source of serum immunoglobulins and of cellular immune response and play an important role in immunologic reactions. All lymphocytes are manufactured in the bone marrow. B lymphocytes mature in the bone marrow, and T lymphocytes mature in the thymus gland. B cells control the antigen-antibody response that is specific to the offending antigen and is said to have “memory.” The T cells, the master immune cells, include CD4⁺ helper T cells, killer cells, cytotoxic cells, and CD8⁺ suppressor T cells.

Plasma cells (fully differentiated B cells) are similar in appearance to lymphocytes. They have abundant blue cytoplasm and an eccentric, round nucleus. Plasma cells are not normally present in blood.

This test measures the number of lymphocytes in the peripheral blood. Lymphocytosis is present in various diseases and is especially prominent in viral disorders. Lymphocytes and their derivatives, the plasma cells, operate in the immune defenses of the body.

**Reference Values**

- **Normal** Lymphocytes: 25%–40% of total leukocyte count (relative value) or 1500–4000 cells/mm³ or 1.5–4.0 × 10⁹/L
- Plasma cells: 0% or none CD4 count: total WBC × lymphocytes (%) × lymphocytes (%) stained with CD4 CD4/CD8 ratio: >1.0

**Procedure**

1. Obtain 5 mL of EDTA-anticoagulated blood. Place the specimen in a biohazard bag.
2. Count lymphocytes as part of the differential count.
1. **Lymphocytosis:** >4000/mm$^3$ or >4.0 × 10$^9$/L in adults; >7200/mm$^3$ or >7.2 × 10$^9$ in children; and >9000/mm$^3$ or >9.0 × 10$^9$/L in infants occurs in:
   a. Lymphatic leukemia (acute and chronic) lymphoma
   b. Infectious lymphocytosis (occurs mainly in children)
   c. Infectious mononucleosis:
      1. Caused by Epstein-Barr virus
      2. Most common in adolescents and young adults
      3. Characterized by atypical lymphocytes (Downey cells) that are large, deeply indented, with deep blue (basophilic) cytoplasm
      4. Differential diagnosis—positive heterophil test
d. Other viral diseases:
   1. Viral infections of the upper respiratory tract (pneumonia)
   2. Cytomegalovirus
   3. Measles, mumps, chickenpox
   4. Acute HIV infection
   5. Infectious hepatitis (acute viral hepatitis)
   6. Toxoplasmosis
e. Some bacterial diseases such as tuberculosis, brucellosis (undulant fever), and pertussis
f. Crohn's disease, ulcerative colitis (rare)
g. Serum sickness, drug hypersensitivity
h. Hypoadrenalism, Addison's disease
i. Thyrotoxicosis (relative lymphocytosis)
j. Neutropenia with relative lymphocytosis
d. Other viral diseases:
   1. Chemotherapy, radiation treatment (immunosuppressive medications)
   2. After administration of ACTH or cortisone (steroids); with ACTH-producing pituitary tumors
   c. Increased loss via gastrointestinal tract owing to obstruction of lymphatic drainage (eg, tumor, Whipple's disease, intestinal lymphectasia)
d. Aplastic anemia
e. Hodgkin's disease and other malignancies
f. Inherited immune disorders, acquired immunodeficiency syndrome (AIDS), and AIDS-immune dysfunction
g. Advanced tuberculosis ("miliary" tuberculosis), renal failure, SLE
h. Severe debilitating illness of any kind
i. Congestive heart failure
2. **Lymphopenia:** <1000 cells/mm$^3$ or <1.0 × 10$^9$/L in adults; <2500 cells/mm$^3$ or <2.5 × 10$^9$/L in children occurs in:
   a. Chemotherapy, radiation treatment (immunosuppressive medications)
   b. After administration of ACTH or cortisone (steroids); with ACTH-producing pituitary tumors
   c. Increased loss via gastrointestinal tract owing to obstruction of lymphatic drainage (eg, tumor, Whipple's disease, intestinal lymphectasia)
d. Aplastic anemia
e. Hodgkin's disease and other malignancies
f. Inherited immune disorders, acquired immunodeficiency syndrome (AIDS), and AIDS-immune dysfunction
g. Advanced tuberculosis ("miliary" tuberculosis), renal failure, SLE
h. Severe debilitating illness of any kind
i. Congestive heart failure
3. **CD4 count:** the number of CD4$^+$ lymphocytes is equal to the absolute number of lymphocytes (total WBC × differential [%] of lymphocytes) times the percentage of lymphocytes staining positively for CD4. A severely depressed CD4 count is the single best indicator of imminent opportunistic infection.
   a. Decreased CD4 lymphocytes
      1. Immune dysfunction, especially AIDS
      2. Acute minor viral infections
   b. Increased CD4 lymphocytes
      1. Therapeutic effect of drugs
      2. Diurnal variation: peak evening values may be two times morning values.
4. **Plasma cells** (not normally present in blood) are increased in:
   a. Plasma cell leukemia
   b. Multiple myeloma
c. Hodgkin's disease
d. Chronic lymphatic leukemia
e. Cancer of liver, breast, prostate
f. Cirrhosis
g. Rheumatoid arthritis, SLE
h. Serum reaction
i. Some bacterial, viral, and parasitic infections

**Interfering Factors**
1. Physiologic pediatric lymphocytosis is a condition in newborns that includes an elevated WBC and abnormal-appearing lymphocytes that can be mistaken for malignant cells.
2. Exercise, emotional stress, and menstruation can cause an increase in lymphocytes.
3. African Americans normally have a relative (not absolute) increase in lymphocytes.
4. See Appendix J for drugs that affect outcomes.

**Abnormal Lymphocytes**

<table>
<thead>
<tr>
<th>Abnormality</th>
<th>Description</th>
<th>Associated Diseases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atypical lymphocytes</td>
<td>Lymphocytes, some with vacuolated cytoplasm, irregularly shaped nucleus, increased numbers of cytoplasmic azurophilic granules, and peripheral basophilia; or some with more abundant basophilic cytoplasm, grossly indented cytoplasm</td>
<td>Infectious mononucleosis, viral hepatitis, other viral infections, tuberculosis, drug sensitivity, posttransfusion syndrome</td>
</tr>
<tr>
<td>Reactive lymphocytes</td>
<td>Downey cells</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Turk cells</td>
<td></td>
</tr>
</tbody>
</table>
Explain test purpose and procedure.
Patients with an absolute helper T-lymphocyte count <200/mm³
Decreased
Refer to standard
Standard immunosuppressive drug therapy usually
Explain purpose and specimen collection procedure. A recent viral cold can cause a decrease in total T cells, as
can medications such as corticosteroids. Nicotine and strenuous exercise have also been shown to decrease
lymphocyte counts.

Clinical Alert
A decreased lymphocyte count <500/mm³ (<0.5 × 10⁹/L) means that a patient is dangerously susceptible to infection, especially viral infections. Institute measures to protect patient from infection.

Posttest Patient Aftercare
1. Interpret test outcomes and monitor appropriately for lymphocytosis or lymphopenia.
2. Refer to standard posttest care for hemogram, CBC, and differential count on page 47. Also, follow Chapter 1
   guidelines for safe, effective, informed posttest care.

Lymphocyte Immunophenotyping (T and B Cells)
Lymphocytes are divided into two categories, T and B cells, according to their primary function within the immune system. In the body, T and B cells work together to help provide protection against infections, oncogenic agents, and foreign tissue, and they play a vital role in regulating self-destruction or autoimmunity.

Most circulating lymphocytes are T cells with a life span of months to years. The life span of B cells is measured in days. B cells (antibody) are considered “bursa or bone marrow dependent” and are responsible for humoral immunity (in which antibodies are present in the serum). T cells (cellular) are thymus derived and are responsible for cellular immunity. T cells are further divided into helper T (CD3⁺, CD4⁺) cells and suppressor T (CD3⁺, CD8⁺) cells.

Evaluation of lymphocytes in the clinical laboratory is performed by quantitation of the lymphocytes and their subpopulations and by assessment of their function activity. These laboratory analyses have become an essential component of the clinical assessment of two major disease states: lymphoproliferative states (eg, leukemia, lymphoma), in which characterization of the malignant cell in terms of lineage and stage of differentiation provides valuable information to the oncologist to guide prognosis and appropriate therapy; and immunodeficient states (eg, HIV infection, organ transplantation), in which the alterations in the immune system that occur secondary to infection are evaluated.

The method of lymphocyte quantitation and characterization is based on the detection of cell surface markers by very specific monoclonal antibodies. For cell surface immunophenotyping, flow cytometry has become the method of choice. Cell surface phenotyping is accomplished by reacting cells from an appropriate specimen with one or more labeled monoclonal antibodies and passing them through a flow cytometer, which counts the proportion of labeled cells.

Reference Values
Normal for Adult Peripheral Blood by Flow Cytometry

- T and B surface markers: Total T cells (CD3⁺): 53%–88%
- Helper T cells (CD3⁺, CD4⁺): 32%–61% Suppressor T cells (CD3⁺, CD8⁺): 16%–42% B cells (CD19⁺): 5%–20%
- Natural killer cells (CD16⁺): 4%–32% Absolute counts (based on pathologist’s interpretation): Total lymphocytes: 660–4600/mm³ (0.6–4.6 × 10⁹/L) Total T cells (CD3⁺): 812–2318/mm³ Helper T cells (CD3⁺, CD4⁺): 589–1505/mm³ Suppressor T cells (CD3⁺, CD8⁺): 325–997/mm³ B cells (CD19⁺): 92–426/mm³ Natural killer cells (CD16⁺): 78–602/mm³ Lymphocyte ratio: Helper-to-suppressor T-cell ratio >1.0

Procedure
1. Obtain a 5-mL EDTA-anticoagulated blood sample (lavender-topped tube).
2. Do not refrigerate or freeze the sample; it should remain at room temperature until testing is performed. Collect a separate 5-mL venous EDTA-anticoagulated blood sample for hematology at the same time. Because the interpretation of data is based on absolute values, it is imperative that a WBC and differential count also be performed so that the appropriate data can be obtained.

Clinical Implications
1. Standard immunosuppressive drug therapy usually decreases lymphocyte totals.
2. Patients with an absolute helper T-lymphocyte count <200/mm³ are at greatest risk for developing clinical AIDS.
3. Decreased T cells occur in congenital immunodeficiency diseases (eg, DiGeorge syndrome, thymic hypoplasia).
4. Decreased T cells occur in kidney and heart transplant patients receiving OKT-3, an immunomodulatory drug used to prevent rejection.
5. A marked increase in B cells occurs in lymphoproliferative disorders (eg, chronic lymphocytic leukemia). In the typical case of chronic lymphocytic leukemia, the B cells would be positive for either ? or ? light chains (indicating mononclonality) and would express CD19 (a B-cell antigen).

Interventions
Pretest Patient Care
1. Explain purpose and specimen collection procedure. A recent viral cold can cause a decrease in total T cells, as
can medications such as corticosteroids. Nicotine and strenuous exercise have also been shown to decrease
lymphocyte counts.
Posttest Patient Aftercare

1. Interpret test outcomes and possible need for repeat testing. Lymphocyte immunophenotyping is performed to monitor patients who are HIV positive and have begun medication treatment. Transplantation patients are also retested at regular intervals to assess the threat of organ rejection or host infection. Also, see Chapter 8 for further discussion of CD4 and CD8 cells.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

STAINS FOR LEUKEMIAS

Several special WBC staining methods are used to diagnose leukemia, amyloid disease, lymphoma, and erythroleukemia; to differentiate erythema myelosis from sideroblastic anemia; to monitor progress and response to therapy; and to detect early relapse. Amyloid refers to starchlike substances deposited in certain diseases (eg, tuberculosis, osteomyelitis, leprosy, Hodgkin’s disease, and carcinoma).

Sudan Black B (SBB) Stain

The SBB stain aids in differentiation of the immature cells of acute leukemias, especially acute myeloblastic leukemia. The SBB stains a variety of fats and lipids that are present in myeloid leukemias but not present in the lymphoid leukemias.

Reference Values

**Positive Reactions**
- Granulocytic cells (neutrophils and eosinophils)
- Myeloblasts
- Promyelocytes
- Neutrophilic myelocytes
- Metamyelocytes, bands, and segmented neutrophils
- Eosinophils at all stages
- Monocytes and precursors

**Variable Reactions**
- Basophils

**Negative Reactions (Sudanophobia)**
- Lymphocytes and lymphocytic precursors
- Megakaryocytes and thrombocytes (platelets)
- Erythrocytes
- Erythroblasts may display a few granules that represent mitochondrial phospholipid components

**Procedure**

1. Obtain bone marrow aspirate.
2. Prepare slide, stain with SBB, and scan microscopically. Use normal smear control.

Clinical Implications

1. Positive staining of primitive (blast) cells indicates myelogenous origin of cells. SBB is positive in acute myelocytic leukemia (AML).
2. SBB is negative in acute lymphocytic leukemia, monocytic leukemia, plasma cell leukemia, and megakaryocyte leukemia.
3. SBB is weak to negative in acute monocytic leukemia.

Interfering Factors

There are cases of acute leukemia in which the cytochemical stains are not useful and fail to reveal the differentiating features of any specific cell line.

Interventions

**Pretest Patient Care**

1. Explain test purposes and procedures. If bone marrow aspiration is done, see pages for special care.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes; counsel and monitor appropriately for leukemia, amyloid disease, anemia, and infection.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Periodic Acid–Schiff (PAS) Stain

The PAS stain aids in the diagnosis of acute lymphoblastic leukemia (ALL). Early myeloid precursors and erythrocyte precursors are negative. As granulocytes mature, they increase in PAS positivity, whereas mature RBCs stay negative. The PAS stain cannot be used to distinguish between AAL and AML or between benign and malignant lymphocytic disorders.

Reference Values

**Normal**
- Lymphoblasts: stain (positive)
- Myeloblasts: do not stain (negative)

**Procedure**

1. Obtain bone marrow aspirate.
2. Prepare slide, stain with PAS, and scan microscopically.

Clinical Implications

1. Positive reaction
   a. Blasts in ALL in childhood often have coarse clumps or masses of PAS-positive material within their scent cytoplasm. The staining pattern is usually heterogeneous, with some cells containing PAS-positive clumps and
others virtually unstained.

b. Acute monocytic leukemia
c. Hairy cell leukemia
d. Sézary's syndrome
e. Conspicuous PAS positivity in the erythroid precursors is strongly suggestive of erythroleukemia (M₆).

2. Weakly positive
   a. In acute granulocytic leukemia, the blasts display either a negative or weakly positive, finely granular pattern.
   b. In some cases of thalassemia and in anemias with blocked or deficient iron, the red blood cell precursors also contain PAS-positive material.
   c. Hodgkin's disease, now Hodgkin's lymphoma
d. Infectious mononucleosis

3. Negative stain
   a. Lymphoblasts of Burkitt's lymphoma
   b. Megaloblastic leukemia

Interventions

Pretest Patient Care

1. Explain test purposes and procedures.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes; counsel and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Terminal Deoxynucleotidyl Transferase (TDT) Stain

The thymus is the primary site of TDT-positive cells, and TDT is found in the nucleus of the more primitive T cells. A thymus-related population of TDT-positive cells resides in the bone marrow (normally a minor population, 0%–2%). TDT is increased in >90% of cases of ALL of childhood. A minor (5%–10%) population of patients with acute nonlymphoblastic leukemia have TDT-positive blasts. TDT-positive blasts are prominent in some cases of chronic myelogenous leukemia (CML), relating to the development of an acute blast phase. TDT has been reported to assist in establishing the diagnosis of ALL. TDT-positive cases of blast-phase CML correlate with a positive response to chemotherapy (vincristine and prednisone).

Reference Values

Normal
Negative in nonlymphoblastic leukemia Negative in peripheral blood 0% to 2% positive in bone marrow

Procedure

1. Obtain a 5-mL EDTA-anticoagulated peripheral blood sample or a 2-mL EDTA-anticoagulated bone marrow aspirate.
2. Dry slides (store at room temperature for up to 5 days), process, and stain, then examine under the microscope for positive cells.

Clinical Implications

1. TDT is positive in ALL, lymphoblastic lymphoma, and CML (blast crisis).
2. TDT is negative in patients in remission and in those with CML or chronic lymphatic leukemia.

Interventions

Pretest Patient Care

1. Explain test purposes and procedures.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes; counsel and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Leukocyte Alkaline Phosphatase (LAP) Stain

Neutrophils are the only leukocytes to contain various amounts of alkaline phosphatase.

The LAP stain is used as an aid to distinguish chronic granulocytic leukemia from a leukemoid reaction. A leukemoid reaction is a high WBC that may look like leukemia but is not. In remission of CML, the LAP may return to normal. In the blast phase of CMC, the LAP may be elevated.

Reference Values

Normal 40–100 LAP units
Each laboratory must establish its own normal values.

**Procedure**

1. Obtain specimen by capillary puncture, venous blood (EDTA), or bone marrow aspirate. Prepare smear and air-dry; stain with LAP.
2. Make a count of 100 granulocytes and score (from 0 to 4+) as to the degree of LAP units.

**Clinical Implications**

1. **Decreased** values (0–15 LAP units):
   a. CML
   b. Paroxysmal nocturnal hemoglobinuria (PNH)
   c. Idiopathic thrombocytopenic purpura
   d. Hereditary hypophosphatasia
   e. Progressive muscular dystrophy
   f. Marked eosinophilia
   g. Nephrotic syndrome
   h. Siderocytic anemia
2. **Increased** values:
   a. Leukemoid reactions, all kinds of neutrophilia with elevated WBC
   b. Polycythemia vera
   c. Thrombocytopenia (essential)
   d. Down syndrome (trisomy 21)
   e. Multiple myeloma
   f. Hodgkin’s disease
   g. Hairy cell leukemia
   h. Aplastic leukemia, acute and chronic lymphatic leukemia, chronic granulocytic leukemia
   i. Myelofibrosis, myeloid metaplasia
3. Normal levels of LAP:
   a. Secondary polycythemia
   b. Hemolytic anemia
   c. Infectious mononucleosis
   d. Iron-deficiency anemia
   e. Viral hepatitis
4. Serial LAP tests can be a useful adjunct in evaluating the activity of Hodgkin’s disease and the response to therapy.

**Interfering Factors**

1. Any physiologic stress, such as third-trimester pregnancy, labor, or severe exercise, causes an increased LAP score.
2. Steroid therapy increases LAP score.
3. CML with infection increases the LAP score.

**Interventions**

**Pretest Patient Care**

1. Explain test purposes and procedures.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes; counsel and monitor appropriately for blood diseases.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Tartrate-Resistant Acid Phosphatase (TRAP) Stain**

The malignant mononuclear cells of leukemic reticuloendotheliosis (hairy cell leukemia) are resistant to inhibition by tartaric acid. There is evidence that the reaction is not entirely specific because TRAP reactions have been reported in prolymphocytic leukemia and malignant lymphoma and in some cases of infectious mononucleosis.

**Reference Values**

**Normal** No TRAP activity

**Procedure**

1. Obtain venous blood sample (5 mL) or bone marrow smear.
2. Incubate blood smear with TRAP, counterstain, and examine microscopically.

**Clinical Implications**

1. TRAP is present in the leukemic cells of most patients with hairy cell leukemia; 5% of patients with otherwise typical hairy cell leukemia lack the enzyme.
2. TRAP occasionally occurs in malignant cells of patients with lymphoproliferative disorders other than hairy cell leukemia.
3. Histiocytes have weakly positive reactions.

**Interventions**

**Pretest Patient Care**

1. Explain test purposes and procedures. Assess for history of signs and symptoms of leukemia.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes; counsel and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

### TESTS OF RED BLOOD CELLS

Many tests look at the red blood cells: their number and size, amount of Hb, rate of production, and percent composition of the blood. The red blood cell count (RBC), hematocrit (Hct), and hemoglobin (Hb) are closely related but different ways to look at the adequacy of erythrocyte production. The same conditions cause an increase (or decrease) in each of these indicators.

**Peripheral Red Blood Cell Abnormalities**

<table>
<thead>
<tr>
<th>Abnormality</th>
<th>Description</th>
<th>Associated Diseases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anisocytosis</td>
<td>Abnormal variation in size (normal diameter = 6–8 µm)</td>
<td>Any severe anemia (eg, iron-deficiency, hemolytic hypersplenism)</td>
</tr>
<tr>
<td>Microcytes</td>
<td>Small cells, &lt;6 µm (MCV &lt;80 fL)</td>
<td>Iron-deficiency and iron-loading (sideroblastic) anemia, thalassemia, lead poisoning, vitamin B&lt;sub&gt;12&lt;/sub&gt; deficiency</td>
</tr>
<tr>
<td>Macrocytes</td>
<td>Large cells, &gt;8 µm (MCV &gt;100 fL)</td>
<td>Megaloblastic anemia, alcoholism, liver disease, hemolytic anemia (reticulocytes), hemolytic disease of newborn, myeloma, leukemia, myelophthisic anemia, metastatic carcinoma, hypothyroidism</td>
</tr>
<tr>
<td>Megalocytes</td>
<td>Large (&gt;9 µm) oval cells</td>
<td>Megaloblastic anemia, pernicious anemia, cancer chemotherapy</td>
</tr>
<tr>
<td>Hypochromia</td>
<td>Pale cells with decreased concentration of hemoglobin (MCHC &lt;30 g/dL)</td>
<td>Severe iron-deficiency and iron-loading (sideroblastic) anemia, thalassemia, lead poisoning, transferrin deficiency</td>
</tr>
<tr>
<td>Poikilocytes</td>
<td>Abnormal variation in shape</td>
<td>Any severe anemia (eg, megaloblastic, iron-deficiency, myeloproliferative syndrome, hemolytic); certain shapes are diagnostically helpful (see entries for Spherocytes, Elliptocytes, Stomatocytes, Sickle cells, Target cells, Schistocytes, Burr cells, Acanthocytes and Teardrop cells)</td>
</tr>
<tr>
<td>Spherocytes</td>
<td>Spherical cells without pale centers; often small (ie, microspherocytosis)</td>
<td>Hereditary spherocytosis, Coombs'–positive hemolytic anemia; small numbers are seen in any hemolytic anemia and after transfusion of stored blood</td>
</tr>
<tr>
<td>Elliptocytes</td>
<td>Oval cells—elongated</td>
<td>Hereditary elliptocytosis (&gt;25% on smear), iron deficiency</td>
</tr>
<tr>
<td>Stomatocytes</td>
<td>Red cells with slitlike (instead of circular) areas of central pallor</td>
<td>Congenital stomatocytosis, Rh-null disease, alcoholism, liver disease, artifact</td>
</tr>
<tr>
<td>Sickle cells</td>
<td>Crescent-shaped cells</td>
<td>Sickle cell disease (Hb S)</td>
</tr>
<tr>
<td>Target cells</td>
<td>Cells with a dark center and periphery and a clear ring in between</td>
<td>Liver disease, thalassemia, iron-deficiency anemia, hemoglobinopathies, (S, C, S-C, S-thalassemia), artifact</td>
</tr>
<tr>
<td>Schistocytes (helmet cells)</td>
<td>Irregularly contracted cells (severe poikilocytosis), fragmented cells</td>
<td>Vasculitis, artificial heart valve, disseminated intravascular coagulation, thrombocytopenia purpura and other microangiopathic anemias, toxins (lead, phenyldiazine, snake bite), severe burns, renal graft rejection, and march hemoglobinuria</td>
</tr>
<tr>
<td>Burr cells (echinocytes)</td>
<td>Burr-like cells, spinous processes</td>
<td>Usually artifactual, uremia, stomach cancer, pyruvate kinase deficiency</td>
</tr>
<tr>
<td>Acanthocytes</td>
<td>Small cells with thorny projections</td>
<td>Abetalipoproteinemia (hereditary acanthocytosis or Bassen-Kornzweig disease), postsplenectomy, hemolytic anemia, alcohol cirrhosis, hepatitis of newborns, malabsorption states</td>
</tr>
<tr>
<td>Teardrop cells, (dacrocytes)</td>
<td>Cells shaped like teardrops</td>
<td>Myeloproliferative syndrome, myelophthisic anemia (neoplastic, granulomatous, or fibrotic marrow infiltration), thalassemia, pernicious anemia, tuberculosis</td>
</tr>
<tr>
<td>Nucleated red cells</td>
<td>Erythrocytes with nuclei still present, normoblastic or megaloblastic</td>
<td>Hemolytic anemias, leukemias, myeloproliferative syndrome, polycythemia vera, myelophthisic anemia (neoplastic, granulomatous, or fibrotic marrow infiltration), multiple myeloma, extramedullary hematopoiiesis, megaloblastic anemias, any severe anemia</td>
</tr>
</tbody>
</table>
Howell-Jolly bodies: Spherical purple bodies (Wright's) within or on erythrocytes, nuclear debris. Hyposplenism, postsplenectomy pernicious anemia, thalassemia, sickle cell anemia, other hemolytic anemias.

Heinz inclusion bodies: Small round inclusions of denatured hemoglobin seen under phase microscopy or with supravital staining. Congenital hemolytic anemias (eg, glucose-6-phosphate dehydrogenase deficiency), hemolytic anemia secondary to drugs (dapsone, phenacetin), thalassemia (Hb H), hemoglobinopathies (Hb Zurich, Koln, Ube, I, and so on).

Pappenheimer bodies (siderocytes): Siderotic granules, staining blue with Wright or Prussian blue stain. Iron-loading anemias (eg, sideroblastic anemia), hyposplenism, lead poisoning, iron overload (hemochromatosis).

Cabot's rings: Purple, fine, ringlike, intrerythrocytic structure. Pernicious anemia, lead poisoning, severe hemolytic anemia.

Basophilic stippling: Punctate stippling when Wright stained. Hemolytic anemia, punctate stippling seen in lead poisoning (mitochondrial RNA and iron), thalassemia, megaloblastic anemia, alcoholism.

Pappenheimer bodies (siderocytes): Siderotic granules, staining blue with Wright or Prussian blue stain. Iron-loading anemias (eg, sideroblastic anemia), hyposplenism, lead poisoning, iron overload (hemochromatosis).

Rouleaux: Aggregated erythrocytes regularly stacked on one another—"rows of coins." Multiple myeloma, Waldenström's macroglobulinemia, cord blood, pregnancy, hypergammaglobulinemia, hyperfibrinogenemia.

Polychromatophilia (called reticulocytes when stained with supravital stain): RBCs containing RNA, staining a pinkish-blue color; stains supravitally as reticular network with new methylene blue. Hemolytic anemia, blood loss, uremia, after treatment of iron-deficiency or megaloblastic anemia.

**NOTE**
Not seen with Wright's stain. Must do supravital stain.

### Red Blood Cell Count (RBC; Erythrocyte Count)

The main function of the red blood cell (RBC or erythrocyte) is to carry oxygen from the lungs to the body tissues and to transfer carbon dioxide from the tissues to the lungs. This process is achieved by means of the Hb in the RBCs, which combines easily with oxygen and carbon dioxide and gives arterial blood a bright red appearance. To enable use of the maximal amount of Hb, the RBC is shaped like a biconcave disk; this affords more surface area for the Hb to combine with oxygen. The cell is also able to change its shape when necessary to allow for passage through the smaller capillaries.

The RBC test, an important measurement in the evaluation of anemia or polycythemia, determines the total number of erythrocytes in a microliter (cubic millimeter) of blood.

### Reference Values

**Normal** See Table 2.3

**Table 2.3 Normal Values for Red Blood Cells**

<table>
<thead>
<tr>
<th>Age</th>
<th>Men</th>
<th>Women</th>
</tr>
</thead>
<tbody>
<tr>
<td>Birth–2 wk</td>
<td>4.1–6.1 × 10⁶/mm³ or 4.1–6.1 × 10¹²/L (average, 4.8)</td>
<td></td>
</tr>
<tr>
<td>2–6 mo</td>
<td>4.0–6.0 × 10⁶/mm³ or 4.0–6.0 × 10¹²/L (average, 4.3)</td>
<td></td>
</tr>
<tr>
<td>6 mo–1 y</td>
<td>3.8–5.2 × 10⁶/mm³ or 3.8–5.2 × 10¹²/L</td>
<td></td>
</tr>
<tr>
<td>1–6 y</td>
<td>3.9–5.3 × 10⁶/mm³ or 3.9–5.3 × 10¹²/L</td>
<td></td>
</tr>
<tr>
<td>6–16 y</td>
<td>4.0–5.2 × 10⁶/mm³ or 4.0–5.2 × 10¹²/L</td>
<td></td>
</tr>
<tr>
<td>16–18 y</td>
<td>4.2–5.4 × 10⁶/mm³ or 4.2–5.4 × 10¹²/L</td>
<td></td>
</tr>
<tr>
<td>&gt;18 y (males)</td>
<td>4.5–5.5 × 10⁶/mm³ or 4.5–5.5 × 10¹²/L</td>
<td></td>
</tr>
<tr>
<td>&gt;18 y (females)</td>
<td>4.0–5.0 × 10⁶/mm³ or 4.0–5.0 × 10¹²/L</td>
<td></td>
</tr>
</tbody>
</table>

**Procedure**

1. Obtain 5 mL of EDTA-anticoagulated venous blood. Place the specimen in a biohazard bag.
2. Remember that automated electronic devices are generally used to determine the number of RBCs.
3. Note patient age and time of day on the laboratory slip.

**Clinical Implications**

1. **Decreased RBC values** occur in:
   a. Anemia, a condition in which there is a reduction in the number of circulating erythrocytes, the amount of Hb, or the volume of packed cells (Hct). Anemia is associated with cell destruction, blood loss, or dietary insufficiency.
of iron or of certain vitamins that are essential in the production of RBCs. See Chart 2.1 on page 79 for a classification of anemias based on their underlying mechanisms and the test for reticulocyte count for a discussion of the purpose and clinical implications of the reticulocyte count.

b. Disorders such as:
   1. Hodgkin's disease and other lymphomas
   2. Multiple myeloma, myeloproliferative disorders, leukemia
   3. Acute and chronic hemorrhage
   4. Lupus erythematosus
   5. Addison's disease
   6. Rheumatic fever
   7. Subacute endocarditis, chronic infection
   8. This list is not meant to be all inclusive.

2. Erythrocytosis (increased RBC) occurs in:
   a. Primary erythrocytosis
      1. Polycythemia vera (myeloproliferative disorder)
   b. Secondary erythrocytosis
      1. Renal disease
      2. Extrarenal tumors
      3. High altitude
      4. Pulmonary disease
      5. Cardiovascular disease
      6. Alveolar hypoventilation
      7. Hemoglobinopathy
      8. Tobacco/carboxyhemoglobin
   c. Relative erythrocytosis (decrease in plasma volume)
      1. Dehydration (vomiting, diarrhea)
      2. Gaisböck's syndrome

---

**Clinical Alert**

Refer to page 76 for a discussion of the combined clinical implications of decreased RBC, Hct, and Hb values. The same underlying conditions cause a decrease in each of these three tests of erythrocyte production.

**Clinical Alert**

Please refer to page 75 for a discussion of the combined clinical implications of increased RBC, Hct, and Hb values. The same underlying conditions cause an increase in each of these three tests of erythrocyte production.

---

Interfering Factors

1. Posture: when a blood sample is obtained from a healthy person in a recumbent position, the RBC is 5% lower. (If the patient is anemic, the count will be lower still.)
2. Dehydration: hemoconcentration in dehydrated adults (caused by severe burns, untreated intestinal obstruction, severe persistent vomiting, or diuretic abuse) may obscure significant anemia.
3. Age: the normal RBC of a newborn is higher than that of an adult, with a rapid drop to the lowest point in life at 2 to 4 months. The normal adult level is reached at age 14 years and is maintained until old age, when there is a gradual drop (see normal values).
4. Falsely high counts may occur because of prolonged venous stasis during venipuncture.
5. Stress can cause a higher RBC.
6. Altitude: the higher the altitude, the greater the increase in RBC. Decreased oxygen content of the air stimulates the RBC to rise (erythrocytosis).
7. Pregnancy: there is a relative decrease in RBC when the body fluid increases in pregnancy, with the normal number of erythrocytes becoming more diluted.
8. There are many drugs that may cause decreased or increased RBC. See Appendix J for drugs that affect test outcomes.
9. The EDTA blood sample tube must be at least three fourths filled or values will be invalid because of cell shrinkage caused by the anticoagulant.
10. The blood sample must not be clotted (even slightly) or the values will be invalid.

---

Interventions

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. Refer to standard pretest care for hemogram, CBC, and differential count on page 47.
3. Have the patient avoid extensive exercise, stress, and excitement before the test. These cause elevated counts of doubtful clinical value.
4. Avoid overhydration or dehydration, if possible; either causes invalid results. If patient is receiving IV fluids or therapy, note on requisition.
5. Note any medications patient is taking.
6. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and monitor appropriately for anemia and erythrocytosis.
2. Refer to standard posttest care for hemogram, CBC, and differential count on page 47.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.
4. Resume normal activities and diet.

Hematocrit (Hct); Packed Cell Volume (PCV)

The word hematocrit means "to separate blood," which underscores the mechanism of the test because the plasma and blood cells are separated by centrifugation.

The Hct test is part of the CBC. This test indirectly measures the RBC mass. The results are expressed as the percentage by volume of packed RBCs in whole blood (PCV). It is an important measurement in the determination of anemia or polycythemia.

<table>
<thead>
<tr>
<th>Reference Values</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Normal</strong> Women: 36%–48% or 0.36–0.48 Men: 42%–52% or 0.42–0.52 Children: 0–2 weeks: 44%–64% or 0.44–0.64 2–8 weeks: 39%–59% or 0.39–0.59 2–6 months: 35%–49% or 0.35–0.49 6 months–1 year: 29%–43% or 0.29–0.43 1–6 years: 30%–40% or 0.30–0.40 6–16 years: 32%–42% or 0.32–0.42 16–18 years: 34%–44% or 0.34–0.44</td>
</tr>
</tbody>
</table>

**NOTE**

If blood is drawn from a capillary puncture and a microhematocrit is done, values are slightly higher.

**Procedure**

1. Observe standard precautions. When doing a capillary puncture (finger puncture), the microcapillary tube is filled three fourths full with blood, directly from puncture site. These tubes are coated with an anticoagulative.
2. Centrifuge the tubes in a microcentrifuge and measure the height of packed cells in the tube.
3. Record the measurement as a percentage of the total amount of blood in the capillary tube.
4. Remember that an Hct can be done on automated hematology instruments, in which case a 5-mL EDTA-anticoagulated venous blood sample is obtained.

**Clinical Implications**

1. Decreased Hct values are an indicator of anemia, a condition in which there is a reduction in the PVC. An Hct <30% (<0.30) means that the patient is moderately to severely anemic. Decreased values also occur in the following conditions:
   a. Leukemias, lymphomas, Hodgkin's disease, myeloproliferative disorders
   b. Adrenal insufficiency
   c. Chronic disease
   d. Acute and chronic blood loss
   e. Hemolytic reaction: this condition may be found in transfusion of incompatible blood or as a reaction to chemicals or drugs, infectious agents, or physical agents (eg, severe burns, prosthetic heart valves).
2. The Hct may or may not be reliable immediately after even a moderate loss of blood or immediately after transfusion.
3. The Hct may be normal after acute hemorrhage. During the recovery phase, both the Hct and the RBC drop markedly.
4. Usually, the Hct parallels the RBC when the cells are of normal size. As the number of normal-sized erythrocytes increases, so does the Hct.
   a. However, for the patient with microcytic or macrocytic anemia, this relationship does not hold true.
   b. If a patient has iron-deficiency anemia with small RBCs, the Hct decreases because the microcytic cells pack to a smaller volume. The RBC, however, may be normal or higher than normal.
5. Increased Hct values occur in:
   a. Erythrocytosis
   b. Polycythemia vera
   c. Shock, when hemoconcentration rises considerably

**Clinical Alert**

Please refer to page 76 for a discussion of the combined clinical implications of decreased Hct, Hb, and RBC values. The same underlying conditions cause a decrease in each of these three tests of erythrocyte production.

**Clinical Alert**

Please refer to page 75 for a discussion of the combined clinical implications of increased Hct, Hb, and RBC values. The same underlying conditions cause an increase in each of these three tests of erythrocyte production.

**Interfering Factors**

1. People living at high altitudes have high Hct values as well as high Hb and RBC.
2. Normally, the Hct slightly decreases in the physiologic hydremia of pregnancy.
3. The normal values for Hct vary with age and gender. The normal value for infants is higher because the newborn has many macrocytic red cells. Hct values in females are usually slightly lower than in males.
4. There is also a tendency toward lower Hct values in men and women older than 60 years of age, corresponding to lower RBC values in this age group.
5. Severe dehydration from any cause falsely raises the Hct.

**Interventions**
Pretest Patient Care

1. Explain test purpose and procedure.
2. Refer to standard pretest care for hemogram, CBC, and differential count on page 47. Also, see Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert

An Hct <20% (<0.20) can lead to cardiac failure and death; an Hct >60% (>0.60) is associated with spontaneous clotting of blood.

Posttest Patient Aftercare

1. Interpret test results and monitor for anemia or polycythemia.
2. Refer to standard posttest care for hemogram, CBC, and differential count on page 47. Also, follow Chapter 1 guidelines for safe, effective, informed posttest care.

Hemoglobin (Hb)

Hb, the main component of erythrocytes, serves as the vehicle for the transportation of oxygen and carbon dioxide. It is composed of amino acids that form a single protein called globin, and a compound called heme, which contains iron atoms and the red pigment porphyrin. It is the iron pigment that combines readily with oxygen and gives blood its characteristic red color. Each gram of Hb can carry 1.34 mL of oxygen per 100 mL of blood. The oxygen-combining capacity of the blood is directly proportional to the Hb concentration rather than to the RBC because some RBCs contain more Hb than others. This is why Hb determinations are important in the evaluation of anemia.

The Hb determination is part of a CBC. It is used to screen for disease associated with anemia, to determine the severity of anemia, to monitor the response to treatment for anemia, and to evaluate polycythemia.

Hb also serves as an important buffer in the extracellular fluid. In tissue, the oxygen concentration is lower, and the carbon dioxide level and hydrogen ion concentration are higher. At a lower pH, more oxygen dissociates from Hb. The unoxygenated Hb binds to hydrogen ion, thereby raising the pH. As carbon dioxide diffuses into the RBC, carbonic anhydrase converts carbon dioxide to bicarbonate and protons. As the protons are bound to Hb, the bicarbonate ions leave the cell. For every bicarbonate ion leaving the cell, a chloride ion enters. The efficiency of this buffer system depends on the ability of the lungs and kidneys to eliminate, respectively, carbon dioxide and bicarbonate. Refer to the discussion of arterial blood gases in Chapter 14.

Reference Values

Normal: Women: 12.0–16.0 g/dL or 120–160 g/L Men: 14.0–17.4 g/dL or 140–174 g/L Children: 0–2 weeks: 14.5–24.5 g/dL or 145–245 g/L 2–8 weeks: 12.5–20.5 g/dL or 125–205 g/L 2–6 months: 10.7–17.3 g/dL or 107–173 g/L 6 months–1 year: 9.9–14.5 g/dL or 99–145 g/L 1–6 years: 9.5–14.1 g/dL or 95–141 g/L 6 months–1 year: 9.9–14.5 g/dL or 99–145 g/L 1–6 years: 9.5–14.1 g/dL or 95–141 g/L

Procedure

1. Obtain a venous blood EDTA-anticoagulated sample of 5 mL. Fill the Vacutainer tube at least three fourths full. Automated electronic devices are generally used to determine the Hb; however, a manual colorimetric procedure is also widely used.
2. Do not allow the blood sample to clot, or the results will be invalid. Place the specimen in a biohazard bag.

Clinical Implications

1. Decreased Hb levels are found in anemia states (a condition in which there is a reduction of Hb, Hct, and/or RBC values). The Hb must be evaluated along with the RBC and Hct.
   a. Iron deficiency, thalassemia, pernicious anemia, hemoglobinopathies
   b. Liver disease, hypothyroidism
   c. Hemorrhage (chronic or acute)
   d. Hemolytic anemia caused by:
      1. Transfusions of incompatible blood
      2. Reactions to chemicals or drugs
      3. Reactions to infectious agents
      4. Reactions to physical agents (eg, severe burns, artificial heart valves)
      5. Various systemic diseases:
         a. Hodgkin's disease
         b. Leukemia
         c. Lymphoma
         d. SLE
         e. Carcinomatosis
         f. Sarcoidosis
         g. Renal cortical necrosis
      h. This list is not meant to be all inclusive.
2. Increased Hb levels are found in:
   a. Polycythemia vera
   b. Congestive heart failure
   c. Chronic obstructive pulmonary disease (COPD)
3. Variation in Hb levels:
a. Occurs after transfusions, hemorrhages, burns. (Hb and Hct are both high during and immediately after hemorrhage.)

b. The Hb and Hct provide valuable information in an emergency situation if they are interpreted not in an isolated fashion but in conjunction with other pertinent laboratory data.

Clinical Alert

Please refer to page 76 for a discussion of the combined clinical implications of decreased Hb, Hct, and RBC values. The same underlying conditions cause a decrease in each of these three tests of erythrocyte production.

Clinical Alert

Please refer below for a discussion of the combined clinical implications of increased Hb, Hct, and RBC values. The same underlying conditions cause an increase in each of these three tests of erythrocyte production.

Clinical Implications of Polycythemia: Increased RBC, Hct, and/or Hb

Polycythemia is the term used to describe an abnormal increase in the number of RBCs. Although there are several tests to directly determine the RBC mass, these tests are expensive and somewhat cumbersome. For screening purposes, we rely on the Hct and Hb to evaluate polycythemia indirectly. Polycythemas are classified as follows:

1. Relative polycythemia: an increase in Hb, Hct, or RBC caused by a decrease in the plasma volume (eg, dehydration, spurious erythrocytosis from stress or smoking)
2. Absolute or true polycythemia:
   a. Primary (eg, polycythemia vera, erythemic erythrocytosis)
   b. Secondary
      1. Appropriate (an appropriate bone marrow response to physiologic conditions)
         a. Altitude
         b. Cardiopulmonary disorder
         c. Increased affinity for oxygen
      2. Inappropriate (an overproduction of RBCs not necessary to deliver oxygen to the tissues)
         a. Renal tumor or cyst
         b. Hepatoma
         c. Cerebellar hemangioblastoma

Clinical Implications of Anemia: Decreased RBC, Hct, and/or Hb

Anemia is the term used to describe a condition in which there is a reduction in the number of circulating RBCs, the amount of Hb, and/or volume of packed cells (Hct). A pathophysiologic classification of anemias based on their underlying mechanisms follows. Anemias are further explained in Chart 2.1. Anemias are classified as follows:

1. Hypoproliferative anemias (inadequate production of RBCs):
   a. Marrow aplasias
   b. Myelophthisic anemia
   c. Anemia with blood dyscrasias
   d. Anemia of chronic disease
   e. Anemia with organ failure
2. Maturation defect anemias:
   a. Cytoplasmic: hypochromic anemias
   b. Nuclear: megaloblastic anemias
   c. Combined: myelodysplastic syndromes
3. Hyperproliferative anemias (decreased Hb or Hct despite an increased production of RBCs):
   a. Hemorrhagic: acute blood loss
   b. Hemolytic: a premature, accelerated destruction of RBCs
      1. Immune hemolysis
      2. Primary membrane
      3. Hemoglobinopathies
      4. Toxic hemolysis (physical-chemical)
      5. Traumatic or microangiopathic hemolysis
      6. Hypersplenism
      7. Enzymopathies
      8. Parasitic infections
4. Dilutional anemias:
   a. Pregnancy
   b. Splenomegaly

Interfering Factors

1. People living at high altitudes have increased Hb values as well as increased Hct and RBC.
2. Excessive fluid intake causes a decreased Hb.
3. Normally, the Hb is higher in infants (before active erythropoiesis begins).
4. Hb is normally decreased in pregnancy as a result of increased plasma volume.
5. There are many drugs that may cause a decreased Hb. Drugs that may cause an increased Hb include gentamicin and methyldopa.
6. Extreme physical exercise causes increased Hb.

Interventions
Pretest Patient Care

1. Explain test purpose and procedure. Assess medication history.
2. Refer to standard pretest care for hemogram, CBC, and differential count on page 47. Also, see Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and monitor appropriately for anemia or polycythemia.
2. Refer to standard posttest care for hemogram, CBC, and differential count on page 47. Also, follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

The panic Hb value is <5.0 g/dL (<50 g/L), a condition that leads to heart failure and death. A value >20 g/dL (>200 g/L) leads to clogging of the capillaries as a result of hemoconcentration.

Red Blood Cell Indices

The red cell indices define the size and Hb content of the RBC and consist of the mean corpuscular volume (MCV), the mean corpuscular hemoglobin concentration (MCHC), and the mean corpuscular hemoglobin (MCH).

The RBC indices are used in differentiating anemias. When they are used together with an examination of the erythrocytes on the stained smear, a clear picture of RBC morphology may be ascertained. On the basis of the RBC indices, the erythrocytes can be characterized as normal in every respect or as abnormal in volume or Hb content. In deficient states, the anemias can be classified by cell size as macrocytic, normocytic, or microcytic, or by cell size and color as microcytic hypochromic.

Procedure

1. Remember that these are calculated values. An explanation of each measurement follows.
2. Obtain 5 mL EDTA blood so that RBC, Hb, and Hct determinations can be done for calculations.

Interventions

Pretest Patient Care for MCV, MCHC, and MCH

1. Explain the purpose and procedure for testing. Assess for possible causes of anemia. No fasting is required.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare for MCV, MCHC, and MCH

1. Interpret test results and monitor appropriately for anemia. Counsel appropriately for proper diet, medication, related hormone and enzyme problems, and genetically linked disorders.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Mean Corpuscular Volume (MCV)

Individual cell size is the best index for classifying anemias. This index expresses the volume occupied by a single erythrocyte and is a measure in cubic micrometers (femtoliters, or fL) of the mean volume. The MCV indicates whether the red blood cell size appears normal (normocytic), smaller than normal (<82 µm$^3$, microcytic), or larger than normal (>100 µm$^3$, macrocytic).

Reference Values

Normal 82–98 mm$^3$ or 82–98 fL (higher values in infants and newborns and for elderly patients)

Procedure

1. Calculate the MCV from the RBC count (the number of cells per cubic millimeter of blood) and the Hct (the proportion of the blood occupied by the RBCs).
2. Use the following formula:

   \[
   \text{MCV (fL)} = \frac{\text{Hct} \times 10}{\text{RBC} \times 10^{12}/L}
   \]

Clinical Implications The MCV results are the basis of the classification system used to evaluate anemia. The categorizations shown in Chart 2.1 aid in orderly investigation.
Chart 2.1 Anemias Characterized by Deficient Hemoglobin Synthesis

Microcytic Anemias (MCV 50–82 fL)

**DISORDERS OF IRON METABOLISM** Iron-deficiency anemia: the most prevalent worldwide cause of anemia; the major causes are dietary inadequacy, malabsorption, increased iron loss, and increased iron requirements Anemia of chronic disease, hereditary spherocytosis, congenital dyserythropoietic microcytic anemia with iron overload (Shahidi–Nathan–Diamond syndrome)

**DISORDERS OF Porphyrin and Heme Synthesis** Acquired sideroblastic anemia; idiopathic refractory sideroblastic anemia, complicating other diseases associated with drugs or toxins (ethanol, isoniazid, lead) Hereditary sideroblastic anemias X chromosome–linked, autosomal anemias

**DISORDERS OF GLOBIN SYNTHESIS** Thalassemias, hemoglobinopathies, characterized by unstable hemoglobins

Normocytic Normochromic Anemias (MCV 82–98 fL)

**ANEMIA WITH APPROPRIATE BONE MARROW RESPONSE** Acute posthemorrhagic anemia Hemolytic anemia (may be macrocytic when there is pronounced reticulocytosis)

**ANEMIA WITH IMPAIRED MARROW RESPONSE**

Marrow hypoplasia Aplastic anemia, pure red cell aplasia

Marrow infiltration Infiltration by malignant cells, myelofibrosis, inherited storage diseases

Decreased erythropoietin production Kidney and liver disease, endocrine deficiencies, malnutrition, anemia of chronic disease

Macrocytic Anemias (MCV 100–150 fL)

**COBALAMIN (B 12) DEFICIENCY**

Decreased ingestion Lack of animal products, strict vegetarianism

Impaired absorption Intrinsic factor deficiency, pernicious anemia, gastrectomy (total or partial), destruction of gastric mucosa by caustics, anti-intrinsic factor antibody in gastric juice, abnormal intrinsic factor molecule, intrinsic intestinal disease, familial selective malabsorption, ileitis, sprue, celiac disease, infiltrative intestinal disease (eg, lymphoma, scleroderma) drug-induced malabsorption

Competitive parasites Fish tapeworm infestations (Diphyllobothrium latum); bacteria in diverticulum of bowel, blind loops

Increased requirements Chronic pancreatic disease, pregnancy, neoplastic disease, hyperthyroidism

Impaired utilization Enzyme deficiencies, abnormal serum cobalamin binding protein, lack of transcobalamin II, nitrous oxide administration

FOLATE DEFICIENCY

Decreased ingestion Lack of vegetables, alcoholism, infancy

Impaired absorption Intestinal short circuits, steatorrhea, sprue, celiac disease, intrinsic intestinal disease, anticonvulsants, oral contraceptives, other drugs

Increased requirement Pregnancy, infancy, hypothyroidism, hyperactive hematopoiesis, neoplastic disease, exfoliative skin disease

Impaired utilization Folic acid antagonists: methotrexate, trimethoprim, enzyme deficiencies

Increased loss Hemodialysis

UNRESPONSIVE TO COBALAMIN OR FOLATE

**Metabolic inhibitors**

Purine synthesis: 6-mercaptopurine, 6-thioguanine, azathioprine Pyrimidine synthesis: 6-azauridine Thymidylate synthesis: methotrexate, 5-fluorouracil Deoxyribonucleotide synthesis: hydroxyurea, cytarabine, severe iron deficiency

**Inborn errors**

Lesch-Nyhan syndrome, hereditary orotic aciduria, deficiency of formiminotransferase, methyltransferase, others

Interfering Factors

1. Mixed (bimorphic) population of macrocytes and microcytes can result in a normal MCV. Examination of the blood film confirms this.
2. Increased reticulocytes can increase the MCV.
3. Marked leukocytosis increases the MCV.
4. Marked hyperglycemia increases MCV.
5. Cold agglutinins increase MBV.

Interventions

**Pretest Patient Care**

1. Explain test purposes and procedures.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes; counsel and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Mean Corpuscular Hemoglobin Concentration (MCHC)**

The MCHC measures the average concentration of Hb in the RBCs. The MCHC is most valuable in monitoring therapy
for anemia because the two most accurate hematologic determinations (Hb and Hct) are used in its calculation.

**Reference Values**

**Normal** 32–36 g/dL or 320–360 g/L

**Procedure**

1. Remember that the MCHC is a calculated value. It is an expression of the average concentration of Hb in the red blood cells and, as such, represents the ratio of the weight of Hb to the volume of the erythrocyte.

2. Use the following formula:

\[
\text{MCHC (g/dL)} = \frac{\text{Hb (g/dL)} \times 100}{\text{Hct (%)}}
\]

**Clinical Implications**

1. *Decreased MCHC values* signify that a unit volume of packed RBCs contains less Hb than normal. Hypochromic anemia (MCHC <30 g/dL) occurs in:
   a. Iron deficiency
   b. Microcytic anemias, chronic blood loss anemia
   c. Some thalassemias

2. *Increased MCHC values* (RBCs cannot accommodate more than 37 g/dL or 370 g/L Hb) occur in:
   a. Spherocytosis (hereditary)
   b. Newborns and infants

**Interfering Factors**

1. The MCHC may be falsely high in the presence of lipemia, cold agglutinins, or rouleaux and with high heparin concentrations.

2. The MCHC cannot be greater than 37 g/dL (370 g/L) because the RBC cannot accommodate more than 37 g/dL (370 g/L) Hb. (Check for errors in calculation or in Hb determination. The MCHC can be used for laboratory quality control.)

**Interventions**

**Pretest Patient Care**

1. Explain test purposes and procedures.
2. See Chapter 1 guidelines for [safe, effective, informed pretest care](#).

**Posttest Patient Aftercare**

1. Interpret test outcomes; counsel and monitor appropriately.
2. Follow Chapter 1 guidelines for [safe, effective, informed posttest care](#).

**Mean Corpuscular Hemoglobin (MCH)**

The MCH is a measure of the average weight of Hb per RBC. This index is of value in diagnosing severely anemic patients.

**Reference Values**

**Normal** 26–34 pg/cell or 0.40–0.53 fmol/cell (normally higher in newborns and infants)

**Procedure**

The MCH is a calculated value. The average weight of Hb in the RBC is expressed as picograms of Hb per RBC. The formula is:

\[
\frac{\text{MCH (pg/cell)}}{\text{RBC (10^{12}/L)}} = \frac{\text{Hb (g/dL)} \times 10}{\text{RBC (10^{12}/L)}}
\]

**Clinical Implications**

1. An increase of the MCH is associated with macrocytic anemia and newborns.
2. A decrease of the MCH is associated with microcytic anemia.

**Interfering Factors**

1. Hyperlipidemia falsely elevates the MCH.
2. WBC >50,000/mm³ falsely raises the Hb value and therefore falsely elevates the MCH.
3. High heparin concentrations falsely elevate the MCH.
4. Cold agglutinins falsely elevates MCH.

**Interventions**

**Pretest Patient Care**

1. Explain test purposes and procedures.
2. See Chapter 1 guidelines for [safe, effective, informed pretest care](#).
Posttest Patient Aftercare

1. Interpret test outcomes; counsel and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Red Cell Size Distribution Width (RDW)

This automated method of measurement is helpful in the investigation of some hematologic disorders and in monitoring response to therapy. The RDW is essentially an indication of the degree of anisocytosis (abnormal variation in size of RBCs). Normal RBCs have a slight degree of variation.

Reference Values

**Normal** 11.5–14.5 coefficient of variation (CV) of red cell size

Procedure

1. Remember that the CV of RDW is determined and calculated by the analyzer.
2. Use the CV of RDW with caution and not as a replacement for other diagnostic tests.
3. Use the following calculation:

\[
\text{RDW (CV\%)} = \frac{\text{Standard deviation of RBC size} \times 100}{\text{MCV}}
\]

Clinical Implications

1. The RDW can be helpful in distinguishing uncomplicated heterozygous thalassemia (low MCV, normal RDW) from iron-deficiency anemia (low MCV, high RDW).
2. The RDW can be helpful in distinguishing anemia of chronic disease (low-normal MCV, normal RDW) from early iron-deficiency anemia (low-normal MCV, elevated RDW).
3. Increased RDW occurs in:
   a. Iron deficiency
   b. Vitamin B₁₂ or folate deficiency (pernicious anemia)
   c. Abnormal Hb: S, S-C, or H
   d. S-thalassemia (homogeneous)
   e. Immune hemolytic anemia
   f. Marked reticulocytosis
   g. Fragmentation of RBCs
4. Normal RDW—normal in anemias with homogeneous red cell size
   a. Chronic disease
   b. Acute blood loss
   c. Aplastic anemia
   d. Hereditary spherocytosis
   e. Hb E disease
   f. Sickle cell disease
5. There is no known cause of a decreased RDW.

Interfering Factors

1. This test is not helpful for persons who do not have anemia.
2. Alcoholism elevates RDW.
3. Cold agglutinins

Interventions

**Pretest Patient Care**

1. Explain the purpose and procedure for testing. Assess for possible causes of anemia. No fasting is required.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results and monitor appropriately for anemia. Counsel appropriately for proper diet, medication, related hormone and enzyme problems, and genetically linked disorders.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Stained Red Cell Examination (Film; Stained Erythrocyte Examination)

The stained film examination determines variations and abnormalities in erythrocyte size, shape, structure, Hb content, and staining properties. It is useful in diagnosing blood disorders such as anemia, thalassemia, and other hemoglobinopathies. This examination also serves as a guide to therapy and as an indicator of harmful effects of chemotherapy and radiation therapy. The leukocytes are also examined at this time.

Reference Values

**Normal** Size: normocytic (normal size, 7–8 µm) Color: normochromic (normal) Shape: normocyte (biconcave disk)
Structure: normocytes or erythrocytes (anucleated cells)

Procedure
1. Collect a 5-mL blood sample in EDTA. Stain a thin smear with Wright's stain and study under a microscope to determine size, shape, and other characteristics of the RBCs.

2. Be aware that a capillary smear may also be used and may be preferred for detection of some abnormalities.

Clinical Implications

Variations in staining, color, shape, and RBC inclusions are indicative of RBC abnormalities.

Clinical Alert
Marked abnormalities in size and shape of RBCs without a known cause are an indication for more complete blood studies.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure for testing. Assess for possible causes of anemia. No fasting is required.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Reticulocyte Count

A reticulocyte—young, immature, nonnucleated RBC—contains reticular material (RNA) that stains gray-blue. Reticulum is present in newly released blood cells for 1 to 2 days before the cell reaches its full mature state. Normally, a small number of these cells are found in circulating blood. For the reticulocyte count to be meaningful, it must be viewed in relation to the total number of erythrocytes (absolute reticulocyte count = % reticulocytes × erythrocyte count).

The reticulocyte count is used to differentiate anemias caused by bone marrow failure from those caused by hemorrhage or hemolysis (destruction of RBCs), to check the effectiveness of treatment in pernicious anemia and folate and iron deficiency, to assess the recovery of bone marrow function in aplastic anemia, and to determine the effects of radioactive substances on exposed workers.

Reference Values

Normal
Adults: 0.5%–1.5% of total erythrocytes (women may be slightly higher) Newborns: 3%–6% of total erythrocytes (drops to adult levels in 1–2 months) Absolute count: 25–85 × 10^3/mm^3 or × 10^9 cells/L Reticulocyte index (RI): 1% corrected reticulocyte count (CRC) Hematocrit correction for anemia: RI = reticulocyte count × (patient's Hct/45 × 1/1.85)

Procedure

1. Obtain a 5-mL EDTA-anticoagulated venous blood sample. Place the specimen in a biohazard bag.
2. Mix the blood sample with a supravital stain such as brilliant cresyl blue. Allow the stain to react with the blood, and prepare a smear with this mixture and scan under a microscope. Count and calculate the reticulocytes.

3. Use the following formula:

\[
\text{Retic } \% = \frac{\text{total retics}}{1000 \text{ RBCs} \times 100}
\]

\[
\text{RI (CRC) reticulocyte count} \times \frac{\text{Patient's Hct}}{45} \times \frac{1}{1.85}
\]

45 = normal Hct; 1.85 = number of days for reticulocyte to mature

Clinical Implications

1. Increased reticulocyte count (reticulocytosis) means that increased RBC production is occurring as the bone marrow replaces cells lost or prematurely destroyed. Identification of reticulocytosis may lead to the recognition of an otherwise occult disease, such as hidden chronic hemorrhage or unrecognized hemolysis (eg, sickle cell anemia, thalassemia). Increased levels are observed in the following:
   a. Hemolytic anemia
      1. Immune hemolytic anemia
      2. Primary RBC membrane problems
      3. Hemoglobinopathic and sickle cell disease
      4. RBC enzyme deficits
      5. Malaria
   b. After hemorrhage (3 to 4 days)
   c. After treatment of anemias
      1. An increased reticulocyte count may be used as an index of the effectiveness of treatment.
      2. After adequate doses of iron in iron-deficiency anemia, the rise in reticulocytes may exceed 20%.
      3. There is a proportional increase when pernicious anemia is treated by transfusion or vitamin B_{12} therapy.

2. Decreased reticulocyte count means that bone marrow is not producing enough erythrocytes; this occurs in:
   a. Untreated iron-deficiency anemia
   b. Aplastic anemia (a persistent deficiency of reticulocytes suggests a poor prognosis)
   c. Untreated pernicious anemia
   d. Anemia of chronic disease
   e. Radiation therapy
f. Endocrine problems

3. Reticulocyte index implications
   a. <2% indicates hypoproliferative component to anemia
   b. >2%–3% indicates increased RBC production

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. Pretest and posttest care are the same as for the hemogram (see page 47).
   Also, see Chapter 1 guidelines for safe, effective, informed pretest care.

2. Note medications. Some drugs cause aplastic anemia.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately for anemias.

   2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Sedimentation Rate (Sed Rate); Erythrocyte Sedimentation Rate (ESR)

Sedimentation occurs when the erythrocytes clump or aggregate together in a column-like manner (rouleaux formation). These changes are related to alterations in the plasma proteins. Normally, erythrocytes settle slowly because normal RBCs do not form rouleaux.

The ESR is the rate at which erythrocytes settle out of anticoagulated blood in 1 hour. This test is based on the fact that inflammatory and necrotic processes cause an alteration in blood proteins, resulting in aggregation of RBCs, which makes them heavier and more likely to fall rapidly when placed in a special vertical test tube. The faster the settling of cells, the higher the ESR. The ESR should not be used to screen asymptomatic patients for disease. It is most useful for diagnosis of temporal arteritis, rheumatoid arthritis, and polymyalgia rheumatica. The sedimentation rate is not diagnostic of any particular disease but rather is an indication that a disease process is ongoing and must be investigated. It is also useful in monitoring the progression of inflammatory diseases; if the patient is being treated with steroids, the ESR will decrease with clinical improvement.

Reference Values by Westergren's Method

Normal
   Men: 0–15 mm/h (over age 50 years: 0–20 mm/h)
   Women: 0–20 mm/h (over age 50 years: 0–30 mm/h)
   Children: 0–10 mm/h

Procedure

1. Obtain an EDTA-anticoagulated venous sample of 5 mL or 3.8% sodium citrate. Place the specimen in a biohazard bag.

2. Suction the specimen into a graduated sedimentation tube and allow to settle for exactly 1 hour. The amount of settling is the patient's ESR.

Clinical Implications

1. Increased ESR is found in:
   a. All collagen diseases, SLE
   b. Infections, pneumonia, syphilis, tuberculosis
   c. Inflammatory diseases (eg, acute pelvic inflammatory disease)
   d. Carcinoma, lymphoma, neoplasms
   e. Acute heavy-metal poisoning
   f. Cell or tissue destruction, myocardial infarction
   g. Toxemia, pregnancy (third month to 3 weeks' postpartum)
   h. Waldenström's macroglobulinemia, increased serum globulins
   i. Nephritis, nephrosis
   j. Subacute bacterial endocarditis
   k. Anemia—acute or chronic disease
   l. Rheumatoid arthritis, gout, arthritis, polymyalgia rheumatica
   m. Hypothyroidism and hyperthyroidism

2. Normal ESR (no increase) is found in:
   a. Polycythemia vera, erythrocytosis
   b. Sickle cell anemia, Hb C disease
   c. Congestive heart failure
   d. Hypofibrinogenemia (from any cause)
   e. Pyruvate kinase deficiency
   f. Hereditary spherocytosis
   g. Anemia
      1. ESR is normal in iron-deficiency anemia
      2. ESR is abnormal in anemia of chronic disease alone or in combination with iron-deficiency anemia and can be used to differentiate these
   h. Uncomplicated viral disease and infectious mononucleosis—normal
   i. Active renal failure with heart failure—normal
   j. Acute allergy—normal
k. Peptic ulcer—normal

---

**Clinical Alert**

Extreme elevation of the ESR is found with malignant lymphocarcinoma of colon or breast, myeloma, and rheumatoid arthritis.

**Interfering Factors**

1. Allowing the blood sample to stand >24 hours before the test is started causes the ESR to decrease.
2. In refrigerated blood, the ESR is increased. Refrigerated blood should be allowed to return to room temperature before the test is performed.
3. Factors leading to an increased ESR include:
   a. The presence of fibrinogen, globulins, C-reactive protein, high cholesterol
   b. Pregnancy after 12 weeks until about the fourth postpartum week
   c. Young children
   d. Menstruation
   e. Certain drugs (eg, heparin, oral contraceptives; see Appendix J)
   f. Anemia (low Hct)
   g. Macrocytosis
4. The ESR may be very high (up to 60 mm/h) in apparently healthy women aged 70 to 89 years.
5. Factors leading to reduced ESR include:
   a. High blood sugar, high albumin level, high phospholipids
   b. Decreased fibrinogen level in the blood in newborns, hypofibrinogenemia
   c. Certain drugs (eg, steroids, high-dose aspirin; see Appendix J)
   d. High Hb and RBC—polycythemia
   e. High WBC
   f. Abnormal RBCs (eg, sickle cells, spherocytes, microcytosis)

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. Obtain appropriate medication history. Fasting is not necessary, but a fatty meal can cause plasma alterations.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities and diet.
2. Interpret test outcome; counsel and monitor appropriately for rheumatic disorders and inflammatory conditions.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

**TESTS FOR PORPHYRIA**

Porphyrins are chemical intermediates in the synthesis of Hb, myoglobin, and other respiratory pigments called cytochromes. They also form part of the peroxidase and catalase enzymes, which contribute to the efficiency of internal respiration. Iron is chelated within porphyrins to form heme. Heme is then incorporated into proteins to become biologically functional hemoproteins.

Tests of blood, urine, and stool are done to diagnose porphyria, an abnormal accumulation of porphyrins in body fluids. Porphyrias are a group of diseases caused by a deficit in the enzymes involved in porphyrin metabolism and abnormalities in the production of the metalloporphyrin heme. These tests are indicated in persons who have unexplained neurologic manifestations, unexplained abdominal pain, cutaneous blisters, and/or the presence of a relevant family history. Test results may identify clinical conditions associated with abnormal heme production, including anemia and porphyria (abnormal accumulation of the porphyrins) associated with enzyme disorders that may be genetic (hereditary) or acquired (eg, lead poisoning, alcohol). Accumulation of porphyrins occurs in blood plasma, serum, erythrocytes, urine, and feces. A discussion of erythrocyte totals and fractionation of erythrocytes and plasma follows. For details of urine, serum, and stool testing for porphyrrias, see Chapter 3, Chapter 6, and Chapter 4, respectively.

**Erythropoietic Porphyrias; Free Erythrocyte Protoporphyrin (FEP)**

Normally, there is a small amount of excess porphyrin at the completion of heme synthesis. This excess is cell-free erythrocyte protoporphyrin (FEP). The amount of FEP in the erythrocyte is elevated when the iron supply is diminished.

This test is useful in screening RBC disorders such as iron deficiency and lead exposure, especially in children 6 months to 5 years of age. This is the test of choice to diagnose erythropoietic protoporphoria. This test should not be used for screening for lead poisoning in children.

**Reference Values**

*Normal* <100 µg/dL of packed RBCs

**NOTE**

This depends on the method. Check with your laboratory.
Procedure
1. Obtain a 5-mL sample of anticoagulated venous blood. EDTA, or heparin, may be used. Place the specimen in a biohazard bag.
2. Protect the blood sample from light.
3. Wash the cells and then test for porphyrins.
4. Be aware that the Hct must be known for test interpretation.

Clinical Implications
1. Increased FEP is associated with:
   a. Iron-deficiency anemias (elevated before anemia)
   b. Lead poisoning (chronic)
   c. Halogenated solvents and many drugs (see Appendix J)
   d. Anemia of chronic disease
   e. Acquired idiopathic sideroblastic anemia (most cases)
   f. Erythropoietic protoporphyria
2. FEP is normal in:
   a. Thalassemia minor (and therefore can be used to differentiate this from iron deficiency and other disorders of globin synthesis)
   b. Pyridoxine-responsive anemia
   c. Certain forms of sideroblastic anemia due to proximal block to protoporphyrin

Interventions
Pretest Patient Care
1. Explain test purpose and sampling procedure.
2. Note on laboratory slip or computer any medications the patient is taking that cause intermittent porphyria. Discontinue such medications before testing (after checking with physician).
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Resume normal activities and diet.
2. Interpret test outcome and monitor appropriately for porphyria or lead poisoning.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert
The critical value is FEP >300 µg/dL.

Porphyrs; Fractionation of Erythrocytes and of Plasma
The primary porphyrins of erythrocytes are protoporphyrin, uroporphyrin, and coproporphyrin.

Fractionation of erythrocytes is used to differentiate congenital erythropoietic coproporphyria from erythropoietic protoporphyria and to confirm a diagnosis of protoporphyria. This test establishes a specific type of porphyria by naming the specific porphyrin in plasma. In persons with renal failure, plasma fractionation can help to determine whether the porphyria is caused by a deficiency of uroporphyrinogen decarboxylase or by failure of the renal system to excrete porphyrinogens.

Reference Values
Normal The value is reported in micrograms per deciliter (µg/dL). Check with your laboratory for reference values.
1. Erythrocyte porphyrins:
   a. Protoporphyrin: 16–60 µg/dL packed cells or 0.3–1.7 µmol/L
   b. Uroporphyrin: <2 µg/dL or <24 nmol/L
   c. Hepatocarboxylic: <1 µg/dL or <10 µg/L
   d. Hexacarboxylic: <1 µg/dL or <10 µg/L
   e. Pentacarboxylic: <1 µg/dL or <10 µg/L
   f. Coproporphyrin: <1 µg/dL or <15 µg/L
2. Plasma porphyrins: Total porphyrins should not exceed 1.0 µg/dL or 12 nmol/L

Procedure
1. Draw a 5-mL sample of anticoagulated blood. EDTA or heparin can be used as an anticoagulant. Place the specimen in a biohazard bag.
2. Protect the specimen from light.

Clinical Implications
1. Increased erythrocyte porphyrins are associated with primary porphyrias:
   a. Congenital erythropoietic protoporphyria
b. Protoporphyria (autosomal dominant deficiency of heme synthetase)
c. Hereditary porphobilinogen synthase deficiency
d. Intoxication porphyria

2. *Increased plasma porphyrins* are associated with:
   a. Congenital erythropoietic protoporphyria
   b. Coproporphyria
   c. Porphyria cutanea tarda
   d. Paraproteoporphyria
   e. Chronic renal failure porphyria

Interventions

*Pretest Patient Care*

1. Advise patient of test purpose.
2. Note on the requisition any drugs the patient is taking.
3. Before testing, discontinue drugs that are known to cause intermittent porphyria (after checking with physician).
4. See Chapter 1 guidelines for [safe, effective, informed pretest care](#).

*Posttest Patient Aftercare*

1. Resume medications.
2. Interpret test outcome and monitor appropriately for porphyria or lead poisoning.
3. Advise persons diagnosed with porphyria (with cutaneous manifestations) to avoid sun exposure.
4. Advise persons diagnosed with porphyria (with neurologic symptoms) that attacks can be precipitated by infections, various phases of the menstrual cycle, fasting states, and certain drugs. A listing of drugs (not all inclusive) that may precipitate acute attacks follows:
   a. Barbiturates
   b. Chlordiazepoxide
   c. Chloroquine
   d. Chlorpropamide
   e. Dichloralphenazone
   f. Ergot preparations
   g. Estrogens
   h. Ethanol
   i. Glutethimide
   j. Griseofulvin
   k. Hydantoins
   l. Imipramine
   m. Meprobamate
   n. Methsuximide
   o. Methylparoxetin
   p. Sulfonamides
5. Follow Chapter 1 guidelines for [safe, effective, informed posttest care](#).

**Clinical Alert**

1. A blood test for uroporphyrinogen I synthase (also known as erythrocyte porphobilinogen deaminase) can be done to identify persons at risk for acute intermittent porphyria, to detect latent-phase intermittent porphyria, and to confirm the diagnosis during an acute episode.
2. The normal value is 5.3–9.2 nmol/L in women; 3.4–8.5 nmol/L in men. A value of <3.5 nmol/L is diagnostic of acute intermittent porphyria.

**ADDITIONAL TESTS FOR HEMOLYTIC ANEMIA**

Several RBC enzyme and fragility tests can be done to screen, detect, and confirm the cause of chronic hemolytic anemia. Many persons with hemolytic anemia have no clinical signs or symptoms. Abnormal test outcomes are associated with inherited deficiencies, abnormal hemoglobins, and exposure to chemicals and drugs. Definitive test results indicate some type of injury to the RBC, oxidized activity that interferes with normal Hb function, and/or increased RBC fragility.

**Pyruvate Kinase (PK)**

PK deficiency is a genetic disorder characterized by a lowered concentration of adenosine triphosphate in the RBC and consequent membrane defect. The result is a nonspherocytic, chronic hemolytic anemia. PK deficiency is the most common and most important form of hemolytic anemia resulting from a deficiency of glycolytic enzymes in the RBC.

**Reference Values**

*Normal* 2.8–8.8 U/g Hb or 46.7–146.7 nkat/g Hb To convert to U/mL of packed RBCs: U/g Hb × 0.34 = U/mL packed RBCs Check with your reference lab.

**Procedure**

1. Obtain a venous blood sample of at least 5 mL with EDTA or heparin anticoagulant.
2. Refrigerate immediately.
Clinical Implications  PK is increased in:

1. Congenital PK deficiency: recessive, nonspherocytic hemolytic anemia. Patients tolerate anemia well because of increased 2,3-diphosphoglycerate (2,3-DPG).
2. Acquired PK deficiency caused by (level returns to normal after treating underlying disorder):
   a. Myelodysplastic disorders
   b. Acute leukemias
   c. Anemias

Interfering Factors  In congenital PK, intravascular hemolysis increases during pregnancy or following use of oral contraceptives.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. There should be no exercising before tests.
2. Withhold transfusion until after blood samples are drawn (especially with osmotic fragility).
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and monitor appropriately for hemolytic anemia, hypoxia, or polycythemia.
2. Splenectomy is indicated when anemia is severe enough to require transfusions.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

Many prescribed drugs interfere with the normal functioning of hemoglobin in susceptible persons, especially sulfonamides, antipyretics, analgesics, large doses of vitamin K, and nitrofurans.

Erythrocyte Fragility (Osmotic Fragility and Autohemolysis)

Spherocytes of any origin (including conditions other than hereditary spherocytosis) are more susceptible than normal RBCs to hemolysis in dilute (hypotonic) saline and show increased osmotic fragility. Generally, fully expanded cells (spheroidal cells or spherocytes) have increased osmotic fragility, whereas cells with higher surface area-to-volume ratios (eg, thin cells, hypochromic cells, tart cells) have decreased osmotic fragility.

In hereditary spherocytosis, the osmotic fragility test may be normal initially. Therefore, the test is incubated at 37°C for 24 hours, at which time the test is positive for hereditary spherocytosis.

Reference Values

<table>
<thead>
<tr>
<th>Normal Immediate test</th>
<th>Hemolysis begins at 0.5% NaCl</th>
<th>Hemolysis complete at 0.3% NaCl</th>
</tr>
</thead>
<tbody>
<tr>
<td>24-hour incubation</td>
<td>Hemolysis begins at 0.7% NaCl</td>
<td>Hemolysis complete at 0.4% NaCl</td>
</tr>
</tbody>
</table>

Procedure

1. Obtain a 7-mL venous blood sample using heparin as anticoagulant. Place the specimen in a biohazard bag.
2. Expose erythrocytes to varying dilutions of sodium chloride. Read hemolysis on a spectrophotometer (optical density measurement). Perform studies and measure both before and after 24-hour incubation of the RBCs.

Clinical Implications

1. Increased osmotic fragility is found in:
   a. Hemolytic anemia (acquired immune)
   b. Hereditary spherocytosis (stomatocytosis)
   c. Hemolytic disease of the newborn
   d. Malaria
   e. Severe pyruvate kinase deficiency
2. Decreased osmotic fragility occurs in:
   a. Iron-deficiency anemia (macrocytic hypochromic)
   b. Thalassemias
   c. Asplenia (postsplenectomy)
   d. Liver disease (obstructive jaundice)
   e. Reticulocytosis
   f. Hemoglobinopathies, especially Hb C, Hb S

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. There should be no exercising before tests.
2. Withhold transfusion until after blood samples are drawn (especially with osmotic fragility).
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Follow Chapter 1 guidelines for safe, effective, informed posttest care.
2. Be aware that the usual treatment for hereditary spherocytosis is splenectomy, which removes the agent of RBC destruction and prevents complications such as aplastic anemia.

Glucose-6-Phosphate Dehydrogenase (G6PD)

G6PD is a sex-linked disorder. The major variants occur in specific ethnic groups. In a large group of African American men, the incidence of type A G6PD deficiency was found to be 11%. Approximately 20% of African American women are heterozygous. With some variants, there is chronic lifelong hemolysis, but more commonly, the condition is asymptomatic and results only in susceptibility to acute hemolytic episodes, which may be triggered by certain drugs, ingestion of fava beans, or viral or bacterial infection. G6PD hemolysis is associated with formation of Heinz bodies in peripheral RBCs.

The other two most common types are Mediterranean, which is common in Iraqis, Kurds, Sephardic Jews, and Lebanese and less common in Greeks, Italians, Turks, and North Africans, and the MAHIDOL variant, which is common in Southeast Asians (22% males).

Reference Values

**Normal**

**G6PD screen:** G6PD detected  
**Adults:** 8.6–18.6 U/g Hb or 0.14–0.31 nkat/g Hb  
**Children:** 6.4–15.6 U/g Hb or 0.11–0.26 nkat/g Hb  
**Newborns:** have values up to 50% higher than adults  
If done as a screening test, G6PD activity is reported as within normal limits. Different laboratories have different ways of reporting. To convert U/g Hb to U/mL of RBCs: U/g Hb × 0.34 = U/mL of RBCs

Procedure

1. Obtain a blood sample of at least 5 mL, using EDTA or heparin anticoagulant.  
2. Place on ice in a biohazard bag. Perform a G6PD screen first.

Clinical Implications

1. G6PD is decreased in:  
   a. G6PD deficiency (causes hemolytic episodes after exposure to certain drugs and fava beans)  
   b. Congenital nonspherocytic anemia  
   c. Nonimmunologic hemolytic disease of the newborn (Asian and Mediterranean)
2. G6PD is increased in:  
   a. Untreated megaloblastic anemia (pernicious anemia)  
   b. Thrombocytopenia purpura  
   c. Hyperthyroidism  
   d. Viral hepatitis

Interfering Factors

1. Marked reticulocytosis may give a falsely high G6PD.  
2. G6PD may be falsely normal for 6 to 8 weeks after a hemolytic episode, especially in black persons with the type A variant. Retest after the patient recovers from the episode of anemia.

**Clinical Alert**

In G6PD-Mediterranean, G6PD levels are grossly deficient in all RBCs. Patients with this variant commonly experience hemolysis induced by diabetic acidosis, infections, and oxidant drugs and potentially fatal hemolytic crises after ingestion of fava beans.

Interventions

**Pretest Patient Care**

1. Explain test purpose and procedure. There should be no exercising before tests.  
2. Withhold transfusion until after blood samples are drawn (especially with osmotic fragility).  
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Follow Chapter 1 guidelines for safe, effective, informed posttest care.  
2. Be aware that there are certain drugs and chemicals that should be avoided by persons with G6PD.

**Heinz Bodies; Heinz Stain; Glutathione Instability**

Heinz bodies are insoluble intracellular inclusions of Hb attached to RBC membrane. Heinz bodies are uncommon except with G6PD deficiency immediately after hemolysis and in patients with unstable Hb variants.

Oxidative denaturation of the Hb molecule leads to Heinz body formation and is probably the mechanism for the precipitation of unstable Hb. Heinz bodies are usually removed by the spleen; after splenectomy, they increase in the peripheral blood and may appear in >50% of RBCs.

Reference Values

**Normal** Not seen in normal patients
Procedure

1. Obtain a venous blood sample, anticoagulated with heparin or EDTA. Place the specimen in a biohazard bag.
2. Mix cells with a supravital stain and examine microscopically. They stain as pale blue bodies, as opposed to the dark purple RNA in reticulocytes.

Clinical Implications

1. Increased Heinz bodies are found in:
   a. G6PD deficiency, especially after hemolysis
   b. Congenital Heinz body hemolytic anemia
   c. Unstable Hb variants (eg, Hb Zurich, Hb Philly)
   d. Homozygous ß-thalassemia
2. Heinz bodies are found in blood of normal persons who have been poisoned by certain drugs used in treatment protocols (eg, chlorates, phenylhydrazine, primaquine).
3. Heinz bodies are present in some newborns or in splenectomized patients.

Interfering Factors See Appendix J for drugs that affect test outcomes.

Interventions

Pretest Patient Care

1. Explain test purposes and procedures.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes; counsel and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

2,3-Diphosphoglycerate (2,3-DPG)

2,3-DPG assists in transporting oxygen in the RBC. 2,3-DPG increases in response to hypoxia or anemia and decreases in acidosis. Levels are lower in newborns and even lower in premature newborns.

Reference Values

Normal Adults: 10.4–14.2 µmol/g Hb or 3.6–4.8 µmol/mL RBCs Check with your reference lab.

Procedure

1. Obtain a venous blood sample of at least 3 mL, anticoagulated with heparin.
2. Place on ice immediately (2,3-DPG is stable for only 2 hours) and transport to laboratory as soon as possible in a biohazard bag.

Clinical Implications

1. Increased 2,3-DPG occurs in:
   a. Emphysema, cystic fibrosis with pulmonary involvement (conditions of hypoxia)
   b. Cyanotic heart disease
   c. Pulmonary vascular disease
   d. Sickle cell anemia, iron-deficiency anemia
   e. Pyruvate kinase deficiency
   f. Hyperthyroidism
   g. Chronic renal failure
   h. Cirrhosis
2. Decreased 2,3-DPG occurs in:
   a. Polycythemia vera
   b. Respiratory distress syndrome
   c. 2,3-DPG deficiency
   d. Hexokinase deficiency

Interfering Factors

1. High altitude increases 2,3-DPG.
2. Exercise increases 2,3-DPG.

Clinical Alert

If blood with decreased 2,3-DPG is used for transfusion, the Hb may not release O₂ when needed.

Interventions

Pretest Patient Care for Tests for Hemolytic Anemia

1. Explain test purpose and procedure. There should be no exercising before tests.
2. Withhold transfusion until after blood samples are drawn (especially with osmotic fragility).
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare for Tests for Hemolytic Anemia

1. Interpret test results and monitor appropriately for hemolytic anemia, hypoxia, or polycythemia.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

IRON TESTS

Iron (Fe), Total Iron-Binding Capacity (TIBC), and Transferrin Tests

Iron is necessary for the production of Hb. Iron is contained in several components. Transferrin (also called siderophilin), a transport protein largely synthesized by the liver, regulates iron absorption. High levels of transferrin relate to the ability of the body to deal with infections. Total iron-binding capacity (TIBC) correlates with serum transferrin, but the relation is not linear. A serum iron test without a TIBC and transferrin determination has very limited value except in cases of iron poisoning. Transferrin saturation is a better index of iron saturation; it is evaluated as follows:

\[ \text{Transferrin saturation} \% = \frac{\text{Serum iron} \times 100}{\text{TIBC}} \]

The combined results of transferrin, iron, and TIBC tests are helpful in the differential diagnosis of anemia, in assessment of iron-deficiency anemia, and in the evaluation of thalassemia, sideroblastic anemia, and hemochromatosis.

Reference Values

**Normal**

<table>
<thead>
<tr>
<th>Test</th>
<th>Normal Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iron (Adult men)</td>
<td>65–175 µg/dL or 11.6–31.3 µmol/L</td>
</tr>
<tr>
<td>Iron (Adult women)</td>
<td>50–170 µg/dL or 9.0–30.4 µmol/L</td>
</tr>
<tr>
<td>Iron (Children)</td>
<td>50–120 µg/dL or 9.0–21.5 µmol/L</td>
</tr>
<tr>
<td>Iron (Newborns)</td>
<td>100–250 µg/dL or 17.9–44.8 µmol/L</td>
</tr>
<tr>
<td>Total iron-binding capacity (TIBC)</td>
<td>250–450 µg/dL or 44.8–76.1 µmol/L</td>
</tr>
<tr>
<td>Transferrin (Iron) saturation</td>
<td>10%–50%</td>
</tr>
<tr>
<td>Procedure</td>
<td></td>
</tr>
<tr>
<td>1. Obtain a venous blood sample of 10 mL.</td>
<td></td>
</tr>
<tr>
<td>2. Place the specimen in a biohazard bag. Serum is needed for these tests.</td>
<td></td>
</tr>
</tbody>
</table>

Clinical Implications

1. Increased transferrin is observed in:
   a. Iron-deficiency anemia (uncomplicated)
   b. Pregnancy
   c. Estrogen therapy
2. Decreased transferrin is found in:
   a. Microcytic anemia of chronic disease
   b. Protein deficiency or loss from burns or malnutrition
   c. Chronic infection
   d. Acute liver disease
   e. Renal disease (nephrosis)
   f. Genetic deficiency, hereditary atransferrinemia
   g. Iron-overload states (hemochromatosis)
3. Decreased iron occurs in:
   a. Iron-deficiency anemia
   b. Chronic blood loss
   c. Chronic diseases (eg, lupus, rheumatoid arthritis, chronic infections)
   d. Third-trimester pregnancy and progesterone birth control pills
   e. Remission of pernicious anemia
   f. Inadequate absorption of iron
   g. Hemolytic anemia (PNH)
4. Increased iron occurs in:
   a. Hemolytic anemias, especially thalassemia, pernicious anemia in relapse (not hemolytic anemias)
   b. Acute iron poisoning (children)
   c. Iron-overload syndromes
   d. Hemochromatosis, iron overload
   e. Transfusions (multiple), intramuscular iron, inappropriate iron therapy
   f. Acute hepatitis, liver damage
   g. Vitamin B<sub>6</sub> deficiency
   h. Lead poisoning
   i. Acute leukemia
   j. Nephritis
5. Increased TIBC is found in:
   a. Iron deficiency
   b. Pregnancy (late)
   c. Acute and chronic blood loss
   d. Acute hepatitis
6. Decreased TIBC is observed in:
   a. Hypoproteinemia (malnutrition and burns)
b. Hemochromatosis
c. Non–iron-deficiency anemia (infection and chronic disease)
d. Cirrhosis of liver
e. Nephrosis and other renal diseases
f. Thalassemia
g. Hyperthyroidism
7. The iron saturation index is *increased* in:
   a. Hemochromatosis
   b. Increased iron intake
c. Thalassemia
d. Hemosiderosis
e. Acute liver disease
8. The iron saturation index is *decreased* in:
   a. Iron-deficiency anemias
   b. Malignancy (standard and small intestine)
c. Anemia of infection and chronic disease
d. Iron neoplasms

**Interfering Factors**

1. Many drugs affect test outcomes (see Appendix J).
2. Drugs that may cause increased iron include ethanol, estrogens, and oral contraceptives.
3. Drugs that may cause decreased iron include some antibiotics, aspirin, and testosterone.
4. Hemolysis of the blood sample interferes with testing.
5. Iron contamination of glassware used in testing can give high values.
6. Menstruation causes decreased iron; iron is elevated in the premenstrual period.
7. There is a diurnal variation in iron: normal values in the morning, lower in midafternoon, very low in the evening.
8. Serum iron and TIBC may be normal in iron-deficiency anemia if the Hb is >9.0 g/dL (or >90 g/L).

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. Draw fasting blood in the morning, when levels are higher.
3. Draw iron sample before iron therapy is initiated or blood is transfused.
4. If the patient has received a transfusion, delay iron testing for 4 days.
5. Avoid any iron-chelating drug (eg, deferoxamine [Desferal]).
6. Avoid sleep deprivation and extreme stress, which cause lower iron levels.
7. Note on laboratory slip or computer screen whether the patient is taking oral contraceptives or estrogen therapy or is pregnant.
8. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Resume normal activities.
2. Interpret test outcome and monitor appropriately. The combination of low serum iron, high TIBC, and high transferrin levels indicates iron deficiency. Diagnosis of iron deficiency may lead further to detection of adenocarcinoma of the gastrointestinal tract, a point that cannot be overemphasized. A significant minority of patients with megaloblastic anemias (20%–40%) have coexisting iron deficiency. Megaloblastic anemia can interfere with the interpretation of iron studies; repeat iron studies 1 to 3 months after folate or vitamin B₁₂ replacement.
3. Use Chapter 1 guidelines for safe, effective, informed posttest care.

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**Clinical Alert**

1. Critical iron values: intoxicated child, 280–2550 µg/dL or 50–456 µmol/L; fatally poisoned child, >1,800 µg/dL or >322 µmol/L.
2. Symptoms of iron poisoning include abdominal pain, vomiting, bloody diarrhea, cyanosis, and convulsions.

**Ferritin**

Ferritin, a complex of ferric (Fe²⁺) hydroxide and a protein, apoferritin, originates in the reticuloendothelial system. Ferritin reflects the body iron stores and is the most reliable indicator of total-body iron status. A bone marrow examination is the only better test. Bone marrow aspiration may be necessary in some cases, such as low-normal ferritin and low serum iron in the anemia of chronic disease.

The ferritin test is more specific and more sensitive than iron concentration or TIBC for diagnosing iron deficiency. Ferritin decreases before anemia and other changes occur.

**Reference Values**

**Normal Men:** 20–250 ng/mL or 20–250 µg/L With anemia of chronic disease: <100 ng/mL or <100 µg/L In absence of inflammation: <20 ng/mL or <20 µg/L **Women:** 10–120 ng/mL or 10–120 µg/L With anemia of chronic disease: <20 ng/mL or <20 µg/L In absence of inflammation: <10 ng/mL or <10 µg/L **Children:** 7–140 ng/mL or 7–140 µg/L **Newborns:** 25–200 ng/mL or 25–200 µg/L 1 month: 50–200 ng/mL or 50–200 µg/L 2–5 months: 50–200 ng/mL or 50–200 µg/L **Serum**
**TfR-ferritin index:** 1.5 in absence of anemia of chronic disease, 0.8 with anemia of chronic disease

**NOTE**

TfR is the transferrin receptor.

**Procedure**

1. Obtain a venous sample of 6 mL.
2. Place the specimen in a biohazard bag.

**Ferritin, Iron, and Iron Saturation Changes in Anemias**

<table>
<thead>
<tr>
<th>Anemia</th>
<th>Ferritin</th>
<th>Iron</th>
<th>Iron Saturation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemorrhage, acute</td>
<td>N</td>
<td>D</td>
<td>D</td>
</tr>
<tr>
<td>Hemorrhage, chronic</td>
<td>D</td>
<td>D</td>
<td>D</td>
</tr>
<tr>
<td>Iron-deficiency</td>
<td>D</td>
<td>D</td>
<td>D</td>
</tr>
<tr>
<td>Aplastic</td>
<td>N/I</td>
<td>D/I</td>
<td>D/I</td>
</tr>
<tr>
<td>Megaloblastic</td>
<td>D</td>
<td>I</td>
<td>I</td>
</tr>
<tr>
<td>Hemolytic</td>
<td>I</td>
<td>I</td>
<td>I</td>
</tr>
<tr>
<td>Sideroblastic</td>
<td>I</td>
<td>I</td>
<td>I</td>
</tr>
<tr>
<td>Thalassemia, major</td>
<td>I</td>
<td>N/I</td>
<td>N/I</td>
</tr>
<tr>
<td>Thalassemia, minor</td>
<td>I</td>
<td>N/I</td>
<td>N/I</td>
</tr>
<tr>
<td>Bone marrow neoplasia</td>
<td>N/I</td>
<td>I</td>
<td>I</td>
</tr>
<tr>
<td>Uremia, nephrosis, or nephrotic syndrome</td>
<td>N/I</td>
<td>D/I</td>
<td>D</td>
</tr>
<tr>
<td>Liver disease</td>
<td>N/I</td>
<td>N/I</td>
<td>N/I</td>
</tr>
<tr>
<td>Chronic diseases</td>
<td>I</td>
<td>D</td>
<td>D</td>
</tr>
</tbody>
</table>

N, no change; D, decrease; I, increase.

**Clinical Implications**

1. **Decreased ferritin** (<10 ng/mL or <10 µg/L) usually indicates iron-deficiency anemia.
2. **Increased ferritin** (>400 ng/mL or >400 µg/L) occurs in iron excess and in the following:
   a. Iron overload from hemochromatosis or hemosiderosis
   b. Oral or parenteral iron administration
   c. Inflammatory diseases
   d. Acute or chronic liver disease involving alcoholism
   e. Acute myeloblastic or lymphoblastic leukemia
   f. Other malignancies (Hodgkin’s disease, breast carcinoma, malignant lymphoma)
   g. Hyperthyroidism
   h. Hemolytic anemia, megaloblastic anemia, thalassemia, sideroblastic anemia
   i. Renal cell carcinoma, end-stage renal disease

**Interfering Factors**

1. Recently administered radioactive medications cause spurious results.
2. Oral contraceptives and antithyroid therapy interfere with testing (see Appendix J).
3. Hemolyzed blood may cause high results.
4. Increases with age.
5. Higher in red-meat eaters than vegetarians.
6. Ferritin is not of value to evaluate iron stores in alcoholic persons with liver disease.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. Fasting is not necessary.
2. Radioactive medications may not be given for 3 to 4 days before testing.
3. Refrain from alcohol (higher levels of ferritin occur in alcoholism).
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Clinical Alert**

**Critical value:** Iron deficiency: <10 mg/mL or <10 µg/L

**Posttest Patient Aftercare**

1. Resume normal activities.
2. Interpret test results and monitor appropriately for iron-deficiency anemia and ferritin increases. When iron and TIBC tests are used together with ferritin, they can better distinguish between iron-deficiency anemia and the anemia of chronic disease. Explain possible treatment with vitamin B\(_12\) and folic acid.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.
Iron Stain (Stainable Iron in Bone Marrow; Prussian Blue Stain)

In the bone marrow, normoblasts containing iron granules (stainable) are known as sideroblasts. Erythrocytes (RBCs) that contain stainable iron are called siderocytes. Normally, about 33% of the normoblasts are sideroblasts. Other storage iron is readily identifiable in monophages in bone marrow particles on the marrow slides.

The bone marrow iron stain is the gold standard of iron deficiency: the presence of iron rules out iron deficiency. Marrow iron disappears before peripheral blood changes occur in iron-deficiency anemia. Only patients with decreased marrow iron are likely to benefit from iron therapy.

Reference Values

**Normal** Bone marrow: 33% sideroblasts present Peripheral blood: no siderocytes present

Procedure

1. Make bone marrow slides (bone marrow biopsy material can be used), stain, and examine under the microscope for the presence of iron.
2. Remember that this test may also be done on peripheral blood for the detection of sideroblastic anemias.

Clinical Implications

1. Bone marrow iron is decreased in:
   a. Iron deficiency from all causes of chronic bleeding, hemorrhage, malignancy
   b. Polycythemia vera
   c. Pernicious anemia (early phase of therapy)
   d. Collagen diseases (eg, rheumatoid arthritis, SLE)
   e. Infiltration of marrow by malignant lymphomas, carcinoma
   f. Chronic infection
   g. Myeloproliferative diseases
   h. Uremia
2. Bone marrow iron is increased in:
   a. Hemochromatosis (primary and secondary)
   b. Anemia, especially thalassemia major and minor, PHN, and other hemolytic anemias
   c. Megaloblastic anemia in relapse
   d. Chronic infections
   e. Chronic pancreatic insufficiency

Interfering Factors

Ingestion of iron dextran will bring values to normal despite other evidence of iron-deficiency anemia.

Interventions

Pretest Patient Care

1. See preparation guidelines for bone marrow aspiration (see page 45).
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. See aftercare guidelines for bone marrow aspiration (see page 46).
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

TESTS FOR HEMOGLOBIN DISORDERS

Hemoglobin Electrophoresis

Normal and abnormal Hb can be detected by electrophoresis, which matches hemolyzed RBC material against standard bands for the various Hb types known. The most common forms of normal adult Hb are Hb A₁, Hb A₂, and Hb F (fetal Hb). Of the various types of abnormal Hb (hemoglobinopathies), the best known are Hb S (responsible for sickle cell anemia) and Hb C (results in a mild hemolytic anemia). The most common abnormality is a significant increase in Hb A₂, which is diagnostic of the thalassemias, especially β-thalassemia trait. More than 350 variants of Hb have been recognized and identified.

Clinical Alert

The results may be questionable if a blood transfusion has been given in the months preceding testing.

Reference Values

**Normal** Hb A₁: 90.5%–98.5% or 0.905–0.985 mass fraction Hb A₂: 1.5%–3.5% or 0.015–0.035 mass fraction Hb F: 0.0%–1.0% or 0–0.01 mass fraction

Fetal Hemoglobin (Hemoglobin F; Alkali-Resistant Hemoglobin)

Hb F is a normal Hb manufactured in the RBCs of the fetus and infant; it makes up 50% to 90% of the Hb in the newborn. The remaining portion of the Hb in the newborn is made up of Hb A₁ and Hb A₂, the adult types.
Under normal conditions, the manufacture of Hb F is replaced by the manufacture of adult Hb types during the first year of life. But if Hb F persists and constitutes more than 5% of the Hb after 6 months of age, an abnormality should be expected.

Determination of Hb F is used to evaluate thalassemia (an inherited abnormality in the manufacture of Hb), hemolytic anemias, hereditary persistence of fetal Hb, and other hemoglobinopathies.

**Reference Values**

**Normal**  
Adults: 0%–2% or 0–0.02 mass fraction Hb F  
Newborns: 60%–90% or 0.60–0.90 mass fraction Hb F  
By 6 months of age: 2% or 0.02 mass fraction Hb F

**Procedure**

1. Use a 5-mL venous blood EDTA-anticoagulated sample for Hb electrophoresis.
2. Remember that a blood smear stain may also be done to identify cells containing Hb F (Kleihauer-Betke stain).

**Clinical Implications** *Increased* Hb F is found in:

1. Thalassemias (major and minor)  
2. Hereditary familial fetal hemoglobinemia (persistence of Hb F)  
3. Hyperthyroidism  
4. Sickle cell disease  
5. Hb H disease  
6. Anemia, as a compensatory mechanism (pernicious anemia, PNH, sideroblastic anemia)  
7. Leakage of fetal blood into the maternal bloodstream  
8. Aplastic anemia (acquired)  
9. Juvenile myeloid leukemia with absence of Philadelphia chromosome  
10. Myeloproliferative disorders, multiple myeloma, lymphoma

**Clinical Alert**

In *thalassemia minor*, continued production of Hb F may occur on a minor scale (5%–10%), and the patient usually lives. In *thalassemia major*, the values may reach 40%–90%. This continued production of Hb F leads to severe anemia, and death usually ensues.

**Interfering Factors**

1. If analysis of the specimen is delayed for more than 2 to 3 hours, the level of Hb F may be falsely increased.
2. Infants small for gestational age or with chronic intrauterine anoxia have persistently elevated Hb F.
3. Hb F is increased during anticonvulsant drug therapy.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.  
2. Ensure that the test is done before transfusion.  
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome; counsel and monitor appropriately for thalassemia and anemia.  
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Hemoglobin A\textsubscript{2} (Hb A\textsubscript{2})**

Hb A\textsubscript{2} levels have special application to the diagnosis of β-thalassemia trait, which may be present even though the peripheral blood smear is normal. The microcytosis and other morphologic changes of β-thalassemia trait must be differentiated from iron deficiency. Low MCV may be present in most patients with β-thalassemia trait, but it does not differentiate iron-deficient patients.

This measurement is used in the investigation of hemolytic anemias for hemoglobinopathies, especially thalassemia and β-thalassemia.

**Reference Values**

**Normal**  
Adult: 1.5%–3.5% or 0.015–0.035 mass fraction Newborns: 0%–1.8% or 0–0.018 mass fraction

**Procedure**

1. Draw a 5-mL venous sample of blood with EDTA anticoagulant.  
2. Perform electrophoresis.

**Clinical Implications**

1. *Increased* Hb A\textsubscript{2} occurs in:
a. β-Thalassemia major (3%–11%)
b. Thalassemia minor (3.5%–7.5%)
c. Thalassemia intermedia (6%–8%)
d. Hb A/S (sickle cell trait) (15%–45%)
e. Hb S/S (sickle cell disease) (2%–6%)
f. S-ß-thalassemia (3.0%–8.5%)
g. Megaloblastic anemia
h. Hyperthyroidism
i. Vitamin B₁₂ or folate deficiency

2. Decreased Hb A₂ occurs in:
   a. Untreated iron-deficiency anemia
   b. Sideroblastic anemia
   c. Hb H disease
   d. Erythroleukemia

Interfering Factors

1. Blood transfusions before electrophoresis will interfere with results.
2. High levels of Hb F usually are accompanied by low levels of A₂.
3. Hb C, Hb O, Hb E interfere with the electrophoric migration of A₂.
4. If a patient with β-thalassemia also has iron-deficiency anemia, the A₂ may be normal; therefore, retesting may be needed after iron therapy.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Provide genetic counseling.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome; counsel and monitor appropriately.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Hemoglobin S (Sickle Cell Test; Sickledex)

Sickle cell disease is a term for a group of hereditary blood disorders. Sickle cell anemia is caused by an abnormality of Hb, the red protein in red blood cells that carries oxygen from the lungs to the tissues. People with sickle cell disease make an abnormal Hb, hemoglobin S (Hb S). The red blood cells of a person with sickle cell disease do not last as long as "normal" red blood cells. This result is chronic anemia. Also, these red blood cells lose their normal disk shape. They become rigid and deformed and take on a "sickle" or crescent shape. These oddly shaped cells are not flexible enough to squeeze through small blood vessels. This may result in blood vessels being blocked. The areas of the body served by those blood vessels will then be deprived of their blood circulation, damaging tissues and organs. This homozygous state of Hb S disease is associated with considerable morbidity and mortality. The heterozygous state presents little mortality. This blood measurement is routinely done as a screening test for sickle cell anemia or trait and to confirm these disorders. This test detects Hb S, an inherited, recessive gene. An examination is made of erythrocytes for the sickle-shaped forms characteristic of sickle cell anemia or trait. This is done by removing oxygen from the erythrocyte. In erythrocytes with normal Hb, the shape is retained, but erythrocytes containing Hb S assume a sickle shape. However, the distinction between sickle cell trait and sickle cell disease is done by electrophoresis, which identifies an Hb pattern.

Reference Values

Normal Adult: None present

Procedure

1. Obtain a venous blood sample of 5 mL with EDTA. Place the specimen in a biohazard bag.
2. Perform the Sickledex test or Hb electrophoresis. Electrophoresis is more accurate and should be done in all positive Sickledex screens.

Clinical Implications A positive test (Hb S present) means that great numbers of erythrocytes have assumed the typical sickle cell (crescent) shape. Positive tests are 99% accurate.

1. Sickle cell trait
   a. Definite confirmation of sickle cell trait by Hb electrophoresis reveals the following heterozygous (A/S) pattern: Hb S, 20%–40%; Hb A₁, 60%–80%; Hb F, small amount. This means that the patient has inherited a normal Hb gene from one parent and an Hb S gene from the other (heterozygous pattern). This patient does not have any clinical manifestations of the disease, but some of the children of this patient may inherit the disease if the patient's mate also has the recessive gene pattern.
   b. The diagnosis of sickle cell trait does not affect longevity and is not accompanied by signs and symptoms of sickle cell anemia. A/S occurs in 8.5% of African Americans.
   c. Sickle cell trait can lead to renal papillary necrosis, hematuria, increased risk for pulmonary embolus, and anterior segment ischemia.
2. Sickle cell anemia (Hb S disease)
Definite confirmation of sickle cell anemia by Hb electrophoresis reveals the following homozygous (S/S) pattern: Hb S, 80%–100%; Hb F, most of the rest, Hb A\(_1\), 0% (absent).

This means that an abnormal Hb S gene has been inherited from both parents (homozygous pattern). Such a patient has all the clinical manifestations of the disease.

3. Hb C—Harlem (rare)
4. Hb C—Georgetown
5. Hb S in combination with other disorders, such as \(\beta\)-thalassemia or Hb S-C

Interfering Factors

1. False-negative results occur in:
   a. Infants younger than 3 months of age (maximum amounts reached by 6 months)
   b. Coexisting thalassemias or iron deficiency
   c. The solubility test is unreliable in pernicious anemia and polycythemia
2. False-positive results occur up to 4 months after transfusion with RBCs having sickle cell trait.
3. Hb D and Hb G migrate to same place as Hb F in electrophoresis.

Clinical Alert

A positive Sickledex test must be confirmed by electrophoresis.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Provide genetic counseling.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome; counsel and monitor appropriately.
2. A person with sickle cell disease should avoid situations in which hypoxia may occur, such as very strenuous exercise, traveling to high-altitude regions, or traveling in an unpressurized aircraft.
3. Because of the hypoxia created by general anesthetics and a state of shock, surgical and maternity patients with sickle cell disease need very close observation.
4. See Chapter 1 guidelines for safe, effective, informed posttest care.

Methemoglobin (Hemoglobin M)

Methemoglobin is formed when the iron in the heme portion of deoxygenated Hb is oxidized to a ferric form rather than a ferrous form. In the ferric form, oxygen and iron cannot combine. The formation of methemoglobin is a normal process and is kept within bounds by the reduction of methemoglobin to Hb. Methemoglobin causes a shift to the left of the oxyhemoglobin dissociation curve. When a high concentration of methemoglobin is produced in the RBCs, it reduces their capacity to combine with oxygen; anoxia and cyanosis result.

This test is used to diagnose hereditary or acquired methemoglobinemia in patients with symptoms of anoxia or cyanosis and no evidence of cardiovascular or pulmonary disease. Hb M is an inherited disorder of the Hb that produces cyanosis.

Methemoglobinemia is most commonly encountered as an acquired state as a result of medications such as phenacetin, sulfonamides, or ingestion of nitrates.

Reference Values

Normal 0.4%—1.5% or 0.004—0.015 of total Hb A value of >40% or >0.40 is a critical value.

Procedure

1. Obtain a venous or arterial blood sample, anticoagulated with sodium fluoride.
2. Place on ice immediately and transport to laboratory in a biohazard bag. Methemoglobin is very unstable and must be tested within 8 hours.

Clinical Implications

1. Hereditary methemoglobinemia (uncommon) is associated with:
   a. A hemoglobinopathy, Hb M (40% [or 0.40] of the total Hb)
   b. Deficiency of methemoglobin reductase (autosomal recessive)
   c. Glutathione deficiency (dominant mode of transmission)
2. Acquired methemoglobinemia is associated with:
   a. Black-water fever
   b. Paroxysmal hemoglobinuria
   c. Clostridial infection
3. Toxic effect of drugs or chemicals (most common cause):
   a. Analgesics, phenacetin
   b. Sulfonamide derivatives—sulfonamide S
   c. Nitrates and nitrites; nitroglycerin
d. Antimalarials
e. Isoniazid
f. Quinones
g. Potassium chloride
h. Benzocaine, lidocaine
i. Dapsone (most common drug causing methemoglobinemia)

**Clinical Alert**

Critical (panic) values:

1. HbM of 30% (or 0.30) results in headaches, cyanosis
2. HbM of 70% (or 0.70) is usually fatal

**Interfering Factors**

1. Consumption of sausage, processed meats, or other foods rich in nitrates and nitrates
2. Absorption of silver nitrate used to treat extensive burns
3. Excessive intake of Bromo-Seltzer is a common cause of methemoglobinemia. (The patient appears cyanotic but otherwise feels well.)
4. Smoking
5. Use of bismuth preparations for diarrhea (see Appendix J)

**Interventions**

**Pretest Patient Care**

1. Advise patient of purpose of test. Assess for history of Bromo-Seltzer or toxic drugs or chemicals.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome; counsel for cause of cyanosis and monitor appropriately for anoxia.
2. Be aware that treatment includes intravenous methylene blue and oral ascorbic acid.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Because fetal hemoglobin is more easily converted to methemoglobin than to adult hemoglobin, infants are more susceptible than adults to methemoglobinemia, which may be caused by drinking well water containing nitrates. Bismuth preparations for diarrhea may also be reduced to nitrates by bowel action.

**Sulfhemoglobin**

Sulfhemoglobin is an abnormal Hb pigment produced by the combination of inorganic sulfides with Hb. Sulfhemoglobinemia manifests as a cyanosis. Sulfhemoglobinemia often accompanies drug-induced methemoglobinemia.

This test is indicated in persons with cyanosis. Sulfhemoglobinemia may occur in association with the administration of various drugs and toxins. The symptoms are few, but cyanosis is intense even though the concentration of sulfhemoglobin seldom exceeds 10%.

**Reference Values**

**Normal** None present or 0%–1.0% or 0–0.01 of total Hb

**Procedure**

1. Draw a 5-mL venous blood sample, anticoagulated with EDTA or heparin.
2. Place the specimen in a biohazard bag. Sulfhemoglobin is stable.

**Clinical Implications**

1. Sulfhemoglobin is observed in patients who take oxidant drugs such as phenacetin, Bromo-Seltzer, sulfonamides, and acetanilid. (See Appendix J)
2. Sulfhemoglobin is formed rarely without exposure to drugs or toxins, as in chronic constipation and purging.
3. Sulfhemoglobin can be due to exposure to trinitrotoluene or zinc ethylene bisdithiocarbamate (fungicide).

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. Assess for exposure to drugs and toxins.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
Carboxyhemoglobin; Carbon Monoxide (CO)

Carboxyhemoglobin is formed when Hb is exposed to carbon monoxide (CO). The affinity of Hb for CO is 240 times greater than for oxygen. CO poisoning causes anoxia because the carboxyhemoglobin formed does not permit Hb to combine with oxygen.

This test is done to detect CO poisoning. Because carboxyhemoglobin is not capable of transporting oxygen, hypoxia results, causing headache, nausea, vomiting, vertigo, collapse, or convulsions. Death may result from anoxia and irreversible tissue changes. Carboxyhemoglobin produces a cherry-red or violet color of the blood and skin, but this may not be present in chronic exposure. The most common causes of CO toxicity are automobile exhaust fumes, coal gas, water gas, and smoke inhalation from fires. Smoking is a minor cause.

Reference Values

Normal Nonsmokers: <2.0% of total Hb or <0.02 fraction of Hb saturation Heavy smokers: 6.0%–8.0% or 0.06–0.08 fraction of Hb saturation Light smokers: 4.0%–5.0% or 0.04–0.05 fraction of Hb saturation Newborns: 10%–12% or 0.10–0.12 fraction of Hb saturation

Procedure

1. Draw a heparinized or EDTA venous blood sample of 5 mL heparin or EDTA and put on ice.
2. Keep sample tightly capped and transport to laboratory immediately in a biohazard bag.

Clinical Implications

1. Carboxyhemoglobin is increased in:
   a. CO poisoning from many sources, including smoking, exhaust fumes, fires
   b. Hemolytic disease
   c. Blood in intestines
   d. Newborns, because of fetal hemoglobin breakdown that yields endogenous CO
2. A direct correlation has been found between CO and symptoms of heart disease, angina, and myocardial infarction.

Interventions

Pretest Patient Care

1. Advise patient of purpose of test.
2. Draw blood sample before oxygen therapy has started.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel for cause of headache, dizziness, vomiting, convulsions, or coma.
2. Be aware that treatment consists of removal of the patient from the source of CO.
3. Initiate oxygen therapy either by supplemental oxygen at atmospheric pressure or by hyperbaric oxygen.
4. See Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

1. With values of 10%–20% (0.10–0.20), the patient may be asymptomatic.
2. With 20%–30% (0.20–0.30), headache, nausea, vomiting, and loss of judgment occur.
3. With 30%–40% (0.30–0.40), tachycardia, hyperpnea, hypotension, and confusion occur.
4. With 50%–60% (0.50–0.60), there is loss of consciousness.
5. Values >60% (>0.60) cause convulsions, respiratory arrest, and death.

Myoglobin (Mb)

Myoglobin (Mb) is the oxygen-binding protein of striated muscle. It resembles Hb but is unable to release oxygen except at extremely low tension. Injury to skeletal muscle results in release of myoglobin. It is not specific to myocardial muscle. Myoglobin is not tightly bound to protein and is rapidly excreted in the urine.

The myoglobin test is used as an early marker of muscle damage in myocardial infarction and to detect injury damage or necrosis to skeletal muscle. Serum myoglobin is found earlier than creatine kinase (CK) enzymes in acute myocardial infarction.

Reference Values

Normal 5–70 ng/mL or 5–70 µg/L

Procedure

1. Draw a venous blood sample of at least 5 mL; use serum. Lipemic or grossly hemolyzed specimens are not
acceptable.

2. Remember that two or three samples taken 1–2 hours apart give optimal results in detecting myocardial infarction.

Clinical Implications

1. *Increased myoglobin values* are associated with:
   a. Myocardial infarction (elevates 1 to 3 hours after pain onset, earlier than creatine kinase). Amount of myoglobin correlates with size of infarct.
   b. Angina without infarction
   c. Other muscle injury (trauma, exercise, open heart surgery, intramuscular injections)
   d. Polymyositis and progressive muscular dystrophy
   e. Myositis
   f. Rhabdomyolysis
   g. Inflammatory myopathy (eg, SLE)
   h. Toxin exposure: narcotics, Malayan sea snake toxin
   i. Malignant hyperthermia
   j. Renal failure
   k. Electric shock
   l. Tonic-clonic seizures

2. *Decreased myoglobin values* are found in:
   a. Circulating antibodies to myoglobin (many patients with polymyositis)
   b. Rheumatoid arthritis
   c. Myasthenia gravis

Interfering Factors

1. See Appendix J for drugs that affect test outcomes.
2. Cocaine use elevates myoglobin.
3. Decreased elimination due to kidney insufficiency causes increase of serum levels.

Interventions

**Pretest Patient Care**

1. Advise patient of test purpose.
2. Have patient avoid radioisotopes until after blood is drawn.
3. Avoid vigorous exercise before the test because it may elevate myoglobin.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Resume normal activities.
2. Interpret test outcomes; counsel and monitor appropriately for myocardial infarction, muscle inflammation, and metabolic stress.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Clinical Alert**

Myoglobin is currently the earliest biologic marker of myocardial necrosis. It appears in the peripheral blood 2 to 3 hours after pain onset and reaches peak levels at 6 to 9 hours. Myoglobin is a sensitive indicator of acute myocardial infarction but is not specific for cardiac muscle.

Haptoglobin (Hp)

Haptoglobin (Hp) is a transport glycoprotein synthesized solely in the liver. It is a carrier for free Hb in plasma; its primary physiologic function is the preservation of iron. Haptoglobin binds Hb and carries it to the reticuloendothelial system.

A decrease in Hp (with normal liver function) is most likely to occur with increased consumption of Hp due to intravascular hemolysis. The concentration of Hp is inversely related to the degree of hemolysis and to the duration of hemolytic episode.

**Reference Values**

- *Normal*
  - Newborns: 5–48 mg/dL or 50–480 mg/L (may be absent at birth)
  - Children: reach adult levels by 1 year
  - Adults: 40–200 mg/dL or 0.4–2.0 g/L

**Procedure**

1. Obtain a venous blood sample of at least 2 mL. Place the specimen in a biohazard bag.
2. Measure the serum for Hp by a radial immunodiffusion method. A single determination is of limited value.

**Clinical Implications**

1. *Hp is decreased in acquired disorders* such as:
   a. Intravascular hemolysis from any cause
   b. Autoimmune hemolytic anemia
c. Other hemoglobinemias caused by intravascular hemorrhages, especially artificial heart valves, and acute bacterial endocarditis
d. Transfusion reactions
e. Erythroleukemias, fetalis
f. Malignant infestation
g. PNH
h. Hematoma, tissue hemorrhage
  i. Thrombotic thrombocytopenic purpura
  j. Drug-induced hemolytic anemia (methylprednisolone)
k. Acute or chronic liver disease

2. Hp is decreased in some inherited disorders, such as:
   a. Sickle cell disease
   b. G6PD and pyruvate kinase deficiency
   c. Hereditary spherocytosis
   d. Thalassemia and megaloblastic anemias
   e. Congenital absence is observed in 1% of black and Asian populations

3. Hp is increased in:
   a. Infection and inflammation (acute or chronic)
   b. Neoplasias, lymphomas (advanced)
   c. Biliary obstruction
   d. Acute rheumatic disease and other collagen diseases
   e. Tissue destruction

Interfering Factors

1. Estrogen and oral contraceptives lower Hp.
2. Steroid therapy raises Hp.
3. Androgens increase Hp.

Clinical Alert
Normal Hp results measured during inflammatory episodes or during steroid treatment do not rule out hemolysis.

Interventions

Pretest Patient Care

1. Advise patient of test purpose.
2. Avoid use of oral contraceptives and androgens before blood is drawn. (Check with physician.)
3. Avoid exercise before test.
4. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume normal activities and medications.
2. Interpret test results. Repeat testing may be necessary. Monitor appropriately for abnormal bleeding.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Bart’s Hemoglobin

Bart’s Hb is an unstable Hb with high oxygen affinity. When there is complete absence of production of the chain of Hb and deletion of all four globin genes, the disorder is known as Bart’s hydrops fetalis. Both parents of the affected infant have heterozygous thalassemia; they are almost all Southeast Asians. Affected infants are either stillborn or die shortly after birth.

This test determines the percentage of the abnormal Bart’s Hb in cord blood and identifies a-thalassemia hemoglobinopathies.

Reference Values
Normal
Adults: None
Children: None
Newborns: <0.5% or <0.005 mass fraction of total Hb

Procedure

1. Obtain a sample of cord blood, and perform Hb electrophoresis.
2. Be aware that venous blood anticoagulated with EDTA or heparin can be used.

Clinical Implications
Increased levels are associated with:

1. Homozygous a-thalassemia (hydrops fetalis syndrome, which causes stillbirth)
2. Hb H disease
3. a-Thalassemia minor

Interventions

Pretest Patient Care
1. Explain test purpose and procedure to parents.
2. Be aware that obstetric complications may lead to significant morbidity and mortality for the mothers of these infants.
3. Provide genetic counseling in a sensitive manner.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel parents.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Paroxysmal Nocturnal Hemoglobinuria (PNH) Test; Acid Hemolysis Test; Ham’s Test

PNH was first described by a patient who noted hemoglobinuria after sleep. In many patients, the hemolysis is irregular or occult. PNH is a hemolytic anemia in which there is also production of defective platelets and granulocytes. The diagnostic feature of PNH is an increased sensitivity of the erythrocytes to complement-mediated lysis. Although patients with PNH can present with hemoglobinuria or a hemolytic anemia, they may also present with iron deficiency (because of urinary loss of blood), bleeding secondary to thrombocytopenia, thrombosis, renal abnormalities, or neurologic abnormalities.

These tests are carried out to make a definitive diagnosis of PNH. The basis of these tests is that the cells peculiar to PNH have membrane defects, making them extrasensitive to complement in the plasma. Cells from patients with PNH undergo marked hemolysis after 15 minutes in the laboratory test. The tests are performed for patients who have hemoglobinuria, bone marrow aplasia (hypoplasia), or undiagnosed hemolytic anemia; they may be useful in the evaluation of patients with unexplained thrombosis or acute leukemia.

Reference Values

Normal: Negative or <1% hemolysis

Procedure

1. Obtain a venous blood sample of 5 mL anticoagulated with EDTA. Place the specimen in a biohazard bag.
2. Mix the patient’s RBCs with normal serum and also with the patient’s own serum, acidify, incubate at 37°C, and examine for hemolysis. Normally, there should be no lysis of the RBCs in this test (also called Ham’s test).
3. Be aware that a separate test called the sugar water test or sucrose hemolysis test may also be done at this time.

Clinical Implications

A positive test (hemolysis) is found in:

1. PNH: a positive test (10%–50% lysis) is needed for diagnosis. The sucrose hemolysis test is also positive in PNH.
2. Hereditary erythroblastic multinuclearity associated with a positive acidified serum test (HEMPS): the sucrose hemolysis test is negative.

Interfering Factors

1. False-positive results may be obtained with the following:
   a. Blood containing large numbers of spherocytes (hereditary or acquired)
   b. Dyserythropoietic anemia
   c. Specimen >8 hours old, specimen hemolyzed
   d. Aplastic anemia
   e. Leukemia and myeloproliferative syndromes
2. These conditions can be distinguished from PNH by the fact that hemolysis occurs in both acidified serum and complement. In PNH, hemolysis occurs only in complement (complement dependent).

Interventions

Pretest Patient Care

1. Explain test purpose.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results; counsel and monitor appropriately for anemia.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

OTHER BLOOD TESTS FOR ANEMIA

Vitamin B₁₂ (VB₁₂)

Vitamin B₁₂ (VB₁₂), also known as the antipernicious anemia factor, is necessary for the production of RBCs. It is obtained only from ingestion of animal protein and requires an intrinsic factor for absorption. Both VB₁₂ and folic acid depend on a normally functioning intestinal mucosa for their absorption and are important for the production of red blood cells. Levels of VB₁₂ and folate are usually tested in conjunction with one another because the diagnosis of macrocytic
anemia requires measurement of both.

This determination is used in the differential diagnosis of anemia and conditions marked by high turnover of myeloid cells, as in the leukemias. When binding capacity is measured, it is the unsaturated fraction that is determined. The measurement of unsaturated VB\textsubscript{12}-binding capacity (UBBC) is valuable in distinguishing between untreated polycythemia vera and other conditions in which there is an elevated Hct.

**Reference Values**

**Normal**
- Adults: 200–835 pg/mL or 148–616 pmol/L
- Newborns: 160–1300 pg/mL or 118–959 pmol/L
- UBBC: 600–1400 pg/mL or 443–1033 pmol/L

**Procedure**

1. Obtain a fasting venous blood sample of at least 5 mL.
2. Obtain the specimen before an injection of VB\textsubscript{12} is administered and before a Schilling test is done. Place the specimen in a biohazard bag.

**Clinical Implications**

1. *Decreased VB\textsubscript{12} (<100 pg/mL or <74 pmol/L)* is associated with:
   a. Pernicious anemia (megaloblastic anemia)
   b. Malabsorption syndromes and inflammatory bowel disease
   c. Fish tapeworm infestation
   d. Primary hypothyroidism
   e. Loss of gastric mucosa, as in gastrectomy and resection
   f. Zollinger-Ellison syndrome
   g. Blind loop syndromes (bacterial overgrowth)
   h. Vegetarian diets (dietary insufficiency)
   i. Folic acid deficiency
   j. Iron deficiency may be present in some patients (eg, gastrectomy)
2. *Increased VB\textsubscript{12} (>700 pg/mL or >517 pmol/L)* is associated with:
   a. Chronic granulocytic leukemia, lymphatic and monocytic leukemia
   b. Chronic renal failure
   c. Liver disease (hepatitis, cirrhosis)
   d. Some cases of cancer, especially with liver metastasis
   e. Polycythemia vera
   f. Congestive heart failure
   g. Diabetes
   h. Obesity
   i. COPD
3. *Increased UBBC* is found in:
   a. Sixty percent of cases of polycythemia vera. (This test is normal in secondary relative polycythemia, aiding in the differential diagnosis of these two states.)
   b. Reactive leukocytosis (leukemoid reaction)
   c. Chronic myelogenous leukemia

**Interfering Factors** The following result in increased VB\textsubscript{12} values:

1. Pregnancy
2. Blood transfusion
3. Aged persons
4. High vitamin C and A doses
5. Smoking
6. Drugs capable of interfering with VB\textsubscript{12} absorption (see Appendix J)

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. Alert patient that overnight fasting from food is necessary. Water is permitted.
3. Withhold VB\textsubscript{12} injection before the blood is drawn.
4. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Resume normal activities and diet.
2. Interpret test results; counsel and monitor appropriately for anemia, leukemia, or polycythemia.
3. See Chapter 1 guidelines for safe, effective, informed posttest care. See Appendix F for more information on vitamin testing.

**Clinical Alert**

1. Persons who have recently received therapeutic or diagnostic doses of radionuclides will have unreliable results.
2. See Appendix F for more information on nutritional status of vitamin B\textsubscript{12}.
The Schilling test is used to confirm pernicious anemia and to determine whether vitamin B$_{12}$ deficiency is caused by malabsorption.

Folic Acid (Folate)

Folic acid is needed for normal RBC and WBC function and for the production of cellular genes. Folic acid is a more potent growth promoter than VB$_{12}$, although both depend on the normal functioning of intestinal mucosa for their absorption. Folic acid, like VB$_{12}$, is required for DNA production. Folic acid is formed by bacteria in the intestines, is stored in the liver, and is present in eggs, milk, leafy vegetables, yeast, liver, fruits, and other elements of a well-balanced diet.

This test is indicated for the differential diagnosis of megaloblastic anemia and in the investigation of folic acid deficiency, iron deficiency, and hypersegmental granulocytosis. Measurement of both serum and RBC folate levels constitutes a reliable means of determining the existence of folate deficiency. The finding of low serum folate means that the patient’s recent diet was subnormal in folate content, that the patient’s recent absorption of folate was subnormal, or both. Low RBC folate can mean either that there is tissue folate depletion owing to folate deficiency requiring folate therapy or, alternatively, that the patient has primary VB$_{12}$ deficiency that is blocking the ability of cells to take up folate. Serum levels are commonly high in patients with VB$_{12}$ deficiency because this vitamin is needed to allow incorporation of folate into tissue cells. For thoroughness, the serum VB$_{12}$ should also be determined, because more than 50% of all patients with significant megaloblastic anemia have VB$_{12}$ deficiency rather than folate deficiency.

**Reference Values**

**Normal**
- Adults: 2–20 ng/mL (serum) or 4.5–45.3 nmol/L
- Children: 5–21 ng/mL (serum) or 11.3–47.6 nmol/L
- Infants: 14–51 ng/mL or 31.7–115.5 nmol/L

Red blood cell folate:
- Adults: 140–628 ng/mL or 317–1422 nmol/L
- Children: >160 ng/mL or >362 nmol/L

**Procedure**

1. Obtain a fasting venous sample of 10 mL. Protect the sample from light. Place the specimen in a biohazard bag.
2. If RBC folate is ordered, draw 5 mL of venous blood with EDTA anticoagulant. An Hct determination is also required.

**Clinical Implications**

1. **Decreased** folic acid levels are associated with:
   a. Inadequate intake owing to alcoholism, chronic disease, malnutrition, diet devoid of fresh vegetables, or anorexia
   b. Malabsorption of folic acid (eg, small bowel disease)
   c. Excessive use of folic acid by the body (eg, pregnancy, hypothyroidism)
   d. Megaloblastic (macrocytic) anemia caused by VB$_{12}$ deficiency
   e. Hemolytic anemia (sickle cell, phenocytosis, PNH)
   f. Liver disease associated with cirrhosis, alcoholism, hepatoma
   g. Adult celiac disease, sprue
   h. Vitamin B$_{6}$ deficiency
   i. Carcinomas (mainly metastatic), acute leukemia, myelofibrosis
   j. Crohn's disease, ulcerative colitis
   k. Infantile hyperthyroidism
   l. Intestinal resection, jejunal bypass procedure
   m. Drugs that are folic antagonists (interfere with nucleic acid synthesis):
      1. Anticonvulsants (phenytoin)
      2. Aminopterin and methotrexate
      3. Antimalarials
      4. Alcohol (ethanol)
      5. Oral contraceptives
      6. Heavy usage of antacids

2. **Increased** folic acid levels are associated with:
   a. Blind loop syndrome
   b. Vegetarian diet
   c. Pernicious anemia, VB$_{12}$ deficiency

3. **Decreased** RBC folate occurs with:
   a. Untreated folate deficiency
   b. VB$_{12}$ deficiency (60% of uncomplicated cases)

**Interfering Factors**

1. Drugs that are folic acid antagonists, among others (see Appendix J)
2. Hemolyzed specimens (false elevation)
3. Iron-deficiency anemia (false increase)
Clinical Alert

Elderly persons and those with inadequate diets may develop folate-deficient megaloblastic anemia.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. Obtain pertinent medication history.
2. Alert patient that fasting from food for 8 hours before testing is required; water is permitted.
3. Draw blood before VB_{12} injection.
4. Do not administer radioisotopes for 24 hours before the specimen is drawn.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume normal activities and medications.
2. Interpret test results; counsel and monitor appropriately for anemia.
3. See Chapter 1 guidelines for safe, effective, informed posttest care. See Appendix F for more information on vitamin testing.

Erythropoietin (Ep)

Erythropoietin (Ep) is a glycoprotein hormone that regulates erythropoiesis. The levels of Ep in anemia are primarily determined by the degree of anemia; Ep is inversely related to red blood cell volume and Hct.

Ep is used to investigate obscure anemias. This test is useful in differentiating primary from secondary polycythemia and in detecting the recurrence of Ep-producing tumors. It is also used as an indicator of need for Ep therapy in patients with renal failure (end-stage renal disease).

Reference Values

Normal 5–36 mU/mL or 5–36 U/L

Procedure

1. Obtain a venous blood serum sample of 5 mL. Place the specimen in biohazard bag.
2. Separate serum from cells as soon as possible and place in polypropylene tube (not clear plastic-polystyrene). Freeze.

Clinical Implications

1. Ep is increased appropriately in:
   a. Anemias with very low Hb (eg, aplastic anemia, hemolytic anemia); hematologic cancers have very high levels.
   b. Patients with any iron-deficiency anemia have moderately high levels.
   c. Myelodysplasia, chemotherapy, AIDS
   d. Secondary polycythemia vera caused by tissue hypoxia (eg, high altitude, COPD)
   e. Pregnancy (very high values)
2. Ep is increased inappropriately in erythropoietin-producing tumors:
   a. Renal cysts, renal transplant rejection
   b. Renal adenocarcinoma
   c. Pheochromocytomas
   d. Cerebellar hemangioblastomas
   e. Polycystic kidney disease
   f. Occasionally, adrenal, ovarian, testicular, breast, and hepatic carcinoma
3. Ep is decreased appropriately in:
   a. Rheumatoid arthritis
   b. Multiple myeloma
   c. Cancer
4. Ep is decreased inappropriately in:
   a. Polycythemia vera (primary)
   b. After bone marrow transplantation (weeks 3 and 4)
   c. AIDS before initiating therapy
   d. Autonomic neuropathy
   e. Renal failure and adult nephrotic syndrome

Interfering Factors

1. Ep is increased in:
   a. Pregnancy
   b. Use of anabolic steroids
   c. Administration of thyroid-stimulating hormone, ACTH, epinephrine
   d. Growth hormone (see Appendix J)
2. Ep is decreased in:
   a. Transfusions
   b. Use of some prescribed drugs (see Appendix J)
   c. Drugs that increase renal blood flow (eg, enalapril)
Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Draw blood at the same time for serial determinations: Circadian rhythm is lowest in the morning and 40% higher in late evening.
3. Alert patient that fasting is not necessary, but a morning specimen is needed.
4. Note use of any drugs.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume normal activities and medications.
2. Interpret test results; counsel and monitor appropriately for anemia.
3. See Chapter 1 guidelines for safe, effective, informed posttest care. See Appendix F for more information on vitamin testing.

Tests of Hemostasis and Coagulation

The prime functions of the coagulation mechanism are to protect the integrity of the blood vessels while maintaining the fluid state of blood. Serious medical problems or even death may occur with the inability to stem the loss of blood, or for the inability for a normal clot to form.

Hemostasis and coagulation tests are generally done for patients with bleeding disorders, vascular injury or trauma, or coagulopathies. Reflex vasoconstriction is the normal response to vascular insult once the first-line defenses (skin and tissue) are breached. In larger vessels, vasoconstriction may be the primary mechanism for hemostasis. With smaller vessels, vasoconstriction reduces the size of the area that must be occluded by the hemostatic plug. Part of this cascade of sequential clotting events relates to the fact that platelets adhere to the injured and exposed subendothelial tissues. This phenomenon initiates the complex clotting mechanism whereby thrombin and fibrin are formed and deposited to aid in intravascular clotting (Table 2.4).

Table 2.4 The Complex Chain of Coagulation Reactions

<table>
<thead>
<tr>
<th>Stage</th>
<th>Components of Stages</th>
</tr>
</thead>
<tbody>
<tr>
<td>STAGE I (3–5 MIN)</td>
<td>90% of all coagulation disorders are caused by defects in phase I. Platelet counts &lt;1 × 10^6/mm^3 indicate moderate interference with phase I activity.</td>
</tr>
<tr>
<td>Phase I—platelet activity; platelets serve as a source of thromboplastin.</td>
<td>Calcium Factor V Factor VIII Factor IX Factor X Factor XI Factor XII are involved in the formation of tissue thromboplastin (intrinsic prothrombin activation)</td>
</tr>
<tr>
<td>Phase II—thromboplastin; factor III, an enzyme thought to be liberated by damaged cells, is formed by six different factors plus calcium.</td>
<td>Factor II Factor X Factor VII Factor V are involved in the conversion of fibrinogen to fibrin</td>
</tr>
<tr>
<td>STAGE II (8–15 SEC)</td>
<td>Prothrombin factor II is converted to thrombin in the presence of calcium.</td>
</tr>
<tr>
<td>STAGE III (1 SEC)</td>
<td>Thrombin interacts with fibrinogen (factor I) to form the framework of the clot. At the end stage III, factor XIII functions in the stabilization of the clot.</td>
</tr>
<tr>
<td>STAGE IV</td>
<td></td>
</tr>
</tbody>
</table>
Fibrinolytic system (antagonistic check-and-balance to the clotting mechanism) is activated. Removal of fibrin clot through fibrinolysis. Plasminogen is converted to plasmin, which breaks clot into fibrin split products.

The entire mechanism of coagulation and fibrinolysis (removal of fibrin clot) is one of balance. It may best be understood by referring to the diagrams in this section. Abnormal bleeding does not always indicate coagulopathy, in much the same way that lack of bleeding does not necessarily indicate absence of a bleeding disorder.

The most common causes of hemorrhage are thrombocytopenia (platelet deficiency) and other acquired coagulation disorders, including liver disease, uremia, disseminated intravascular coagulation (DIC), and anticoagulant administration. Together, they account for most hemorrhagic problems. Hemophilia and other inherited factor deficiencies are seen less frequently. Bleeding tendencies are associated with delays in clot formation or premature clot lysis. Thrombosis is associated with inappropriate clot activation or localization of the blood coagulation process. Finally, clotting disorders are divided into two classes: those caused by impaired coagulation and those caused by hypercoagulability.

Hypercoagulability States

Two general forms of hypercoagulability exist: hyperreactivity of the platelet system, which results in arterial thrombosis, and accelerated activity of the clotting system, which results in venous thrombosis. Hypercoagulability refers to an unnatural tendency toward thrombosis. The thrombus is the actual insoluble mass (fibrin or platelets) present in the bloodstream or chambers of the heart.

Conditions and classifications associated with hypercoagulability include the following:

**Platelet Abnormalities.** These conditions are associated with arteriosclerosis, diabetes mellitus, increased blood lipids or cholesterol levels, increased platelet levels, and smoking. Arterial thrombosis may be related to blood flow disturbances, vessel wall changes, and increased platelet sensitivity to factors causing platelet adherence and aggregation.

**Clotting System Abnormalities.** These are associated with congestive heart failure, immobility, artificial surfaces (eg, artificial heart valves), damaged vasculature, use of oral contraceptives or estrogen, pregnancy and the postpartum state, and the postsurgical state. Other influences include malignancy, myeloproliferative (bone marrow) disorders, obesity, lupus disorders, and genetic predisposition.

**Venous Thrombosis.** This can be related to stasis of blood flow, to coagulation alterations, or to increases in procoagulation factors or decreases in anticoagulation factors (Table 2.5).

### Table 2.5 Proteins Involved in Blood Coagulation

<table>
<thead>
<tr>
<th>Protein *</th>
<th>Synonym</th>
<th>Plasma Concentration (mg/dL)</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fibrinogen</td>
<td>Factor I</td>
<td>200–400</td>
<td>Converted to fibrin along with platelets to form clot</td>
</tr>
<tr>
<td>Factor II</td>
<td>Prothrombin (prethrombin)</td>
<td>10–15</td>
<td>Is converted to thrombin (IIa), which splits fibrinogen into fibrin</td>
</tr>
<tr>
<td>Factor V</td>
<td>Proaccelerin; labile factor</td>
<td>0.5–1.0</td>
<td>Supports Xa activation of II to IIa</td>
</tr>
<tr>
<td>Factor VII</td>
<td>Stable factor; proconvertin</td>
<td>0.2</td>
<td>Activates X</td>
</tr>
<tr>
<td>Factor VIII:C</td>
<td>Antihemophilic factor (AHF) platelet cofactor I</td>
<td>1.0–2.0</td>
<td>Supports IXa activation of X</td>
</tr>
<tr>
<td>Factor IX</td>
<td>Christmas factor; plasma thromboplastin component (PTC)</td>
<td>0.3–0.4</td>
<td>Activates X</td>
</tr>
<tr>
<td>Factor X</td>
<td>Stuart-Prower factor (AVTD prothrombin III)</td>
<td>0.6–0.8</td>
<td>Activates II</td>
</tr>
<tr>
<td>Factor XI</td>
<td>Plasma thromboplastin antecedent (antihemophilic factor C)</td>
<td>0.4</td>
<td>Activates XII and prekallikrein</td>
</tr>
<tr>
<td>Factor XII</td>
<td>Hageman factor</td>
<td>2.9</td>
<td>Activates XI and prekallikrein</td>
</tr>
<tr>
<td>Factor XIII</td>
<td>Fibrin-stabilizing factor; Laki-Lorand factor</td>
<td>2.5</td>
<td>Crosslinks fibrin and other proteins</td>
</tr>
<tr>
<td>von Willebrand's factor</td>
<td>Factor VIII–related antigen VIII:VWD</td>
<td>1.0</td>
<td>Stabilizes VIII, mediates platelet adhesion</td>
</tr>
<tr>
<td>Prekallikrein</td>
<td>Fletcher factor</td>
<td>5.0</td>
<td>Activates XII and prekallikrein, cleaves HMWK</td>
</tr>
<tr>
<td>High-molecular-weight kininogen (HMWK)</td>
<td>Fitzgerald factor</td>
<td>4.7–12.2</td>
<td>Supports reciprocal activation of XII, XI and prekallikrein</td>
</tr>
<tr>
<td>Fibronectin</td>
<td>Cold insoluble globulin</td>
<td>20–40</td>
<td>Mediates cell adhesion</td>
</tr>
</tbody>
</table>
Major antithrombin  Antithrombin III  20–40  Inhibits IIa, Xa, Xla, and kallikrein
Protein C  0.5  Complexed with protein S, inactivates V and VIII
Plasminogen  20  Forms plasmin, which lyses the fibrin clot and inhibits other factors
a 2-Antiplasmin  9.6–13.5  Inhibits plasmin
a 1-Antitrypsin  245–335  Weak inhibitor of thrombin, potent inhibitor of Xla
Tissue plasminogen activator inhibitor I  Activates plasminogen
Plasminogen activator inhibitor II  Inactivates tissue plasminogen activator (tPA)
Plasminogen activator inhibitor III  Inactivates urokinase

*The clotting factors of the blood are proteins; they are present in the blood plasma in an inactive form called zymogens.

Disorders of Hemostasis

Congenital Vascular Abnormalities (Vessel Wall Structure Defects). Defects of the actual blood vessel are poorly defined and difficult to test. Hereditary telangiectasia is the most commonly recognized vascular abnormality. Laboratory studies are normal, so the diagnosis must be made from clinical signs and symptoms. Patients frequently report epistaxis and symptoms of anemia. Another abnormality is congenital hemangiomas (Kasabach-Merritt syndrome).

Acquired Abnormalities of the Vessel Wall Structure. Causes include Henoch-Schönlein purpura as an allergic response to infection or drugs, diabetes mellitus, rickettsial diseases, septicemia, and amyloidosis present with some degree of vascular abnormalities. Purpura can also be associated with steroid therapy and easy bruising in females (infectious purpura), or it can be a result of drug use.

Hereditary Connective Tissue Disorders. These include Ehlers-Danlos syndrome (hyperplastic skin and hyperflexible joints) and pseudoxanthoma elasticum (rare connective tissue disorder).

Acquired Connective Tissue Defects. These can be caused by scurvy (vitamin C deficiency) or senile purpura.

Qualitative Platelet Abnormalities. These disorders can be divided into subclasses:

1. Thrombocytopenia (platelet count <150 × 10^3/mm^3) is caused by decreased production of platelets, increased use or destruction of platelets, or hypersplenism. Contributing factors include bone marrow disease, autoimmune diseases, DIC, bacterial or viral infection, chemotherapy, therapy radiation, multiple transfusions, and certain drugs (eg, NSAIDs thiazides, estrogens).

2. Thrombocytosis (elevated platelet count) is caused by hemorrhage, iron-deficiency anemia, inflammation, or splenectomy.

3. Thrombocythemia (platelet count >1000 × 10^3/mm^3 or >1000 × 10^9/L) is caused by granulocytic leukemia, polycythemia vera, or myeloid metaplasia.

Clinical Alert
An increased platelet count predisposes the patient to arterial thrombosis. Paradoxically, a substantially elevated platelet count can also cause easy bleeding after dental surgery, gastrointestinal bleeding, and epistaxis.

Quantitative Platelet Abnormalities. These are associated with Glanzmann's thrombasthenia, a hereditary autosomal-recessive disorder that can produce severe bleeding, especially with trauma and surgical procedures. Platelet factor 3 differences associated with aggregation, adhesion, or release defects may be manifested in storage-pool disease, May-Hegglin anomaly, Bernard-Soulier syndrome, and Wiskott-Aldrich syndrome. Dialysis and use of drugs such as aspirin, other antiinflammatory agents, dipyridamole, and prostaglandin E also can be tied to platelet abnormalities.

Congenital Coagulation Abnormalities. These include hemophilia A and B (deficiencies of factors VIII and IX, respectively), rare autosomal recessive traits (hemophilia C), and autosomal dominant traits (eg, von Willebrand's disease).

Acquired Coagulation Abnormalities. These are associated with several disease states and are much more common than inherited deficiencies.

1. Circulatory anticoagulant activity may be evident in the presence of antifactor VIII, rheumatoid arthritis, immediate
postpartum period, SLE, or multiple myeloma.

2. Vitamin D deficiency may be caused by oral anticoagulants, biliary obstruction and malabsorption syndrome, or intestinal sterilization by antibiotic therapy. Newborns are prone to vitamin D deficiency.

3. DIC causes continuous production of thrombin, which, in turn, consumes the other clotting factors and results in uncontrolled bleeding.

4. Primary fibrinolysis is the situation whereby isolated activation of the fibrinolytic mechanism occurs without prior coagulation activity, as in streptokinase therapy, severe liver disease, prostate cancer, or, more rarely, electroshock.

5. Most coagulation factors are manufactured in liver. Consequently, in liver disease, the extent of coagulation abnormalities is directly proportional to the severity of the liver disease.

Tests for Disseminated Intravascular Coagulation (DIC)

DIC is an acquired hemorrhagic syndrome characterized by uncontrolled formation and deposition of fibrin thrombi. Continuous generation of thrombin causes depletion (consumption) of the coagulation factors and results in uncontrolled bleeding. Also, fibrinolysis is activated in DIC. This further adds to the hemostasis defect caused by the consumption of clotting factors. The many coagulation test abnormalities found in acute DIC include the following:

1. Prolonged
   a. Prothrombin time (PT)
   b. Partial thromboplastin time (PTT) or activated partial thromboplastin time (APTT)
   c. Bleeding time
   d. Thrombin time (TT)

2. Decreased
   a. Fibrinogen
   b. Platelet count
   c. Clotting factors II, V, VIII, and X
   d. Antithrombin III (AT-III)

3. Increased
   a. Fibrinolysin test
   b. Fibrinopeptide A

4. Positive
   a. Fibrin split products
   b. D-Dimer

In chronic DIC, the results are variable, especially the PT, PTT, TT, and fibrinogen, making the diagnosis much more difficult. No single test or group of tests is diagnostic, and diagnosis usually depends on a combination of findings. Normal levels do not rule out DIC, and a repeat profile should be done a few hours later to look for changes in platelet count and fibrinogen.

Causes of DIC include septicemia, malignancies and cancer, obstetric emergencies, cirrhosis of liver, sickle cell disease, trauma or crushing injuries, malaria, incompatible blood transfusion, cold hemoglobinuria or PNH, connective tissue diseases, snake bites, and brown recluse spider bites.

Paradoxically, the treatment of uncontrolled bleeding in DIC is heparin administration. The heparin blocks thrombin formation, which blocks consumption of the other clotting factors and allows hemostasis to occur.

Laboratory Investigation of Hemostasis

Usually, a blood sample of at least 20 mL is obtained by the two-tube technique. In the first tube, a 5-mL blood sample is obtained and discarded. Then 15 to 20 mL of blood is drawn into Vacutainer tubes with sodium citrate as the anticoagulant. A butterfly needle may be used to prevent backflow or to make sampling easier in the case of a difficult draw. Coagulation studies (coagulation profiles, coag panels, coagulograms) are used for screening or as diagnostic tools for evaluation of symptoms such as easy or spontaneous bruising, petechiae, prolonged bleeding (eg, from cuts), abnormal nosebleeds, heavy menstrual flow, family history of coagulopathies, or gastrointestinal bleeding (Table 2.6).

<table>
<thead>
<tr>
<th>Name of Test</th>
<th>Vascular Function</th>
<th>Platelet Function</th>
<th>Stage I</th>
<th>Stage II</th>
<th>Stage III</th>
<th>Stage IV</th>
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<td>Bleeding time</td>
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<tr>
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<td></td>
<td></td>
<td></td>
<td>X</td>
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<tr>
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<tr>
<td>Activated clotting</td>
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<td></td>
<td></td>
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<tr>
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<td></td>
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</table>
Fibrinogen assay X
Factor XIII assay X
Euglobulin lysis time X
Thrombin time—diluted X X
Plasminogen assay X
Protamine sulfate (ethanal gelation) X X
D-Dimer X
Fibrin monomer X
Fibrinopeptide A X
Latex agglutination for fibrin split products X

*These tests measure all facets of hemostasis: vascular function, platelets, and clotting factors.

**Activates factor X.

Many of the more common screening tests are now automated and easily done. Platelet counts are included in the automated CBC with most instruments. Protime, and PTT can be done on photooptical instruments that sense the change in optical density when a clot forms. Tests for fibrinogen are on instruments that detect fibrin strands. Many patients can undergo testing at the same time with the help of automation. Some of the more specialized tests still must be done manually or using semiautomated methods.

1. These five primary screening tests are initially performed to diagnose suspected coagulation disorders:
   a. Platelet count, size, and shape
   b. Bleeding time—reflects data about the ability of platelets to function normally and the ability of the capillaries to constrict their walls
   c. PTT—determines the overall ability of the blood to clot
   d. PT—measures the function of second-stage clotting factors
   e. Fibrinogen level

2. Factor assays are definitive coagulation studies of a specific clotting factor (eg, factor VIII for hemophilia). These are done if the screening test indicates a problem with a specific factor or factors.

3. Fibrinolysis is used to address problems of the fibrinolytic system and includes the following studies:
   a. Euglobulin clot lysis—identifies increased plasminogen activator activity. (Plasmin is not usually present in the blood plasma.)
   b. Factor XIII (fibrin-stabilizing factor)
   c. Fibrin split products (eg, protamine sulfate test)

4. The investigation of hypercoagulable status (thrombotic tendency, thromboembolic disorders) covers both primary causes (deficiencies of AT-III, protein C, protein S, and factor XII; fibrinolytic mechanisms) and secondary causes (acquired platelet disorders and acquired diseases of coagulation and fibrinolytic impairment) and includes the following tests:
   a. PT
   b. PTT
   c. Fibrinogen test
   d. Antiplatelet factors (eg, prostacyclin)
   e. Anticoagulant factors (eg, AT-III, protein C, protein S, lupus anticoagulant)
   f. Fibrinolysis tests (eg, fibrin degradation products [FDPs], euglobulin lysis time, fibrin monomers)
   g. TT

NOTE

The lupus inhibitor (lupus anticoagulant) is an antibody (against the phospholipid used in the PT and PTT tests) that is responsible for inhibition of the PT, PTT, Russell viper venom time (dRVVT), and kaolin clotting time (kCT). To demonstrate its presence, 1 mL of the patient's plasma is mixed with 1 mL of normal plasma, and a PTT test of the mixture is done. When an inhibitor of any sort is present, the PTT will not return to normal range. An inhibitor of the lupus type can be shown by correcting the PTT through use of platelets as a phospholipid source or by demonstrating a characteristic pattern in the PTT that results from sequential dilution of the phospholipid reagent. Lupus anticoagulants may be associated with false-positive Venereal Disease Research Laboratory (VDRL) test reports and with another antiphospholipid—the anticardiolipin antibody (β 2-glycoprotein I).

Clinical Alert

Conditions associated with the presence of the lupus anticoagulant include:

1. SLE (one fifth of patients)
2. Multiple myeloma
3. Other autoimmune diseases (rheumatoid arthritis, Raynaud's syndrome)
4. Spontaneous abortions (associated with presence of anticardiolipin autoantibody) and postpartum complications
5. Lupus anticoagulant is more often associated with thromboembolism than with bleeding problems.
6. Most lupus anticoagulant antibodies are directed against prothrombin or β 2-glycoprotein I.
Clinical Alert

1. All patients with hemorrhagic or thrombotic tendencies, or undergoing coagulation studies, should be observed closely for possible bleeding emergencies. A comprehensive history and physical examination should be done.
2. Blood samples for coagulation studies should be drawn last if other blood studies are indicated.
3. Procedure alert: when a blood sample is obtained for PT, PTT, and TT, sodium citrate is used as the anticoagulant in the sampling tubes.

Patient Assessment for Bleeding Tendency

1. Examine all skin for bruising.
2. Record petechiae associated with use of blood pressure cuffs or tourniquets.
3. Note bleeding from the nose or gums with no apparent cause.
4. Estimate blood quantity in vomitus, expectorated mucus, urine, stools, and menstrual flow.
5. Note prolonged bleeding from injection sites.
6. Watch for symptoms, especially changes in levels of consciousness or neurologic checks that may signal an intracranial bleed.
7. Determine whether the patient is taking anticoagulants or aspirin.

Bleeding Time (Ivy Method; Template Bleeding Time)

Bleeding time measures the primary phase of hemostasis: the interaction of the platelet with the blood vessel wall and the formation of a hemostatic plug. Bleeding time is the best single screening test for platelet function disorders and is one of the primary screening tests for coagulation disorders.

This test is of value in detecting vascular abnormalities and platelet abnormalities or deficiencies. It is not recommended for routine presurgical workup.

A small stab wound is made in either the earlobe or the forearm; the bleeding time (the amount of time it takes to form a clot) is recorded. The duration of bleeding from a punctured capillary depends on the quantity and quality of platelets and the ability of the blood vessel wall to constrict.

The principal use of this test today is in the diagnosis of von Willebrand's disease, an inherited defective molecule of factor VIII and a type of pseudohemophilia. It has been established that aspirin may cause abnormal bleeding in some normal persons, but the bleeding time test has not proved consistently valuable in identifying such persons.

Reference Values

Normal 3–10 minutes in most laboratories Duke method (earlobe): 5 minutes (not recommended—not very reproducible with a wide range of normal values) Ivy method (forearm with template): 25–90 minutes Mielke's method (Surgicut):
Adults: 1–7 minutes Teens: 3.0–8 minutes Children: 2.5–13 minutes

Procedure (Ivy Method)

1. Cleanse the area three fingerwidths below the antecubital space with alcohol and allow to dry.
2. Place a blood pressure cuff on the arm above the elbow and inflate to 40 mm Hg.
3. Select a cleansed area of the forearm without superficial veins. Stretch the skin laterally and tautly between the thumb and forefinger.
4. Start a stopwatch. Use the edge of a 4? × 4? filter paper to blot the blood through capillary action by gently touching the drop every 30 seconds. Do not disturb the wound itself. Remove the blood pressure gauge when bleeding stops and a clot has formed. Apply a sterile dressing when the test is completed.
5. Remember that the end point (by the Ivy or the earlobe method) is reached when blood is no longer blotted from the forearm puncture. Report in minutes and half minutes (eg, 5 minutes, 30 seconds).

Clinical Implications

1. Bleeding time is prolonged when the level of platelets is decreased or when platelets are qualitatively abnormal:
   a. Thrombocytopenia (platelet count <80 × 10^3/mm^3)
   b. Platelet dysfunction syndromes
   c. Decrease or abnormality in plasma factors (eg, von Willebrand's factor, fibrinogen)
   d. Abnormalities in the walls of the small blood vessels, vascular disease
   e. Advanced renal failure
   f. Severe liver disease
   g. Leukemia, other myeloproliferative diseases
   h. Scurvy
      i. DIC disease (owing to the presence of FDPs)
2. In von Willebrand's disease, bleeding time can be variable; it will definitely be prolonged if aspirin is taken before testing (aspirin tolerance test).
3. A single prolonged bleeding time does not prove the existence of hemorrhagic disease. Because a larger vessel can be punctured, the puncture should be repeated on an alternate body site, and the two values obtained should be averaged.
4. Bleeding time is normal in the presence of coagulation disorders other than platelet dysfunction, vascular disease, or von Willebrand's disease.
5. Aspirin therapy (antiplatelet function therapy): when thrombus formation is thought to be mediated by platelet activation, the patient frequently is given agents to interrupt normal platelet function, which may be monitored by bleeding times or platelet aggregation studies. Aspirin is the most commonly used inhibitor; it inhibits platelet
adhesion or "stickiness."

Interfering Factors

1. Normal values for bleeding time vary when the puncture site is not of uniform depth and width.
2. Touching the puncture site during this test will break off fibrin particles and prolong the bleeding time.
3. Excessive alcohol consumption (as in alcoholic patients) may cause increased bleeding time.
4. Prolonged bleeding time can reflect ingestion of 10 g of aspirin as long as 5 days before the test.
5. Other drugs that may cause increased bleeding times include dextran, streptokinase-streptodornase (fibrinolytic agents), mithramycin, pantotethanol alcohol (see Appendix J).
6. Extreme hot or cold conditions can alter the results.
7. Edema of patient's hands or cyanotic hands will invalidate the test.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. See Patient Assessment for Bleeding Tendency on page 131.
2. Instruct patient to abstain from aspirin and aspirin-like drugs for at least 7 days before the test.
3. Advise the patient to abstain from alcohol before the test.
4. Inform the patient that scar tissue may form at the puncture site (keloid formation).
5. If the patient has an infectious skin disease, postpone the test.
6. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately for prolonged bleeding. See Patient Assessment for Bleeding Tendency on page 131.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

1. The critical value for bleeding time is >15 minutes.
2. If the puncture site is still bleeding after 15 minutes, discontinue the test and apply pressure to the site.
   Document and report the results to the clinician.

Platelet Count; Mean Platelet Volume (MPV)

Platelets (thrombocytes) are the smallest of the formed elements in the blood. These cells are nonnucleated, round or oval, flattened, disk-shaped structures. Platelet activity is necessary for blood clotting, vascular integrity and vasoconstriction, and the adhesion and aggregation activity that occurs during the formation of platelet plugs that occlude (plug) breaks in small vessels. Thrombocyte development takes place primarily in the bone marrow. The life span of a platelet is about 7.5 days. Normally, two thirds of all the body platelets are found in the circulating blood and one third in the spleen.

The platelet count is of value for assessing bleeding disorders that occur with thrombocytopenia, uremia, liver disease, or malignancies and for monitoring the course of disease associated with bone marrow failure. This test is indicated when the estimated platelet count (on a blood smear) appears abnormal. It is also part of a coagulation profile or workup.

The mean platelet volume (MPV) is sometimes ordered in conjunction with a platelet count. The MPV indicates the uniformity of size of the platelet population. It is used for the differential diagnosis of thrombocytopenia.

Reference Values

**Normal Platelet count:** Adults: 140–400 × 10^3/mm^3 or 140–400 × 10^9/L Children: 150–450 × 10^3/mm^3 or 150–450 × 10^9/L

**Mean platelet volume:** Adults: 7.4–10.4 µm^3 or fl Children: 7.4–10.4 µm^3 or fl

Procedure

1. Mix a 7-mL venous blood sample with an EDTA anticoagulant tube.
2. Count the platelets by phase microscopy or by an automated counting instrument. The MPV is also calculated by many instruments at the time of the platelet count.
3. Make a blood smear and note the size, shape, and clumping of the platelets.
4. Place the specimen in a biohazard bag.

Clinical Implications

1. Abnormally increased numbers of platelets (thrombocytethemia, thrombocytosis) occur in:
   a. Essential thrombocytethemia
   b. Chronic myelogenous and granulocytic leukemia, myeloproliferative diseases
   c. Polycythemia vera and primary thrombocytosis
   d. Splenectomy
   e. Iron-deficiency anemia
   f. Asphyxiation
   g. Rheumatoid arthritis and other collagen diseases, SLE
   h. Rapid blood regeneration caused by acute blood loss, hemolytic anemia
i. Acute infections, inflammatory diseases
j. Hodgkin’s disease, lymphomas, malignancies
k. Chronic pancreatitis, tuberculosis, inflammatory bowel disease
l. Renal failure
m. Recovery from bone marrow suppression (thrombocytopenia)

2. Abnormally decreased numbers of platelets (thrombocytopenia) occur in:
   a. Idiopathic thrombocytopenic purpura, neonatal purpura
   b. Pernicious, aplastic, and hemolytic anemias
   c. After massive blood transfusion (dilution effect)
   d. Viral, bacterial, and rickettsial infections
   e. Congestive heart failure, congenital heart disease
   f. Thrombopoietin deficiency
   g. During cancer chemotherapy and radiation, exposure to dichlorodiphenyl-trichloroethane (DDT) and other chemicals
   h. HIV infection
   i. Lesions involving the bone marrow (eg, leukemias, carcinomas, myelofibrosis)
   j. DIC and thrombotic thrombocytopenic purpura
   k. Inherited syndromes such as Bernard-Soulier syndrome, May-Hegglin anomaly, Wiskott-Aldrich syndrome, Fanconi’s syndrome
   l. Toxemia of pregnancy, eclampsia
   m. Alcohol toxicity, ethanol abuse
   n. Hypersplenism
   o. Renal insufficiency
   p. Antiplatelet antibodies

3. Increased MPV is observed in:
   a. Idiopathic thrombocytopenic purpura (autoimmune)
   b. Thrombocytopenia caused by sepsis
   c. Prosthetic heart valve
   d. Massive hemorrhage
   e. Myeloproliferative disorders
   f. Acute and chronic myelogenous leukemia
   g. Splenectomy
   h. Vasculitis
   i. Megaloblastic anemia

4. Decreased MPV occurs in Wiskott-Aldrich syndrome.

Clinical Alert

1. In 50% of patients who exhibit unexpected platelet increases, a malignancy is found.
2. In patients with an extremely elevated platelet count (>1000 × 10^3/mm^3 or >1000 × 10^9/L) as a result of a myeloproliferative disorder, assess for bleeding caused by abnormal platelet function.

NOTE

Many drugs have toxic effects. The dosage does not have to be high to be toxic. Toxic thrombocytopenia depends on the inability of the body to metabolize and secrete the toxic substance.

Clinical Alert

1. Panic values: a decrease in platelets to <20 × 10^3/mm^3 or <20 × 10^9/L is associated with a tendency for spontaneous bleeding, prolonged bleeding time, petechiae, and ecchymosis.
2. Platelet counts >50 × 10^3/mm^3 or >50 × 10^9/L are not generally associated with spontaneous bleeding.

Interfering Factors

1. Platelet counts normally increase at high altitudes; after strenuous exercise, trauma, or excitement; and in winter.
2. Platelet counts normally decrease before menstruation and during pregnancy.
3. Clumping of platelets may cause falsely lowered results.
4. Oral contraceptives cause a slight increase.
5. See Appendix J for drugs that affect test outcomes.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Avoid strenuous exercise before blood is drawn.
3. Note what medications and what treatments the patient is receiving.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately. Observe for signs and symptoms of gastrointestinal bleeding, hemolysis, hematuria, petechiae, vaginal bleeding, epistaxis, and bleeding from the gums. When hemorrhage is apparent, use emergency measures to control bleeding and notify the attending physician.
2. Use platelet transfusions if the platelet count is <20 × 10^3/mm^3 (<20 × 10^9/L) or if there is a specific bleeding
lesion. One unit of platelet concentrate raises the count by $15 \times 10^3 \text{/mm}^3 (15 \times 10^5 \text{/L})$.

3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Platelet Aggregation

Platelet aggregation is used to evaluate congenital qualitative functional disorders of adhesion, release, or aggregation. It is rarely used to evaluate acquired bleeding disorders.

Reference Values

Normal: Full platelet aggregation in response to the following: Adenosine diphosphate, Collagen, Epinephrine, Thrombin, Ristocetin

Procedure

1. Obtain a 5-mL venous blood sample (anticoagulated in a tube containing sodium citrate).
2. Place it in a biohazard bag. The sample is kept at room temperature (never refrigerate) and must be run within 30 minutes after the blood is drawn.
3. Increase the transmission of light through a sample of platelet-rich plasma when platelets aggregate. This increase in light transmission can be used as an index to the aggregation in response to various agonists.

Clinical Implications

1. Decreased platelet aggregation occurs in congenital diseases:
   a. Bernard-Soulier syndrome
   b. Glanzmann's thrombasthenia
   c. Storage pool diseases (eg, Chédiak-Higashi syndrome, gray platelet disease)
   d. Cyclooxygenase deficiency
   e. Wiskott-Aldrich syndrome
   f. Albinism
   g. β-Thalassemia major
   h. May-Hegglin anomaly
   i. Various connective tissue disorders (eg, Marfan's syndrome)
   j. von Willebrand's disease
2. Decreased platelet aggregation also occurs in acquired disorders:
   a. Uremia
   b. Antiplatelet antibodies
   c. Cardiopulmonary bypass
   d. Myeloproliferative disorders
   e. Dysproteinemias (macroglobulinemia)
   f. Idiopathic thrombocytopenic purpura
   g. Polycythemia vera
   h. Use of drugs and aspirin, some antibiotics, anti-inflammatory drugs, psychotropic drugs, and others (see Appendix J)
   i. DIC
3. Increased aggregation occurs in primary and secondary Raynaud's syndrome.

Interfering Factors

1. Platelet count <100,000/mm$^3$
2. Patient cannot be taking drugs that interfere with aggregation (see Appendix J).
3. Lipemia will interfere with testing.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Be aware that for 10 days before test, drugs that inhibit platelet aggregation are contraindicated. These include aspirin, antihistamines, steroids, cocaine, anti-inflammatory drugs, theophylline, and antibiotics.
3. On the day of the test, avoid caffeine.
4. Avoid warfarin (Coumadin) for 2 weeks and heparin therapy for 1 week before testing.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel appropriately for congenital disorders.
2. Resume medications and normal diet.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Thrombin Time (TT); Thrombin Clotting Time (TCT)

Stage III fibrinogen defects can be detected by the TT test. It can detect DIC and hypofibrinogenemia and may also be used for monitoring streptokinase therapy. The test actually measures the time needed for plasma to clot when thrombin is added. Normally, a clot forms rapidly; if it does not, a stage III deficiency is present (Fig. 2.1). A TT test is often included as part of a panel for coagulation defects.
FIGURE 2.1 Intrinsic, extrinsic, and common pathways of coagulation. Vessel injury initiates intrinsic pathway through contact activation by exposed collagen. Extrinsic pathway is initiated by endothelial release of tissue factor (ie, tissue thromboplastin). Extrinsic and intrinsic pathways each initiate common pathway to create stable fibrin clot. (Lotspeich-Steininger, C. A., et al. [1992]. Clinical Hematology, Philadelphia: JB Lippincott Co.)

Reference Values
Normal 7.0–12.0 seconds (varies widely by laboratory) Check with your laboratory for values.

Procedure
1. Use the procedure for two-tube specimen collection to anticoagulate a 7-mL venous blood sample with sodium citrate and put on ice. Take care not to contaminate the specimen with heparin from IV apparatus or other sources.
2. Ensure that the specimen is tested within 2 hours, or it must be frozen for later testing.

Clinical Implications
1. Prolonged TT occurs in:
   a. Hypofibrinogenemia
   b. Therapy with heparin or heparin-like anticoagulants
   c. DIC
   d. Fibrinolysis
   e. Multiple myeloma
   f. Presence of large amounts of fibrin split products (FSPs) or FDPs, as in DIC
   g. Uremia
   h. Severe liver diseases
2. Shortened TT occurs in:
   a. Hyperfibrinogenemia
   b. Elevated Hct (>55%)
3. Therapy with plasminogen activators—streptokinase, urokinase, or tissue plasminogen activator (tPA).

Anticoagulant therapy is an attempt either to prevent thrombus formation or to promote thrombus lysis. The type and location of the thrombus usually determine the type of anticoagulant to be administered and the treatment protocol. The newest treatment for life-threatening thrombus formation uses plasminogen activators to accelerate fibrinolysis, which is the enzymatic dissolution of already organized clots (Fig. 2.2). The action of some of these agents produces a lytic state that can be monitored by the TT.

Although several tests are sensitive to the effects of thrombolytic drugs, many require lengthy assay procedures or special techniques. Of the laboratory procedures that have been recommended (PT, TT, APTT, quantitative fibrinogen, euglobulin clot lysis, and plasminogen levels), the TT has become widely accepted because it is fast and practical, does not require special equipment, and can detect the decrease in fibrinogen levels as well as the presence of fibrin and FDPs. The half-life for these activators is relatively short (10–90 minutes); therefore, the antidote for overdose is to hold giving the next dose.
Clinical Alert
TT is severely prolonged in the presence of afibrinogenemia (<80 mg/dL or <0.8 g/L of fibrinogen). Critical value: >60 seconds.

Interfering Factors
1. Heparin prolongs thrombin time. Interpret test results within this context.
2. Plasminogen activator therapy prolongs TCT.
3. See Appendix J for drugs that affect test outcomes.

Interventions
Pretest Patient Care
1. Explain test purpose and procedure.
2. If possible, ensure that no heparin is taken for 2 days before testing.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Resume normal activities and medications as ordered.
2. Interpret test outcomes and monitor appropriately. Check for excess bleeding. If plasminogen activator is being monitored, see Posttest Patient Aftercare for APTT, pages 142–143.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Partial Thromboplastin Time (PTT); Activated Partial Thromboplastin Time (APTT)
The PTT, a one-stage clotting test, screens for coagulation disorders. Specifically, it can detect deficiencies of the intrinsic thromboplastin system and also reveals defects in the extrinsic coagulation mechanism pathway.

NOTE
The PTT and APTT test for the same functions. APTT is a more sensitive version of PTT that is used to monitor heparin therapy.

The APTT is used to detect deficiencies in the intrinsic coagulation system, to detect incubating anticoagulants, and to monitor heparin therapy. It is part of a coagulation panel workup.

Reference Values
Normal APTT: 21–35 seconds Check with your laboratory for therapeutic range values during heparin therapy (2–2.5 times normal).

Procedure
1. Obtain a 5-mL venous blood sample and anticoagulate with sodium citrate. Use the two-tube method. Place the specimen in a biohazard bag.
2. Do not draw blood samples from a heparin lock or heparinized catheter.
3. Be aware that the sample may be transported at room temperature, but the vacuum must be intact (do not remove stopper). It is stable for 12 hours.

Clinical Implications
1. Prolonged APTT occurs in:
   a. All congenital deficiencies of intrinsic system coagulation factors, including hemophilia A and hemophilia B
   b. Congenital deficiency of Fitzgerald's factor, Fletcher's factor (prekallikrein)
   c. Heparin therapy, streptokinase, urokinase
   d. Warfarin (Coumadin)-like therapy
   e. Vitamin K deficiency
   f. Hypofibrinogenemia
   g. Liver disease
   h. DIC (chronic or acute)
   i. Fibrin breakdown products
2. When APTT is performed in conjunction with PT, a further clarification of coagulation defects is possible. For example, a normal PT with an abnormal APTT means that the defect lies within the first stage of the clotting cascade (factors VIII, IX, X, XI, and/or XII). The pattern of a normal PTT with an abnormal PT suggests a possible factor VII deficiency. If both PT and APTT are prolonged, a deficiency of factor I, II, V, or X is suggested. Used together, APTT and PT will detect approximately 95% of coagulation defects.
3. Shortened APTT occurs in:
   a. Extensive cancer, except when the liver is involved
   b. Immediately after acute hemorrhage
   c. Very early stages of DIC
4. Circulating anticoagulants (inhibitors) usually occur as inhibitors of a specific factor (eg, factor VIII). These are most commonly seen in the development of anti–factor VIII or anti–factor IX in 5% to 10% of hemophiliac patients. Anticoagulants that develop in the treated hemophiliac are detected through prolonged APTT. Circulating anticoagulants are also associated with other conditions:
a. After many plasma transfusions
b. Drug reactions
c. Tuberculosis
d. Chronic glomerulonephritis
e. SLE
f. Rheumatoid arthritis

5. Heparin therapy: In deep vein thrombosis or acute myocardial infarction, the usual protocol requires injection of heparin (monitored by the APTT), followed by long-term therapy with oral anticoagulants (monitored by the PT, APTT, or both).
   a. In the blood, heparin combines with an a-globulin (heparin cofactor) to form a potent antithrombin. It is a direct anticoagulant.
   b. Intravenous heparin injection produces an immediate anticoagulant effect; it is chosen when rapid anticoagulant effects are desired.
   c. Because the half-life of heparin is 3 hours, the APTT is measured 3 hours after heparin administration, or 1 hour before the next dose.
   d. Therapeutic APTT levels are ordinarily maintained at 2 to 2.5 times the normal values.
   e. To evaluate heparin effects, blood is tested:
      1. For baseline values before therapy is initiated
      2. One hour before the next dose is due (when a 4-hour administration cycle is ordered)
      3. According to the patient's status (eg, bleeding)

NOTE
Mixing equal parts of patient plasma and normal plasma corrects the APTT if it is caused by a coagulation factor defect but does not correct the APTT to normal if it is caused by a circulating inhibitor. A more sensitive test is the Russell viper venom test, which demonstrates the presence of the lupus anticoagulant. This test is unaffected by inhibitors of factor VIII or deficiencies of factors VIII, IX, XI, or is affected by deficiencies of factors II, V, or X and by the use of sodium, warfarin, or heparin. Because lupus-type anticoagulants vary greatly in their reactivity in various test systems, it is recommended that this test be done in conjunction with the APTT and the anticardiolipin antibody assay. The reference range is 33.5–41.5 seconds.

Clinical Alert
Panic value: APTT >70 seconds signifies spontaneous bleeding.

NOTE
Not all individuals respond ideally or predictably to heparin. Anaphylaxis and erythematous may occur. There is no shortcut to adequate and safe anticoagulation.

Interfering Factors
1. See Appendix J for drugs that affect test outcomes.
2. Hemolized plasma shortens APTT in normal patients but not in abnormal (heparinized) patients.
3. Very increased or decreased Hct
4. Incorrect ratio of blood to citrate ("short" fill of blood in collection tube)

Interventions

Pretest Patient Care
1. Explain test purpose, procedure, benefits, and risks.
2. See Chapter 1 guidelines for safe, effective, informed, pretest care.
3. Draw blood sample 1 hour before next dose of heparin. The heparin dose given relates to the APTT result.

Posttest Patient Aftercare
1. Interpret test outcome and monitor appropriately. Protamine sulfate is the antidote for heparin overdose or for reversal of heparin anticoagulation therapy.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.
3. Watch for signs of spontaneous bleeding; notify physician immediately and treat accordingly.
4. Alert the patient to watch for bleeding gums, hematuria, oozing from wounds, and excessive bruising.
5. Instruct the patient to use an electric shaver instead of a blade and to exercise caution in all activities.
6. Avoid use of aspirin or ASA-like drugs (unless specifically prescribed) because they contribute to bleeding tendencies.
7. Be aware that long-term use of heparin can cause development of osteoporosis with fractures.
8. Remember that thrombocytopenia can also develop with high-dose heparin therapy, along with progressive thromboembolic syndrome. This platelet abnormality quickly reverses when heparin is discontinued.

Activated Coagulation Time (ACT)
The ACT test evaluates coagulation status. The ACT responds linearly to heparin level changes and responds to wider ranges of heparin concentrations than does the APTT. The ACT however, assays overall coagulation activity. Therefore, prolonged values may not be exclusively the result of heparin.

The ACT can be a bedside procedure and requires only 0.4 mL of blood. Heparin infusion or reversal with protamine can
then be titrated almost immediately according to the ACT results. ACT also is routinely used during dialysis, coronary artery bypass procedures, arteriograms, and percutaneous transluminal coronary arteriography. This test is hard to standardize, and no controls are available; therefore, it is used with caution mainly in cardiac surgery. The results are backed up by the APTT.

Reference Values

**Normal** ACT: 70–120 seconds Therapeutic range: 180–240 seconds (two times normal range)

### Prothrombin Time (Pro Time; PT)

Prothrombin is a protein produced by the liver for clotting of blood. Prothrombin production depends on adequate vitamin K intake and absorption. During the clotting process, prothrombin is converted to thrombin. The prothrombin content of the blood is reduced in patients with liver disease.

The PT is one of the four most important screening tests used in diagnostic coagulation studies. It directly measures a potential defect in stage II of the clotting mechanism (extrinsic coagulation system) through analysis of the clotting ability of five plasma coagulation factors (prothrombin, fibrinogen, factor V, factor VII, and factor X). In addition to screening for deficiency of prothrombin, the PT is used to evaluate disfibrinogenemia, evaluate the heparin effect and coumarin effect, liver failure, and vitamin K deficiency.

Reference Values

**Normal** PT: 11.0–13.0 seconds (can vary by laboratory) Therapeutic levels are at a P/C ratio of 2.0–2.5. Recommended therapeutic ranges are shown in Table 2.7.

<table>
<thead>
<tr>
<th>Table 2.7 Therapeutic Context</th>
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<tbody>
<tr>
<td>P/C ratio (prothrombin time ratio): the observed patient PT divided by the laboratory PT mean normal value INR (International Normalized Ratio): a comparative rating of PT ratios (representing the observed PT ratio adjusted by the International Reference Thromboplastin) ISI (International Sensitivity Index): a comparative rating of thromboplastin (supplied by the manufacturer of the reagent)</td>
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</table>

**Procedure**

1. Draw a 5-mL venous blood sample (by the two-tube technique) into a tube containing a calcium-binding anticoagulant (sodium citrate). The ratio of sodium citrate to blood is critical.
2. Use blue-topped vacuum tubes that keep prothrombin levels stable at room temperature for 12 hours if left capped (vacuum intact). Place the specimen in biohazard bag.

**Oral Anticoagulant Therapy** Oral anticoagulant drugs (eg, Coumadin, dicumarol) are commonly prescribed to treat blood clots. These are indirect anticoagulants (compared with heparin, which is a direct anticoagulant). However, if necessary, heparin is the anticoagulant of choice for initiating treatment because it acts rapidly and also partially lyses the clot.

1. These drugs act via the liver to delay coagulation by interfering with the action of the vitamin K–related factors (II, VII, IX, and X), which promote clotting.
2. Oral anticoagulants delay vitamin K formation and cause the PT to increase as a result of decreased factors II, VII, IX, and X.
3. The usual procedure is to run a PT test every day when beginning therapy. The anticoagulant dose is adjusted until the therapeutic range is reached. Then, weekly to monthly PT testing continues for the duration of therapy.
4. Coumadin takes 48 to 72 hours to cause a measurable change in the PT (3–4 days of drug therapy).

**Drug Therapy and PT Protocols**

1. Patients with cardiac problems are usually maintained at a PT level 2 to 2.5 times the normal (baseline) values.
2. Use of the INR values allows more sensitive control. A reasonable INR target for virtually all thromboembolic problems is 2.0 to 3.0. See Table 2.7 more specific guidelines.
3. For treatment of blood clots, the PT is maintained within the 2 to 2.5 times normal range. If the PT drops below this range, treatment may be ineffective, and old clots may expand or new clots may form. Conversely, if the PT rises above 30 seconds, bleeding or hemorrhage may occur.

**Clinical Implications**

1. Conditions that cause increased PT include:
a. Deficiency of factors II (prothrombin), V, VII, or X
b. Vitamin K deficiency, newborns of mother with vitamin K deficiency
c. Hemorrhagic disease of the newborn
d. Liver disease (eg, alcoholic hepatitis), liver damage
e. Current anticoagulant therapy with warfarin (Coumadin)
f. Biliary obstruction
g. Poor fat absorption (eg, sprue, celiac disease, chronic diarrhea)
h. Current anticoagulant therapy with heparin
i. DIC
j. Zollinger-Ellison syndrome
k. Hypofibrinogenemia (factor I deficiency), dysfibrinogenemia
l. Circulating anticoagulants, lupus anticoagulant
m. Premature newborns

2. Conditions that do not affect the PT include:
   a. Polycythemia vera
   b. Tannin disease
   c. Christmas disease (factor IX deficiency)
   d. Hemophilia A (factor VIII deficiency)
   e. von Willebrand’s disease
   f. Platelet disorders (idiopathic thrombocytopenic purpura)

Interfering Factors

1. Diet: ingestion of excessive green, leafy vegetables increases the body's absorption of vitamin K, which promotes blood clotting.
2. Alcoholism or excessive alcohol ingestion prolongs PT levels.
3. Diarrhea and vomiting decrease PT because of dehydration.
4. Quality of venipuncture: PT can be shortened if technique is traumatic and tissue thromboplastin is introduced to the sample and if collection tube is not filled properly.
5. Influence of prescribed medications: antibiotics, aspirin, cimetidine, isoniazid, phenothiazides, cephalosporins, cholesteramines, phenylbutazone, metronidazole, oral hypoglycemics, phenytoin
6. Prolonged storage of plasma at 4°C—activates factor VII and shortens PT

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, and need for frequent testing. Emphasize the need for regular monitoring through frequent blood testing if long-term therapy is prescribed. Do not refer to anticoagulants as "blood thinners." One explanation might be, “Your blood will be tested periodically to determine the pro time, which is an indication of how the blood clots.” The anticoagulant dose will be adjusted according to PT results.
2. Caution against self-medication. Ascertain what drugs the patient has been taking. Many drugs, including over-the-counter medications, alter the effects of anticoagulants and the PT value. Aspirin, acetaminophen, and laxative products should be avoided unless specifically ordered by the physician.
3. Instruct the patient never to start or discontinue any drug without the doctor's permission. This will affect PT values and may also interfere with the healing process.
4. Counsel regarding diet. Excessive amounts of green, leafy vegetables (eg, spinach, broccoli) will increase vitamin K levels and could interfere with anticoagulant metabolism. Caution against using razor blades; electric shavers should be used.
5. Remember that these guidelines also apply to aftercare.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately with follow-up testing and observation.
2. Avoid intramuscular injections during anticoagulant therapy because hematomas may form at the injection site. As the PT increases to upper limits (>30 seconds), assess carefully for bleeding from different areas; this may require neurologic assessment (if cranial bleeding is suspected), lung assessment and auscultation, gastrointestinal and genitourinary assessments, or other assessments as appropriate. Instruct the patient to observe for bleeding from gums, blood in the urine, or other unusual bleeding. Advise that care should be exercised in all activities to avoid accidental injury.
3. Alert patients who are being monitored by PT for long-term anticoagulant therapy that they should not take any other drugs unless they have been specifically prescribed.
4. Remember that when unexpected adjustments in anticoagulant doses are required to maintain a stable PT, or when there are erratic changes in PT levels, a drug interaction should be suspected and further investigation should take place.
5. Make changes in exercise intensity gradually or not at all. Active sports and contact sports should be avoided because of the potential for injury.
6. See Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

1. Critical value: If P/C is >2.5 or >30.0 seconds, notify clinician (for patients on anticoagulant).
2. If PT is excessively prolonged (>30 seconds), vitamin K may be ordered.
3. Critical value: >20 seconds (for nonanticoagulated persons)
4. Baseline PT levels should be determined before anticoagulant administration.
5. Critical value: INR >3.6; notify clinician (for patients on anticoagulants)
Coagulant Factors (Factor Assay)

Assay of specific factors of coagulation is done in the investigation of inherited and acquired bleeding disorders. For example, tests of factor VIII–related antigen are used in the differential diagnosis of classic hemophilia and von Willebrand's disease in cases in which there is no family history of bleeding and bleeding times are borderline or abnormal. A test for ristocetin cofactor is done to help diagnose von Willebrand's disease by determining the degree or rate of platelet aggregation that is taking place.

Reference Values

| Normal | Factor II—prothrombin: 80%–120% of normal Factor V—labile factor: 50%–150% of normal Factor VII—stable factor: 65%–140% of normal or 65–135 AU Factor VIII—antihemophilic factor: 55%–145% of normal or 55–145 AU Factor IX—Christmas factor: 60%–140% of normal or 60–140 AU Factor X: 45%–155% of normal or 45–155 AU Factor XI: 65%–135% of normal or 65–135 AU Factor XII—Hageman factor: 50%–150% of normal or 50–150 AU Ristocetin (von Willebrand's factor): 45%–140% of normal or 45–140 AU Factor VIII antigen: 100 µg/L or 50%–150% of normal or 50–150 AU Factor VIII–related antigen: 45%–185% of normal or 45–185 AU Fletcher's factor (prekallikrein): 80%–120% of normal or 0.80–1.20 Critical value for any coagulation factor: <10% of normal |

Procedure

1. Draw a 5-mL venous blood sample by the two-tube method and add to a collection tube containing sodium citrate as the anticoagulant.
2. Cap samples, put on ice, and send to the laboratory as soon as possible.

Clinical Implications

1. Inherited deficiencies:
   a. Any of the specific factors—I, II, V, VII, IX, X, XI, XII, and XIII—may be deficient on a familial basis.
   b. Factor VII is decreased in hypoproconvertinemia (autosomal recessive).
   c. Factor VIII is decreased in classic hemophilia A and von Willebrand's disease (inherited autosomally).
   d. Factor IX is decreased in Christmas disease or hemophilia B (sex-linked recessive).
   e. Factor XI is decreased in hemophilia C (autosomal dominant, occurring predominantly in Jews).
2. Acquired disorders:
   a. Factor II is decreased in:
      1. Liver disease
      2. Vitamin K deficiency
      3. Oral anticoagulants (last factor to decrease after starting Coumadin therapy)
      4. Normal newborns
      5. Circulating inhibitors or lupus-like anticoagulants
   b. Factor V is decreased in:
      1. Liver disease
      2. Factor V inhibitors
      3. Myeloproliferative disorders
      4. DIC and fibrinolysis
      5. Normal newborns (mildly decreased)
   c. Factor VII is decreased in:
      1. Liver disease
      2. Treatment with coumarin-type drugs (first factor to decrease)
      3. Normal newborns
      4. Kwashiorkor
   d. Factor VIII is increased in:
      1. Late normal pregnancy
      2. Thromboembolic conditions
      3. Liver disease
      4. Postoperative period
      5. Rebound activity after sudden cessation of a coumarin-type drug
      6. Normal newborns
   e. Factor VIII is decreased in:
      1. Presence of factor VIII inhibitors (anticoagulants capable of specifically neutralizing a coagulation factor and thereby disrupting hemostasis), associated with hemophilia A and immunologic reactions, and postpartum
      2. von Willebrand's disease
      3. DIC, fibrinolysis
      4. Myeloproliferative disorders
   f. Factor IX is decreased in:
      1. Uncompensated cirrhosis, liver disease
      2. Nephrotic syndrome
      3. Development of circulating anticoagulants against factor IX (rare)
      4. Normal newborns
      5. Dicumarol and related anticoagulant drugs
      6. DIC
      7. Vitamin K deficiency
   g. Factor X is decreased in:
      1. Vitamin K deficiency
      2. Liver disease
      3. Oral anticoagulants
      4. Amyloidosis
      5. DIC
6. Normal newborns

h. Factor XI is decreased in:
   1. Liver disease
   2. Intestinal malabsorption (vitamin K)
   3. Occasional development of circulatory anticoagulants against factor IX
   4. DIC
   5. Newborns (do not reach adult levels for up to 6 months)

i. Factor XII is decreased in:
   1. Nephrotic syndrome
   2. Liver disease
   3. Chronic granulocytic leukemia
   4. Normal newborns

j. Factor XIII is decreased in:
   1. Postoperative patients
   2. Liver disease
   3. Persistent increased fibrinogen levels
   4. Obstetric complications with hypofibrinogenemia
   5. Acute myelogenous leukemia
   6. Circulating anticoagulants
   7. DIC

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Plasminogen (Plasmin; Fibrinolysin)

Plasminogen is a glycoprotein, synthesized in the liver, present in plasma. Under normal circumstances, plasminogen is a part of any clot because of the tendency of fibrin to absorb plasminogen from the plasma. When plasminogen activators perform their function, plasmin is formed within the clot; this gradually dissolves the clot while leaving time for tissue repair. Free plasmin also is released to the plasma; however, antiplasmins there immediately destroy any plasmin released from the clot (see Fig. 2.2).

This test is done to determine plasminogen activity in persons with thrombosis or DIC. When pathologic coagulation processes are involved, excessive free plasmin is released to the plasma. In these situations, the available antiplasmin is depleted, and plasmin begins destroying components other than fibrin, including fibrinogen, factors V and VIII, and other factors. Plasmin acts more quickly to destroy fibrinogen because of fibrinogen's instability.

For therapeutic destruction of thrombi, urokinase, a trypsin-like protease purified from urine, may be administered to a patient to activate plasminogen to plasmin and induce fibrinolysis. Streptokinase is another therapeutic agent used for the same purpose.

Reference Values

Normal Plasminogen activity Males: 76%–124% of normal for plasma or 0.76–1.24 fraction of normal Females: 65%–153% of normal for plasma or 0.65–1.53 fraction of normal Infants: 27%–59% of normal for plasma or 0.27–0.59 fraction of normal

Procedure

1. Add a 5-mL venous blood sample to a collection tube containing sodium citrate. Use the two-tube method. Place the specimen in a biohazard bag.
2. Put the sample on ice and transport to the laboratory immediately.
3. Be aware that the test must be started within 30 minutes after the blood is drawn.

Clinical Implications

1. Decreased plasminogen activity occurs in:
   a. Some familial or isolated cases of idiopathic deep vein thrombosis
   b. DIC and systemic fibrinolysis
   c. Liver disease and cirrhosis
   d. Neonatal hyaline membrane disease
   e. Therapy with plasminogen activators
2. Decreased levels of plasminogen or abnormally functioning plasminogen can lead to venous and arterial clotting (thrombosis).
3. Increased plasminogen activity occurs in:
   a. Pregnancy (third trimester)
   b. Regular vigorous physical exercise

Interfering Factors See Appendix J for drugs that affect test outcomes.
Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately for thrombotic tendency.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Fibrinolysis (Euglobulin Lysis Time; Diluted Whole Blood Clot Lysis)

Primary fibrinolysis, without any sign of intravascular coagulation, is extremely rare. Secondary fibrinolysis is usually seen and follows or occurs simultaneously with intravascular coagulation. This secondary fibrinolysis is a protective mechanism against generalized clotting.

This test is done to evaluate a fibrinolytic activity. Shortened time indicates excessive fibrinolytic activity. Lysis is marked and rapid with primary fibrinolysis but can be minimal in secondary fibrinolysis. The diluted whole blood is used to monitor urokinase and streptokinase therapy.

Reference Values

Normal Euglobulin lysis—no lysis of plasma clot at 37°C in 60–120 minutes. The clot is observed for 24 hours. Diluted whole blood clot lysis: no lysis of clot in 120 minutes at 37°C.

Procedure

1. Collect a 5-mL venous blood sample in a tube containing sodium citrate using the two-tube method. Place the specimen in a biohazard bag.
2. Put the sample on ice and transport to the laboratory immediately, or start at bedside.
3. Be aware that the test must be started within 90 minutes after the blood is centrifuged.

Clinical Alert

A lysis time <1 hour signifies that abnormal fibrinolysis is occurring.

Clinical Implications

1. Increased fibrinolysis time occurs in the following conditions:
   a. Primary fibrinolysis
   b. Within 48 hours after surgery
   c. Cancer of prostate or pancreas
   d. Circulatory collapse, shock
   e. During lung and cardiac surgery
   f. Obstetric complications (eg, antepartum hemorrhage, amniotic embolism, septic abortion, death of fetus, hydatidiform mole)
   g. Long-term DIC (may be normal if plasminogen is depleted)
   h. Liver disease
      i. Administration of plasminogen activators (tPA, streptokinase, urokinase)
2. Heparin does not interfere with the euglobulin lysis test.

Interfering Factors

1. Increased fibrinolysis occurs with moderate exercise and increasing age.
2. Decreased fibrinolysis occurs in arterial blood, compared with venous blood. This difference is greater in arteriosclerosis (especially in young persons).
3. Decreased fibrinolysis occurs in postmenopausal women and in normal newborns.
4. FDPs interfere with fibrinolysis.
5. Normal results can occur if fibrinolysis is far advanced (plasminogen depleted).
6. Fibrinolysis is increased by very low fibrinogen levels (<80 mg/dL or <0.8 g/L) and decreased by high fibrinogen levels.
7. Increased fibrinolysis can be caused by traumatic venipuncture or a tourniquet that is too tight.
8. See Appendix J for drugs that affect test outcomes.

Interventions

Patient Preparation

1. Advise patient of test purpose and procedure; no exercise before test.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.
**Patient Aftercare**

1. Interpret test results and monitor appropriately for fibrinolytic crisis.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Fibrin Split Products (FSPs); Fibrin Degradation Products (FDPs)**

When fibrin is split by plasmin, positive tests for fibrin degradation (split) products, identified by the letters X, Y, D, and E, are produced. These products have an anticoagulant action and inhibit clotting when they are present in excess in the circulation. Increased levels of FDPs may occur with a variety of pathologic processes in which clot formation and lysis occur. This test is done to establish the diagnosis of DIC and other thromboembolic disorders.

**Reference Values**

**Normal** Negative at 1:4 dilution or <10 µg/mL (<10 mg/L)

**Procedure**

1. Place a venous blood sample of at least 4.5 mL in a tube containing thrombin and an inhibitor of fibrinolysis (reptilase, aprotinin, and calcium). Place the specimen in a biohazard bag.
2. Be aware that blood must be completely clotted before the test is started for the test to be valid because fibrinogen is broken down into identical fragments. Therefore, no fibrinogen can be present when the test is done.

**Clinical Implications** Increased FSP and FDP is associated with any condition associated with DIC (see page 128 for examples) and in:

1. Primary fibrinolysis
2. Venous thrombosis
3. Thoracic and cardiac surgery or renal transplantation
4. Acute myocardial infarction
5. Pulmonary embolism
6. Carcinoma
7. Liver disease

**Interfering Factors**

1. Because all of the laboratory methods are sensitive to fibrinogen as well as FDP, it is essential that no unclotted fibrinogen be left in the serum preparation. False-positive reactions can result if any fibrinogen is present.
2. False-positive results occur with heparin therapy.
3. The presence of rheumatoid factor (rheumatoid arthritis) may cause falsely high FSP and FDP values.
4. See Appendix J for drugs that affect test outcomes.

**Interventions**

**Patient Preparation**

1. Explain test purpose and procedure.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Patient Aftercare**

1. Interpret test results and monitor appropriately for DIC and thrombosis.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

<table>
<thead>
<tr>
<th>Clinical Alert</th>
</tr>
</thead>
</table>
| 1. Patients with very high levels of FSP/FDP have blood that does not clot or clots poorly.  
2. Critical value: >40 µg/mL (>40 mg/L) |

**D-Dimer**

D-Dimers are produced by the action of plasmin on cross-linked fibrin. They are not produced by the action of plasmin on unclotted fibrinogen or FDPs and therefore are specific for fibrin. The presence of D-dimer confirms that both thrombin generation and plasmin generation have occurred.

This test is used in the diagnosis of DIC disease and to screen for venous thrombosis and acute myocardial infarction. The D-dimer test is more specific for DIC than are tests for FSPs. The test verifies in vivo fibrinolysis because D-dimers are produced only by the action of plasmin on cross-linked fibrin, not by the action of plasmin on unclotted fibrinogen. A positive D-dimer test is presumptive evidence for DIC but is not diagnostic.

**Reference Values**

**Normal** <250 µg/L or <1.37 nmol/L Qualitative: no D-dimer fragments present

**Procedure** A venous blood sample or 5 mL is collected into a tube containing sodium citrate and apotinin. Place the specimen in biohazard bag and return to lab immediately.
Clinical Implications

1. Increased D-dimer values are associated with:
   a. DIC (secondary fibrinolysis)
   b. Arterial or venous thrombosis (deep vein thrombosis)
   c. Renal or liver failure
   d. Pulmonary embolism
   e. Late in pregnancy, preeclampsia
   f. Myocardial infarction
   g. Malignancy, inflammation, and severe infection
2. D-Dimer values are increased with tPA anticoagulant therapy.

NOTE

D-Dimer analysis of spinal fluid can rapidly and accurately differentiate cases of subarachnoid hemorrhage (SAH) from a traumatic tap. Positive in SAH.

Interfering Factors

1. False-positive tests are obtained with high titers of rheumatoid factor.
2. False-positive D-dimer levels increase as the tumor marker CA-125 for ovarian cancer increases.
3. The D-dimer test will be positive in all patients after surgery or trauma.
4. False-positive results found in estrogen therapy, normal pregnancy

Interventions

Patient Preparation

1. Explain test purpose and procedure.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Patient Aftercare

1. Interpret test outcome and monitor appropriately for DIC or thrombin.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Fibrinopeptide A (FPA)

Fibrinopeptides A and B are formed by the action of thrombin on fibrinogen; therefore, the presence of FPA indicates that thrombin has acted on fibrinogen.

The measurement is the most sensitive assay done to determine thrombin action. FPA reflects the amount of active intravascular blood clotting; this occurs in a subclinical DIC, which is common in patients with leukemia of various types and may be associated with tumor progression. FPA elevations can occur without intravascular thrombosis, decreasing the value of a positive test.

Reference Values

Normal Male: 0.4–2.6 mg/mL Female: 0.7–31 mg/mL

Procedure

1. Collect a venous blood sample of 5 mL in special Vacutainer tube containing aprotinin EDTA and thrombin to prevent activation in vitro. Use a two-tube method of draining blood.
2. Draw the specimen in a prechilled tube and place immediately on ice.
3. Place the specimen in a biohazard bag. Clean venipuncture and gentle handling of specimen are required. The specimen must be transported to the lab within 30 minutes.

Clinical Implications

1. Increased FPA occurs in:
   a. DIC
   b. Leukemia of various types
   c. Venous thrombosis and pulmonary embolus
   d. Myocardial infarction
   e. Postoperative patients
   f. Patients with widespread solid tumors, malignancies
2. Decreased FPA occurs in:
   a. Clinical remission of leukemia achieved with chemotherapy
   b. Therapeutic heparinization

Interfering Factors

1. A traumatic venous puncture may result in falsely increased levels.
2. The biologic half-life (stable for 2 hours or more) imposes limitations on the interpretation of a negative FPA test.
DIC occurs commonly in association with death of tumor cells in acute promyelocytic leukemia. For this reason, heparin is used prophylactically and in association with the initiation of chemotherapy for promyelocytic leukemia. DIC occurs less commonly during the treatment of acute myelomonocytic leukemia and acute lymphocyte leukemia. Evidence of DIC should be sought in every patient with leukemia before initiation of treatment.

Interventions

**Patient Preparation**

1. Explain test purpose and procedure.
2. Avoid prolonged use of tourniquet.
3. See Chapter 1 guidelines for *safe, effective, informed pretest care*.

**Patient Aftercare**

1. Interpret test outcome and monitor appropriately for DIC and thrombosis.
2. See Chapter 1 guidelines for *safe, effective, informed posttest care*.
3. Resume normal activities.

**Prothrombin Fragment (F1 + 2)**

The prothrombin F1 + 2 fragment is liberated from the prothrombin molecule when it is activated by factor Xa to form thrombin. Thrombin may be rapidly inactivated by antithrombin III. The F1 + 2 fragment, however, has a half-life of about 1.5 hours, making it a useful marker for activated coagulation.

Prothrombin F1 + 2 is used to detect activation of the coagulation system before actual thrombosis occurs. It is used to identify patients with low-grade intravascular coagulation (DIC) and to judge the effectiveness of oral anticoagulant therapy. F1 + 2 levels may assist in the study of the hypercoagulable states and in the assessment of thrombotic risk.

**Reference Values**

*Normal* 7.4–102.9 µg/L or 0.2–2.78 nmol/L Levels rise slightly with age over 45 years.

**Procedure**

1. Draw a 5-mL sample of venous blood into a blue-topped (sodium citrate anticoagulant) Vacutainer.
2. Use the two-tube technique. (Some methods may use lithium heparin.)

**Clinical Implications**

*Increased* prothrombin F1 + 2 is found in:

1. DIC (early)
2. Congenital deficiencies of antithrombin III
3. Congenital deficiencies of protein S and protein C
4. Leukemias
5. Severe liver disease
6. Post–myocardial infarction

**NOTE**

Failure to achieve a reduction in prothrombin F1 + 2 levels during oral anticoagulant therapy, despite an adequately prolonged PT, suggests inadequate anticoagulation.

**Interfering Factors**

1. Levels will be high in the immediate postoperative period.
2. Decreased with oral anticoagulants (Coumadin)
3. Decreased in patients treated with AT-III

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Avoid prolonged use of tourniquet.
3. See Chapter 1 guidelines for *safe, effective, informed pretest care*.

**Posttest Patient Aftercare**

1. Interpret test outcome and monitor appropriately for DIC and thrombosis.
2. See Chapter 1 guidelines for *safe, effective, informed posttest care*.
3. Resume normal activities.

**Fibrin Monomers (Protamine Sulfate Test; Fibrin Split Products)**

A positive test result reflects the presence of fibrin monomers, indicative of thrombin activity and consistent with a diagnosis of intravascular coagulation. A negative result does not mean that intravascular coagulation is not present. A
positive result may also be seen in some cases of severe liver disease and in inflammatory disorders caused by accumulation of products of coagulation in the circulation.

The detection of fibrin monomers and early-stage FSPs in plasma is useful in the diagnosis of DIC. Heparin therapy does not interfere with this test.

Reference Values
Normal Negative; no fibrin monomer present

Procedure
1. Obtain a 5-mL venous blood sample anticoagulated with sodium citrate (blue-topped tube). The two-tube technique is used.
2. Place the specimen on ice and transport to the laboratory. The test must be performed within 1 hour after collection.

Clinical Implications
1. A positive test is indicative of DIC.
2. Patients with deep vein thrombosis occasionally have positive results.
3. The test may be positive in severe liver disease or metastatic cancer.

Interfering Factors False-positive results may occur in the following situations:
1. Traumatic venipuncture
2. During or immediately before menstruation
3. During streptokinase therapy (thrombolytic therapy)

Interventions
Pretest Patient Preparation
1. Explain test purpose and procedure. If possible, drain blood before heparin therapy is started.
2. Avoid prolonged use of tourniquet.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcome and monitor appropriately for DIC and thrombosis.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.
3. Resume normal activities.

Fibrinogen
Fibrinogen is a complex protein (polypeptide) that, with enzyme action, is converted to fibrin. The fibrin, along with platelets, forms the network for the common blood clot. Although it is of primary importance as a coagulation protein, fibrinogen is also an acute-phase protein reactant. It is increased in diseases involving tissue damage or inflammation.

This test is done to investigate abnormal PT, APTT, and TT and to screen for DIC and fibrin-fibrinogenolysis. It is part of a coagulation panel.

Reference Values
Normal 200–400 mg/dL or 2.0–4.0 g/L

Procedure
1. Obtain a 5-mL venous blood sample using the two-tube technique with a collection tube containing sodium citrate.
2. Place the specimen in a biohazard bag.

Clinical Implications
1. Increased fibrinogen values occur in:
   a. Inflammation and infections (rheumatoid arthritis, pneumonia, tuberculosis, streptomycin)
   b. Acute myocardial infarction
   c. Nephrotic syndrome
   d. Cancer, multiple myeloma, Hodgkin's disease
   e. Pregnancy, eclampsia
   f. Various cerebral accidents and diseases
2. Decreased fibrinogen values occur in:
   a. Liver disease
   b. DIC (secondary fibrinolysis)
   c. Cancer
   d. Primary fibrinolysis
   e. Hereditary and congenital hypofibrinogenemia
   f. Dysfibrinogenemia

Interfering Factors
1. High levels of heparin interfere with test results.
2. High levels of FSP and FDP cause low fibrinogen values.
3. Oral contraceptives cause high fibrinogen values.
4. Elevated AT-III may cause decreased fibrinogen.
5. See Appendix J for other drugs that affect test outcomes.

**Clinical Alert**

<100 mg/dL or 1.0 g/L—possible panic value, notify physician

1. Values <50 mg/dL or <0.5 g/L can result in hemorrhage after traumatic surgery.
2. Values >700 mg/dL or >7.0 g/L constitute a significant risk for both coronary artery and cerebrovascular disease.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Have the patient avoid aggressive muscular exercise before the test.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome and monitor appropriately for DIC and response to treatment. If fibrinogen is low, cryoprecipitate is the preferred product for therapeutic replacement.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Protein C (PC Antigen)**

Protein C, a vitamin K–dependent protein that prevents thrombosis, is produced in the liver and circulated in the plasma. It functions as an anticoagulant by inactivating factors V and VIII. Protein C is also a profibrinolytic agent (ie, it enhances fibrinolysis). The protein C mechanism therefore functions to prevent extension of intravascular thrombi. This test is used for evaluation of patients suspected of having congenital protein C deficiency. Resistance to protein C is caused by an inherited defect in the factor V gene (factor V Leiden) and causes significant risk for thrombosis. It is the underlying defect in up to 60% of patients with unexplained thrombosis and is the most common cause of pathologic thrombosis. If functional protein C is abnormal, a protein C resistance test should be performed.

This test evaluates patients with severe thrombosis and those with an increased risk or predisposition to thrombosis. Patients with partial protein C or partial protein S deficiency (heterozygotes) may experience venous thrombotic episodes, usually in early adult years. There may be deep vein thromboses, episodes of thrombophlebitis or pulmonary emboli (or both), and manifestations of a hypercoagulable state. Patients who are heterozygous may have type I protein C deficiency, with decreased protein C antigen, or type II deficiency, with normal protein C antigen levels but decreased functional activity.

**NOTE**

The protein S level should always be determined when a protein C test is ordered.

<table>
<thead>
<tr>
<th>NOTE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein C resistance (factor V Leiden) should be tested in all patients with abnormal protein C activity.</td>
</tr>
</tbody>
</table>

**Reference Values**

Normal Qualitative: 70%–150% or 0.70–1.50 of increased functional activity Quantitative: 60%–125% or 0.60–1.25 of normal PC antigen

**Procedure**

1. Anticoagulate a 5-mL venous blood sample with sodium citrate (blue-topped tube). The two-tube method is used.
2. Cap the specimen and place on ice.

**Clinical Implications**

1. Decreased protein C is associated with:
   a. Severe thrombotic complications in the neonatal period (neonatal purpura fulminans)
   b. Increased risk for venous thrombotic episodes
   c. Warfarin (Coumadin)-induced skin necroses (pathognomonic for protein C deficiency)
   d. DIC, especially when it occurs with cancer (presumably owing to consumption by cofactor thrombin-thrombomodulin catalyst activities)
   e. Thrombophlebitis and pulmonary embolism, especially in early adult years
   f. Other acquired causes of protein C deficiency include:
      1. Liver disease
      2. Acute respiratory distress syndrome
      3. L-Asparaginase therapy
      4. Malignancies
5. Vitamin K deficiency
2. A deficiency of protein C may also be congenital (35%–58%).

Clinical Alert

Homozygous protein C–deficient patients have absent or almost absent protein C antigen and usually succumb in infancy with the picture of purpura fulminans neonatalis, including lower extremity skin ecchymoses, anemia, fever, and shock.

Interfering Factors

1. Decreased protein C is found in the postoperative state.
2. Pregnancy or use of oral contraceptives decreases protein C.
3. A transient drop in protein C occurs with a high loading dose of warfarin (Coumadin).
4. Protein C decreases with age.
5. High doses of heparin decrease protein C.
6. Lipemic serum may interfere with the assay.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. Patient should be fasting.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately for thrombosis. In the case of a protein C deficiency, educate the patient concerning the symptoms and implications of the disease. The risk factors include obesity, oral contraceptives, varicose veins, infection, trauma, surgery, pregnancy, immobility, and congestive heart failure.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Protein S

Both protein S and protein C are dependent on vitamin K for their production and function. A deficiency of either one is associated with a tendency toward thrombosis. Protein S serves as a cofactor to enhance the anticoagulant effects of activated protein C. Slightly more than half of protein S is complexed with C4 binding protein and is inactive. Activated protein C in the presence of protein S rapidly inactivates factors V and VIII.

This test is done to differentiate acquired from congenital protein S deficiency. Congenital deficiency of protein S is associated with a high risk for thromboembolism. Acquired deficiency of protein S can be seen in various autoimmune disorders and inflammatory states owing to elevation of C4-binding protein. This protein forms an inactive complex with protein S. C4-binding protein levels should be determined in all patients who demonstrate a reduced level of protein S.

Reference Values

Normal
Males: 60%–130% or 0.60–1.30 of normal activity
Females: 50%–120% or 0.50–1.20 of normal activity
Newborns: 15%–50% or 0.15–0.50 of normal activity

Procedure

1. Anticoagulate a 5-mL venous blood sample with sodium citrate (blue-topped tube). The two-tube method is used.
2. Keep the specimen capped and on ice. Place in biohazard bag and take to laboratory immediately.

Clinical Implications

1. Decreased values are associated with protein S deficiency. Familial protein S deficiency is associated with recurrent thrombosis. Abnormal plasma distribution of protein S occurs in functional protein S deficiency. In type I, free protein S is decreased, although the level of total protein may be normal; in type II, total protein is markedly reduced.
2. Hypercoagulable-state acquired protein S deficiency is found in:
   a. Diabetic nephropathy
   b. Chronic renal failure caused by hypertension
   c. Cerebral venous thrombosis
   d. Coumarin-induced skin necrosis
   e. DIC
   f. Thrombotic thrombocytopenia purpura
   g. Acute inflammation

Interfering Factors The following factors cause decreased protein S:

1. Heparin therapy or specimen contaminated with heparin
2. Patient on unstable warfarin (Coumadin should be discontinued for 30 days for a true protein S determination)
3. Pregnancy
4. Contraceptives (oral)
5. First month of life
6. L-Asparaginase therapy

**NOTE**

This test is not useful in diagnosing DIC.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome and monitor appropriately for thrombotic tendency.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Antithrombin III (AT-III; Heparin Cofactor Activity)**

AT-III inhibits the activity of activated factors XII, XI, IX, and X as well as factor II. AT-III is the main physiologic inhibitor of activated factor X, on which it appears to exert its most critical effect. AT-III is a “heparin cofactor.” Heparin interacts with AT-III and thrombin, increasing the rate of thrombin neutralization (inhibition) but decreasing the total quantity of thrombin inhibited.

This test detects a decreased level of antithrombin that is indicative of thrombotic tendency. Only the test of functional activity gives a direct clue to thrombotic tendency. In some families, several members may have a combination of recurrent thromboembolism and reduced plasma antithrombin (30%–60%). A significant number of patients with mesenteric venous thrombosis have AT-III deficiency. It has been recommended that patients with such thrombotic disease be screened for AT-III levels to identify those patients who may benefit from coumarin anticoagulant prophylaxis rather than heparin therapy.

**Reference Values**

**Normal**

Functional assay
- Infants (1–30 days): 26%–61% or 0.26–0.61 (premature); 44%–76% or 0.44–0.76 (full-term)
- Adults and infants older than 6 months: 80%–120% or 0.80–1.20

Immunologic assay
- Adults and infants older than 6 months: 17–30 mg/dL or 170–300 mg/L

**Procedure**

1. Anticoagulate a venous blood sample (5 mL) with sodium citrate. Mix gently.
2. Use the two-tube method.
3. Place the sample on ice and transport to laboratory immediately.

**Clinical Implications**

1. *Increased AT-III values* are associated with:
   a. Acute hepatitis
   b. Renal transplantation
   c. Inflammation, patients with increased ESR
   d. Menstruation
   e. Use of warfarin (Coumadin) anticoagulant
   f. Hyperglobulinemia
2. *Decreased AT-III values* are associated with:
   a. Congenital deficiency (hereditary)
   b. Liver transplantation and partial liver removal, cirrhosis, nephrotic syndrome, liver failure
   c. DIC, fibrinolytic disorders (not diagnostically useful)
   d. Acute myocardial infarction
   e. Active thrombotic disease (deep vein thrombosis), thrombophlebitis
   f. Carcinoma, trauma, severe inflammations
   g. Pulmonary embolism
   h. Heparin failure (low levels of AT-III exhibit heparin resistance)
   i. Protein-wasting diseases

**Interfering Factors**

1. Antithrombin decreases after 3 days of heparin therapy.
2. Use of oral contraceptives interferes with the test (decreased values).
3. Results are unreliable in the last trimester of pregnancy and in the early postpartum period.
4. Decreased after surgery, prolonged bed rest.
5. Decreased in L-asparaginase therapy.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. See Chapter 1 guidelines for **safe, effective, informed pretest care.**

**Posttest Patient Aftercare**

1. Interpret test outcome and monitor appropriately for thrombotic tendency.
2. If patient has decreased levels of AT-III, coumarin anticoagulant would be used as a prophylaxis.
3. See Chapter 1 guidelines for **safe, effective, informed posttest care.**

**BIBLIOGRAPHY**


Freeman J, Rodgers BA: Lupus: A Patient Care Guide for Nurses and Other Health Professionals. Bethesda, MD, National Institutes of Health, National Institute of Arthritis and Musculoskeletal and Skin Diseases, 1999


Kjeldsberg C: Practical Diagnosis of Hematologic Disorders, revised ed. Chicago, ASCP Press, 1991


Leavelle D: Interpretive Data for Diagnostic Laboratory Tests. Rochester, MN, Mayo Clinic Laboratories, 2001


Turgeon ML: Clinical Hematology Theory & Procedures, 3rd ed. Philadelphia, Lippincott Williams & Wilkins, 1999

OVERVIEW OF URINE STUDIES
Urine Formation
Types of Urine Specimens

URINE TESTING
Chart 3.1 Urinary System and Related Tests
Laboratory Testing for Routine Urinalysis

Dipstick Testing

NOTE
COLLECTION OF URINE SPECIMENS

Single, Random Urine Specimen
Long-Term, Timed Urine Specimen (2-Hour, 24-Hour)

ROUTINE URINALYSIS (UA) AND RELATED TESTS

Urine Volume
Urine Specific Gravity (SG)

Urine Test for Osmolality

Urine Blood or Hemoglobin (Hb)
Urine Protein (Albumin), Qualitative and 24-Hour
Urine N2-Microglobulin
Urine Glucose (Sugar)
Urine Ketones (Acetone; Ketone Bodies)
Urine Nitrite (Bacteria)
Urine Leukocyte Esterase

Urine Bilirubin
Urine Urobilinogen, Random and Timed

MICROSCOPIC EXAMINATION OF URINE SEDIMENT
Clinical Alert
Urine Red Blood Cells and Red Blood Cell Casts
Urine White Blood Cells and White Blood Cell Casts
Urine Epithelial Cells and Epithelial Casts
Urine Hyaline Casts
Urine Granular Casts
Urine Waxy Casts or Broad Casts (Renal Failure Casts) and Fatty Casts
Urine Crystals

Urine Strips

URINE CHEMISTRY
Urine Pregnancy Test; Human Chorionic Gonadotropin (hCG) Test
Urine Estrogen, Total and Fractions (Estradiol [E2] and Estriol [E3]), 24-Hour Urine and Total Estrogen—Blood

URINE DRUG INVESTIGATION SPECIMENS

Chart 3.2 Common Urine Drug Tests

Witnessed Urine Sampling for Suspected Substance Abuse

TIMED URINE TESTS
Urine Chloride (Cl), Quantitative (24-Hour)
Urine Sodium (Na), Quantitative (24-Hour)
Urine Potassium (K), Quantitative (24-Hour) and Random
Urine Uric Acid, Quantitative (24-Hour)
Urine Calcium (Ca), Quantitative (24-Hour)
Urine Magnesium (Mg), Quantitative (24-Hour)
Urine Oxalate, Quantitative (24-Hour)
Urine Pregnanediol (24-Hour)

Urine Pregnanetriol (24-Hour)

Urine 5-Hydroxyindoleacetic Acid (5-HIAA) (24-Hour)

Urine Vanillylmandelic Acid (VMA); Catecholamines (24-Hour)

Urine Urinary Porphyrins and Porphobilinogens (24-Hour and Random); 7-Aminolevulinic Acid (ALA, 7ALA)

Urine Amylase Excretion and Clearance (Random, Timed Urine, and Blood)

Phenylketonuria (PKU); Urine Phenylalanine (Random Urine and Blood)

Phenylalanine Absorption (Timed Urine and Blood)

Urine Creatinine, Creatinine Clearance (Timed Urine and Blood)

Urine Cystine (Random and 24-Hour)

Urine Hydroxyproline (Timed Urine and Blood)

NOTE

Urine Lysozyme (Random, 24-Hour Urine, and Blood)

Urine Amino Acids, Total and Fractions (Random, 24-Hour Urine, and Blood)

BIBLIOGRAPHY
OVERVIEW OF URINE STUDIES

Urine Formation

Urine is continuously formed by the kidneys. It is actually an ultrafiltrate of plasma from which glucose, amino acids, water, and other substances essential to body metabolism have been reabsorbed. The physiologic process by which approximately 170,000 mL of filtered plasma is converted to the average daily urine output of 1200 mL is complex.

Urine formation takes place in the kidneys, two fist-sized organs located outside the peritoneal cavity on each side of the spine, at about the level of the last thoracic and first two lumbar vertebrae. The kidneys, together with the skin and the respiratory system, are the chief excretory organs of the body. Each kidney is a highly discriminatory organ that maintains the internal environment of the body by selective secretion or reabsorption of various substances according to specific body needs.

The main functional unit of the kidney is the nephron. There are about 1 to 1.5 million nephrons per kidney, each composed of two main parts: a glomerulus, which is essentially a filtering system, and a tubule through which the filtered liquid passes. Each glomerulus consists of a capillary network surrounded by a membrane called Bowman's capsule, which continues on to form the beginning of the renal tubule. The kidney's ability to clear waste products selectively from the blood while maintaining the essential water and electrolyte balances in the body is controlled in the nephron by renal blood flow, glomerular filtration, and tubular reabsorption and secretion.

Blood is supplied to the kidney by the renal artery and enters the nephron through the efferent arteriole. It flows through the glomerulus and into the efferent arteriole. The varying size of these arterioles creates the hydrostatic pressure difference necessary for glomerular filtration and serves to maintain glomerular capillary pressure and consistent renal blood flow within the glomerulus. (The smaller size of the efferent arteriole produces an increase in the glomerular capillary pressure, which aids in urine formation.)

As the filtrate passes along the tubule, more solutes are added by excretion from the capillary blood and secretions from the tubular epithelial cells. Essential solutes and water pass back into the blood through the mechanism of tubular reabsorption. Finally, urine concentration and dilution occur in the renal medulla. The kidney has the remarkable ability to dilute or concentrate urine, according to the needs of the individual, and to regulate sodium excretion. Blood chemistry, blood pressure, fluid balance, and nutrient intake, together with the general state of health, are key elements in this entire metabolic process.

Urine Constituents

In general, urine consists of urea and other organic and inorganic chemicals dissolved in water. Considerable variations in the concentrations of these substances can occur as a result of the influence of factors such as dietary intake, physical activity, body metabolism, endocrine function, and even body position. Urea, a metabolic waste product produced in the liver from the breakdown of protein and amino acids, accounts for almost half of the total dissolved solids in urine. Other organic substances include primarily creatinine and uric acid. The major inorganic solid dissolved in urine is chloride, followed by sodium and potassium. Small or trace amounts of many additional inorganic chemicals are also present in urine.

The concentrations of these inorganic compounds are greatly influenced by dietary intake, making it difficult to establish normal levels. Other substances found in urine include hormones, vitamins, and medications. Although they are not a part of the original plasma filtrate, the urine may also contain formed elements such as cells, casts, crystals, mucus, and bacteria. Increased amounts of these formed elements are often indicative of disease.

Types of Urine Specimens

During the course of 24 hours, the composition and concentration of urine changes continuously. Urine concentration varies according to water intake and pretest activities. To obtain a specimen that is truly representative of a patient's metabolic state, it is often necessary to regulate certain aspects of specimen collection, such as time of collection, length of collection period, patient's dietary and medicinal intake, and method of collection. It is important to instruct patients when special collection procedures must be followed. See Appendix A: Standard Precautions, Appendix B: Latex Precautions, and Appendix E: Guidelines for Specimen Transport for additional guidelines.

URINE TESTING

Urinalysis (UA) is an essential procedure for patients undergoing hospital admission or physical examination. It is a useful indicator of a healthy or diseased state and has remained an integral part of the patient examination. Two unique characteristics of urine specimens can account for this continued popularity:

1. Urine is a readily available and easily collected specimen.
2. Urine contains information about many of the body's major metabolic functions, and this information can be obtained by simple laboratory tests.

These characteristics fit in well with the current trends toward preventive medicine and lower medical costs. By offering an inexpensive way to test large numbers of people, not only for renal disease but also for the asymptomatic beginnings of conditions such as diabetes mellitus and liver disease, the UA can be a valuable metabolic screening procedure.

Should it be necessary to determine whether a particular fluid is actually urine, the specimen can be tested for its urea and creatinine content. Inasmuch as both of these substances are present in much higher concentrations in urine than in...
other body fluids, the demonstration of a high urea and creatinine content can identify a fluid as urine (Chart 3.1).

**Chart 3.1 Urinary System and Related Tests**

**Organs and Function**

The kidneys, ureter, bladder, and urethra compose the urinary system. Kidneys must be able to excrete dietary and waste products (not eliminated by other organs) through the urine. Urine is formed within the functional unit of the kidneys, the nephron, which consists of glomeruli and tubules.

**KIDNEY GLOMERULUS**

Formation of filtrate

**KIDNEY TUBULE**

Filtration

Secretion of waste products

Reabsorption of waste products needed by the body

Reabsorption of water, sodium chloride, bicarbonates, potassium, and calcium, among others

**KIDNEY: PELVIS, URETERS, AND BLADDER**

Excretion and storage of formed urine

Main urine constituents: water, urea, uric acid, creatinine, sodium, potassium, chloride, calcium, magnesium, phosphates, sulfates, and ammonia

**Examples of Selective Filtration, Reabsorption, and Excretion by the Urinary System**

<table>
<thead>
<tr>
<th>Constituent</th>
<th>Filtered (g/24 h)</th>
<th>Reabsorbed (g/24 h)</th>
<th>Excreted (g/24 h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium</td>
<td>540</td>
<td>537</td>
<td>3.3</td>
</tr>
<tr>
<td>Chloride</td>
<td>630</td>
<td>625</td>
<td>5.3</td>
</tr>
<tr>
<td>Bicarbonate</td>
<td>300</td>
<td>300</td>
<td>0.3</td>
</tr>
<tr>
<td>Potassium</td>
<td>28</td>
<td>24</td>
<td>3.9</td>
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<tr>
<td>Glucose</td>
<td>140</td>
<td>140</td>
<td>0.0</td>
</tr>
<tr>
<td>Urea</td>
<td>53</td>
<td>28</td>
<td>25</td>
</tr>
<tr>
<td>Creatinine</td>
<td>1.4</td>
<td>0.0</td>
<td>1.4</td>
</tr>
<tr>
<td>Uric acid</td>
<td>8.5</td>
<td>7.7</td>
<td>0.8</td>
</tr>
</tbody>
</table>

**Laboratory Testing for Routine Urinalysis**

First, the physical characteristics of the urine are noted and recorded. Second, a series of chemical tests is run. A chemically impregnated dipstick can be used for many of these tests. Standardized results can be obtained by processing the urine-touched dipstick through special automated instruments. Third, the urine sediment is examined under the microscope to identify the components of the urinary sediment.

**Dipstick Testing**

Although laboratory facilities allow for a wide range of urine tests, some types of tablet, tape, and dipstick tests are available for UA outside the laboratory setting. They can be used and read directly by patients and clinicians.

Similar in appearance to pieces of blotter paper on a plastic strip, dipsticks actually function as miniature laboratories. Chemically impregnated reagent strips (UA Chemstrip Screen) provide quick determinations of pH, protein, glucose, ketones, bilirubin, hemoglobin (blood), nitrite, leukocyte esterase, urobilinogen, and specific gravity. The dipstick is impregnated with chemicals that react with specific substances in the urine to produce color-coded visual results. The depth of color produced relates to the concentration of the substance in the urine. Color controls are provided against which the actual color produced by the urine sample can be compared. The reaction times of the impregnated chemicals are standardized for each category of dipstick; it is vital that color changes be matched to the control chart at the correct elapsed time after each stick is dipped into the urine specimen. Instructions that accompany each type of dipstick outline the procedure. When more than one type of test is incorporated on a single stick (eg, pH, protein, and glucose), the chemical reagents for each test are separated by a water-impermeable barrier made of plastic so that results do not become altered (Table 3.1). An example of a form used to record dipstick (UA Chemstrip Screen) testing results is shown on the following page.

**Table 3.1 Urine Testing by Dipstick/Reagent Strip**

<table>
<thead>
<tr>
<th>Measurement</th>
<th>False-Positive</th>
<th>False-Negative</th>
<th>Correlations with Other Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Possible Reaction Interference</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### pH
None
Runover from the protein pad may lower
Nitrite
Leukocytes
Microscopic examination
Blood
Nitrite
Leukocytes
Microscopic examination

### Protein
Highly alkaline urine, ammonium compounds (antisepsics), detergents
High salt concentration
Nitrite
Leukocytes
Microscopic examination

### Glucose
Peroxide, oxidizing detergents
Ascorbic acid, 5-HIAA, homogentisic acid, aspirin, levodopa, ketones, high specific gravity with low pH
Ketones

### Ketones
Levodopa, phenylketones
None
Glucose

### Blood
Oxidizing agents, vegetable and bacterial peroxidases
Ascorbic acid, nitrite, protein pH < 5.0, high specific gravity, captopril
Protein

### Bilirubin
Lodine, pigmented urine, indican
Ascorbic acid, nitrite
Urobilinogen

### Urobilinogen
Ehrlich-reactive compounds (Multistix), medication color
Nitrile, formalin
Bilirubin

### Nitrite
Pigmented urine on automated readers
Ascorbic acid, high specific gravity
Protein

### Leukocytes
Oxidizing detergents
Glucose, protein, high specific gravity, oxalic acid, gentamycin, tetracycline, cephalixin, cephalothin
Protein

### Specific gravity
Protein
Alkaline urine
None

In addition to dipsticks, reagent strips, tablets, and treated slides for special determinations such as bacteria, phenylketonuria (PKU), mucopolysaccharides, salicylate, and cystinuria are available for urine analysis.

**NOTE**

Tablets are becoming obsolete but are still used for certain tests, such as glucose and reducing agents.

**Procedure**

1. Use a fresh urine sample within 1 hour of collection or a sample that has been refrigerated; bring to room temperature and mix specimen.
2. Read or review directions for use of the reagent. Periodically check for changes in procedure.
3. Dip a reagent strip into well-mixed urine, then remove it, blot, and compare each reagent area on the dipstick with the corresponding color control chart within the established time frame. Correlate color comparisons as closely as possible using good lighting.

**Interfering Factors**

1. If the dipstick is kept in the urine sample too long, the impregnated chemicals in the strip might be dissolved and could produce inaccurate readings and values.
2. If the reagent chemicals on the impregnated pad become mixed, the readings will be inaccurate. To avoid this, blot off excess urine after withdrawing the dipstick from the sample.
Clinical Alert

1. Precise timing is essential. If the test is not timed correctly, color changes may produce invalid or false results.
2. When not in use, the container of dipsticks should be kept tightly closed and stored in a cool, dry environment. If the reagents absorb moisture from the air before they are used, they will not produce accurate results. A desiccant comes with the reagents and should be kept in the container.
3. Quality control protocols must be followed:
   a. The expiration date must be honored even if there is no detectable deterioration of strips.
   b. Bottles must be discarded 6 months after opening, regardless of expiration date.
   c. Known positive and negative (abnormal and normal) controls must be run for each new bottle of reagent strips when it is opened and whenever there is a question of deterioration.

COLLECTION OF URINE SPECIMENS

Standard UA specimens can be collected any time, whereas first morning, fasting, and timed specimens require collection at specific times of day. Patient preparation and education needs vary according to the type of specimen required (Table 3.2) and the patient’s ability to cooperate with specimen collection. Clear instructions and assessment of the patient's understanding of the process are key to a successful outcome. Assess the patient's usual urinating patterns and encourage fluid intake (unless contraindicated). Provide verbal and written directions for self-collection of specimens. Assess for presence of interfering factors: failure to follow collection instructions, inadequate fluid intake, certain medications, and patient use of illegal drugs may affect test results. Certain foods, or any type of food consumption in some instances, may also affect test results.

<table>
<thead>
<tr>
<th>Type of Specimen</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIRST MORNING SPECIMEN</td>
<td>Most concentrated, Free of dietary influences, Bladder-incubated, Best for nitrate, protein, pregnancy tests; microscopic examination; routine screening</td>
</tr>
<tr>
<td>SECOND (DOUBLE-VOIDED) SPECIMEN</td>
<td>The first morning specimen is discarded; the second specimen is collected and tested</td>
</tr>
<tr>
<td>POSTPRANDIAL</td>
<td>Used for glucose determination, diabetic monitoring</td>
</tr>
<tr>
<td>TIMED</td>
<td>Requires collection at certain time, Total specimen must be collected</td>
</tr>
<tr>
<td>TIMED 2-HOUR VOLUME</td>
<td>Used for urobilinogen determination, All urine saved for 2-hour period</td>
</tr>
<tr>
<td>TIMED 24-HOUR VOLUME</td>
<td>Necessary for accurate quantitative results, All urine saved for 24-hour period</td>
</tr>
<tr>
<td>CATHETER SPECIMEN</td>
<td>Clamp catheter 15 to 30 minutes before collection, Cleanse sample port with alcohol, Insert needle into sample port; after aspirating sample, transfer to specimen container</td>
</tr>
<tr>
<td>ALERT: Unclamp catheter</td>
<td></td>
</tr>
<tr>
<td>SUPRAPUBIC ASPIRATION</td>
<td>Sterile bladder urine</td>
</tr>
</tbody>
</table>

*Table 3.2 Collection of Urine Specimens*

*Place urine specimens in a biohazard bag.*

Single, Random Urine Specimen

This is the most commonly requested specimen. Because the composition of urine changes over the course of the day, the time of day when the specimen is collected may influence the findings. The first voided morning specimen is particularly valuable because it is usually more concentrated and therefore more likely to reveal abnormalities as well as
the presence of formed substances. It is also relatively free of dietary influences and of changes caused by physical activity because the specimen is collected after a period of fasting and rest.

Procedure

1. Instruct the patient to void directly into a clean, dry container or bedpan. Transfer the specimen directly into an appropriate container. Disposable containers are recommended. Women should always have a clean-catch specimen if a microscopic examination is ordered (see Chap. 7).
2. Collect specimens from infants and young children into a disposable collection apparatus consisting of a plastic bag with an adhesive backing around the opening that can be fastened to the perineal area or around the penis to permit voiding directly into the bag. The specimen bag is carefully removed, and the urine is transferred to an appropriate specimen container.
3. Cover all specimens tightly, label properly, and send immediately to the laboratory. Place the label on the cup, not on the lid.
4. Obtain a clean specimen using the same procedure as for bacteriologic examination (see Chap. 7) if a urine specimen is likely to be contaminated with drainage, vaginal discharge, or menstrual blood.
5. If a urine specimen is obtained from an indwelling catheter, it may be necessary to clamp off the catheter for about 15 to 30 minutes before obtaining the sample. Clean the specimen port (in the tubing) with antiseptic before aspirating the urine sample with a needle and syringe.
6. Observe standard precautions when handling urine specimens (see Appendix A).
7. If the specimen cannot be delivered to the laboratory or tested within 1 hour, it should be refrigerated or have an appropriate preservative added.

Interfering Factors

1. Feces, discharges, vaginal secretions, and menstrual blood will contaminate the urine specimen. A clean voided specimen must be obtained.
2. If the specimen is not refrigerated within 1 hour of collection, the following changes in composition may occur:
   a. Increased pH from the breakdown of urea to ammonia by urease-producing bacteria
   b. Decreased glucose from glycolysis and bacterial utilization
   c. Decreased ketones because of volatilization
   d. Decreased bilirubin from exposure to light
   e. Decreased urobilinogen as a result of its oxidation to urobilin
   f. Increased nitrite from bacterial reduction of nitrate
   g. Increased bacteria from bacterial reproduction
   h. Increased turbidity caused by bacterial growth and possible precipitation of amorphous material
   i. Disintegration of red blood cells (RBCs) and casts, particularly in dilute alkaline urine
   j. Changes in color caused by oxidation or reduction of metabolites

Long-Term, Timed Urine Specimen (2-Hour, 24-Hour)

Some diseases or conditions require a second morning specimen or a 2-hour or 24-hour urine specimen to evaluate kidney function accurately (see Table 3.2). Substances excreted by the kidney are not excreted at the same rate or in the same amounts during different periods of the day and night; therefore, a random urine specimen might not give an accurate picture of the processes taking place over a 24-hour period. For measurement of total urine protein, creatinine, electrolytes, and so forth, more accurate information is obtained from a long-term specimen. All urine voided in a 24-hour period is collected into a suitable receptacle; depending on the intended test, a preservative is added, the collection is kept refrigerated, or both (Table 3.3).

<table>
<thead>
<tr>
<th>Test Element and Purpose</th>
<th>Preservative</th>
<th>Specimen Handling and Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acid mucopolysaccharides inherited enzyme deficiency in infants with mental retardation or failure to thrive</td>
<td>20 mL toluene (add at start of collection)</td>
<td>Refrigerate during collection; include patient's age</td>
</tr>
<tr>
<td>Aldosterone (cause of hypertension)</td>
<td>1 g boric acid per 100 mL urine</td>
<td>Refrigerate</td>
</tr>
<tr>
<td>Amino acids, quantitative (aminocaiduria, screen for inborn errors of metabolism and genetic abnormalities)</td>
<td>None</td>
<td>Refrigerate during collection</td>
</tr>
<tr>
<td>Aminolevulinic acid (porphyria and lead poisoning)</td>
<td>25 mL of 50% acetic acid; for children &lt; 5 y, use 15 mL of 50% acetic acid</td>
<td>Refrigerate or ice; protect from light</td>
</tr>
<tr>
<td>Amylase (differentiates acute pancreatitis from other abdominal diseases)</td>
<td>None</td>
<td>Refrigerate during collection</td>
</tr>
<tr>
<td>Arsenic (arsenic poisoning—occupational exposure)</td>
<td>20 mL of 6N HNO₃ in a metal-free container</td>
<td>Refrigerate during collection</td>
</tr>
<tr>
<td>Cadmium (toxic levels, including occupational exposure)</td>
<td>20 mL of 6N HNO₃ in a metal-free container</td>
<td>Refrigerate during collection</td>
</tr>
</tbody>
</table>
Calcium, quantitative Sulkowitch (hypercalciuria as in hyperparathyroidism, hyperthyroidism, vitamin D toxicity, Paget's disease, osteolytic diseases, and renal tubular acidosis) 30 mL of 6N HCl Refrigerate during collection

Catecholamine fractions, urinary free catecholamines (measure adrenomedullary function, to diagnose pheochromocytoma) 25 mL of 50% acetic acid; for children < 5 y, use 15 mL of 50% acetic acid Refrigerate or freeze, pH 1–3

Chloride (electrolyte imbalance, dehydration, metabolic alkalosis) None Refrigerate during collection

Chromium (toxic levels, including occupational exposure) 20 mL of 6N HNO₃ in a metal-free container Refrigerate

Citrate/citric acid (renal disease) 10 g boric acid Refrigerate

Copper (Wilson's disease) 20 mL of 6N HNO₃ in a metal-free container Refrigerate during collection

Cortisol, free (hydrocortisone levels in adrenal hormone function) 30 mL of 6N HCl Refrigerate during collection

Creatinine (to evaluate disorders of kidney function) None Refrigerate during collection

Creatinine clearance (measures kidney function, primarily glomerular filtration) None Refrigerate during collection

Cystine, quantitative (to diagnose cystinuria, inherited disease characterized by bladder calculi) 20 mL of toluene Refrigerate during collection, pH 2–3; if not acidified—freeze

?-Aminolevulinic acid (porphyria and lead poisoning) 30 mL of 33% glacial acetic acid Protect from light; refrigerate during collection

Electrolytes, sodium, potassium (electrolyte imbalance) None, or 1.0 g boric acid Refrigerate

Estriol, estradiol (menstrual and fertility problems, male feminization characteristics, estrogen-producing tumors, and pregnancy) 1.0 g boric acid Refrigerate during collection

Estrogens, total, nonpregnancy or third trimester (estrogen levels for menstrual and fertility problems, pregnancy and estrogen-producing tumors) 1.0 g boric acid Refrigerate during collection

Follicle-stimulating/luteinizing hormone (gonadotropic hormones, FSH and LH to determine cause of gonadal deficiency) 1.0 g boric acid or none Store frozen

Glucose (glucosuria to screen, confirm, or monitor diabetes mellitus, rapid intestinal absorption) 1.0 g boric acid or NaF Store in dark bottle

Histamine (chronic myelogenous leukemia, carcinoids, polycythemia vera) None Refrigerate; freeze portion after collection

Homovanillic acid (to diagnose neuroblastoma, pheochromocytoma, ganglioblastoma) 15 mL of 50% acetate acid <5 yrs of age, 25 mL of 50% acetic acid >5 years of age, to maintain pH 2.0–4.0 Refrigerate during collection; freeze portion after collection

17-Hydroxycorticosteroids (adrenal function by measuring urine excretion of steroids to diagnose endocrine disturbances of the adrenal androgens, Cushing's, Addison's, and so forth) 1.0 g boric acid Refrigerate, pH 5–7; freeze portion after collection

5-Hydroxyindoleacetic acid, serotonin (carcinoid tumors) 5-HIAA 15 mL of 50% acetic acid <5 yrs of age, 25 mL of 50% acetic acid >5 years of age, to maintain pH 2.0–4.0 Refrigerate during collection; freeze portion after collection

Hydroxyproline, free (measures the free hydroxyproline [less than 10% normally]; rapid growth and increased collagen turnover) 10 mL 6N HCl per liter of urine, maintain pH < 3 Refrigerate during collection; store frozen

Hydroxyproline, total 24-hour collection (bone collagen reabsorption and the degree of bone destruction from bone tumors) 10 mL 6N HCl per liter of urine, maintain pH < 3 Refrigerate during collection; use gelatin-free and low-collagen diet

Immunofixation electrophoresis (measures immune status and competence by identifying monoclonal and particle protein band immunoglobulins) None Refrigerate

?- and ? chains quantitative, also in serum (monoclonal gammapathies, myeloma tumor burden) None Refrigerate

17-Ketogenic steroids (Porter-Silber and Cushing's syndrome, adenogenital syndrome) 1.0 g boric acid Do not refrigerate

17-Ketosteroid, fractions (adrenal and gonadal abnormalities) 1.0 g boric acid Do not refrigerate

Lead (lead poisoning and chelation therapy) 20 mL of 6N HNO₃ in a metal-free container Refrigerate
Lipase (acute pancreatitis and to differentiate pancreatitis from other abdominal disorders) None Refrigerate
Lysozyme, muramidase (to differentiate acute myelogenous or monocytic leukemia from acute lymphatic leukemia) None Refrigerate
Magnesium (magnesium metabolism, electrolyte status, and nephrolithiasis) 20 mL of 6N HCl in a metal-free container Refrigerate
Manganese (toxicity, parenteral nutrition) None Refrigerate during collection
Mercury (toxicity, industrial and dental overexposure; inorganic mercury) 20 mL of 6N HNO$_3$ in a metal-free container Refrigerate; pH 2 with nitric acid
Metanephrine, total (assays of catecholamines and vanillylmandelic acid; frequently to diagnose pheochromocytoma) 30 mL of 6N HCl pH 1–3
Metanephrine, fractions (to diagnose and monitor pheochromocytoma and ganglioneuroblastoma) 30 mL of 6N HCl, final pH < 3 Refrigerate; no caffeine before or during testing
Metanephrine, total (pheochromocytoma, children with neuroblastoma, ganglioneuroma) 25 mL of 50% acetic acid; for children <5 y, use 15 mL of 50% acetic acid; or 30 mL of 6N HCl Refrigerate; no caffeine before or during testing

MHPG (3-methoxy-4-hydroxyphenylglycol) (to classify bipolar manic depression for drug therapy) None Refrigerate, ship frozen
Microalbumin, 24-hour (diabetic nephropathy) None Refrigerate
Osmolality, 24-hour (diabetes insipidus, primary polypepsia) None Refrigerate
Oxalate (nephrolithiasis and inflammatory bowel diseases) 20 mL of 6N HCl Refrigerate; pH 2–3
Phosphorous, 24-hour (renal losses; hyperparathyroidism and hypoparathyroidism) Acid-washed, detergent-free container Refrigerate during collection; acidify after collection
Porphobilinogens None Refrigerate during collection; freeze a portion; protect from light
Porphyruins, quantitative (to diagnose porphyrias and lead poisoning) 5 g sodium carbonate (do not use sodium bicarbonate) Refrigerate; protect specimen from light
Porphyruins (to diagnose porphyrias and lead poisoning) None (preservative is added on receipt in laboratory) Refrigerate; protect specimen from light
Potassium, 24-hour (electrolyte imbalance, renal and adrenal disorders) None Refrigerate during collection
Pregnanediol, 24-hour (measures ovarian and placental function) Boric acid Refrigerate during collection
Pregnanetriol (adrenogenital syndrome) 25 mL of 50% acetic acid; for children <5 y, use 15 mL of 50% acetic acid Refrigerate during collection; pH 4–4.5 after receipt in laboratory
Protein electrophoresis, 24-hour None Refrigerate during collection
Protein, total (proteinuria, differential diagnosis of renal disease) None Refrigerate during collection
Selenium (nutritional deficiency, industrial exposure) None Refrigerate; transport entire specimen to laboratory
Sodium, 24-hour (electrolyte imbalance, acute renal failure, oliguria and hyponatremia, sodium excreted for diagnosis of renal and adrenal imbalances) None Refrigerate during collection
Substance abuse screen (specific drugs and alcohol involved in substance abuse) None Refrigerate or freeze
Thallium (thallium toxicity, occupational exposure) None Refrigerate
Thiocyanate (short-term nitroprusside therapy, cyanide poisoning) None Refrigerate during collection
Total protein (renal disease) None Refrigerate during collection
Urea nitrogen, 24-hour (kidney function, hyperalimentation) 10 g boric acid Refrigerate
Uric acid, 24-hour (uric acid metabolism in gout and renal calculus formation) None Refrigerate during collection
Urobilinogen (liver function and liver cell damage) 5 g sodium carbonate and 100 mL petroleum ether (do not use sodium bicarbonate) Refrigerate during collection; protect specimen from light; check with laboratory
Vanillylmandelic acid, quantitative (adrenomedular pheochromocytoma, hypertension) 15 mL of 50% acetic acid <5 yrs of age, 25 mL of 50% acetic acid >5 years of age Refrigerate, pH 1–3; protect from light
Zinc (industrial exposure, toxicity, nutritional, acrodermatitis enteropathies) 20 mL 6N HNO$_3$ in a metal-free container Refrigerate

Procedure
1. Ask the patient to void at the beginning of a 24-hour timed urine specimen collection (or any other timed specimen collection). Discard this first specimen, and note the time.

2. Mark the time the test begins and the time the collection should end on the container. As a reminder, it may be helpful to post a sign above the toilet (eg, “24-Hour Collection in Progress”), with the beginning and ending times noted.

3. Collect all urine voided over the next 24 hours into a large container (usually glass or polyethylene), and label it with the patient's name, the timeframe for collection, the test ordered, and other pertinent information. It is not necessary to measure the volume of individual voidings, unless specifically ordered.

4. Ask the patient to void 24 hours after the first voiding, to conclude the collection. Add urine from this last voiding to the specimen in the container.

5. Storage
   a. Keep nonrefrigerated samples in a specified area or in the patient's bathroom.
   b. Refrigerate the collection bottle immediately after the patient has voided or place it into an iced container if refrigeration is necessary.

**NOTE**

Because the patient may not always be able to void on request, the last specimen should be obtained as closely as possible to the stated end-time of the test.

**Special Considerations**

1. In a health care facility, responsibility for the collection of urine specimens should be specifically assigned.

2. When instructing a patient about 24-hour urine collections, make certain the patient understands that the bladder must be emptied just before the 24-hour collection starts and that this preliminary specimen must be discarded; then, all urine voided until the ending time is saved.

3. Do not predate and pretime requisitions for serial collections. It is difficult for some patients to void at specific times. Instead, mark the actual times of collection on containers.

4. Documentation of the exact times at which the specimens are obtained is crucial to many urine tests.

5. Instruct the patient to urinate as near to the end of the collection period as possible.

6. When a preservative is added to the collection container (eg, HCl preservative in 24-hour urine collection for vanillylmandelic acid [VMA]), the patient must take precautions against spilling the contents and receiving an acid burn. Instructions regarding spillage need to be provided before the test begins.

7. The preservative used is determined by the urine substance to be tested for. The laboratory usually provides the container and the proper preservative when the test is ordered. If in doubt, verify this with the laboratory personnel.

**Interfering Factors**

1. Failure of patient or attending personnel to follow the procedure is the most common source of error.
   a. The patient should be given both verbal and written instructions. If the patient is unable to comprehend these directions, a significant other should be instructed in the process.
   b. If required, the proper preservative must be used.

2. Instruct the patient to use toilet paper after transferring the urine to the 24-hour collection container. Toilet paper placed in the specimen decreases the actual amount of urine available and contaminates the specimen.

3. The presence of feces contaminates the specimen. Patients should void first and transfer the urine to the collection receptacle before defecating.

4. If heavy menstrual flow or other discharges or secretions are present, the test may have to be postponed, or an indwelling catheter may need to be inserted to keep the specimen free of contamination. In some cases, thorough cleansing of the perineal or urethral area before voiding may be sufficient. If in doubt, communicate with laboratory personnel and the patient's physician.

**Interventions**

**Pretest Patient Preparation**

Most 24-hour urine specimen collections start in the early morning at about 7:00 a.m. (0700). Instruct the patient to do the following:

1. Empty the bladder completely on awakening and then discard that urine specimen. Record the time the voided specimen is discarded and the time the test is begun.

2. Save all urine voided during the next 24 hours, including the first specimen voided the next morning.

3. Add the urine voided the next morning (as close to the ending time as possible) to the collection container. The 24-hour test is then terminated, and the ending time is recorded.

4. Use a urinal, wide-mouth container, special toilet device, bedpan, or the collection container itself to catch urine. It is probably easier for women to void into another wide-mouth receptacle first and then to transfer the entire specimen carefully to the collection bottle. Men may find it simpler to void directly into the 24-hour collection container.

5. It is most important that all urine be saved in the 24-hour container. Ideally, the container should be refrigerated or placed on ice.

6. Test results are calculated on the basis of a 24-hour output. Unless all urine is saved, results will not be accurate. Moreover, these tests are usually expensive, complicated, and necessary for the evaluation and treatment of the patient's condition.

7. If the laboratory requests an aliquot, record total amount, mix well, and aliquot the requested amount.

8. Always check with your laboratory as to the preservative needed—different laboratories may have different requirements.
Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

ROUTINE URINALYSIS (UA) AND RELATED TESTS

The process of UA determines the following properties of urine: color, odor, turbidity, specific gravity, pH, glucose, ketones, blood, protein, bilirubin, urobilinogen, nitrite, leukocyte esterase, and other abnormal constituents revealed by microscopic examination of the urine sediment. A 10-mL urine specimen is usually sufficient for conducting these tests (Table 3.4).

Table 3.4 Normal Values in Urinalysis

<table>
<thead>
<tr>
<th>General Characteristics and Measurements</th>
<th>Chemical Determinations</th>
<th>Microscopic Examination of Sediment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color: pale yellow to amber</td>
<td>Glucose: negative</td>
<td>Casts negative: occasional hyaline casts</td>
</tr>
<tr>
<td>Appearance: clear to slightly hazy</td>
<td>Ketones: negative</td>
<td>Red blood cells: negative or rare</td>
</tr>
<tr>
<td>Specific gravity: 1.005–1.025 with a normal fluid intake</td>
<td>Blood: negative</td>
<td>Crystals: negative (none)</td>
</tr>
<tr>
<td>pH: 4.5–8.0; average person has a pH of about 5 to 6</td>
<td>Protein: negative</td>
<td>White blood cells: negative or rare</td>
</tr>
<tr>
<td>Volume: 600–2,500 mL/24 h; average 1200 mL/24 h</td>
<td>Bilirubin: negative</td>
<td>Epithelial cells: few; hyaline casts 0–1/lpf</td>
</tr>
<tr>
<td></td>
<td>Urobilinogen: 0.5–4.0 mg/d</td>
<td>(low-power field)</td>
</tr>
<tr>
<td></td>
<td>Nitrate for bacteria:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>protein: negative</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Leukocyte esterase:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>negative</td>
<td></td>
</tr>
</tbody>
</table>

Urine Volume

Urine volume measurements are part of the assessment for fluid balance and kidney function. The normal volume of urine voided by the average adult in a 24-hour period ranges from 600 to 2500 mL; the typical amount is about 1200 mL. The amount voided over any period is directly related to the individual's fluid intake, the temperature and climate, and the amount of perspiration that occurs. Children void smaller quantities than adults, but the total volume voided is greater in proportion to their body size.

The volume of urine produced at night is <700 mL, making the day-to-night ratio approximately 2:1 to 4:1.

Urine volume depends on the amount of water excreted by the kidneys. Water is a major body constituent; therefore, the amount excreted is usually determined by the body's state of hydration. Factors that influence urine volume include fluid intake, fluid loss from nonrenal sources, variations in the secretion of antidiuretic hormone (ADH), and the necessity to excrete increased amount of solutes such as glucose or salts. Polyuria is marked increase of urine production. Oliguria is decreased urinary output. The extreme form of this process is anuria, a total lack of urine production.

Reference Values

Normal

600–2500 mL in 24 hours or 600–2500 mL/day

Procedure

1. Collect a 24-hour urine specimen and keep it refrigerated or on ice.
2. Record the exact collection starting time and collection ending time on the specimen container and in the patient's health care record.
3. Transfer the specimen container to the laboratory refrigerator when the collection is completed. Complete the proper forms and document accordingly.
4. Ascertain volume by measuring the entire urine amount in a graduated and appropriately calibrated pitcher or other receptacle. The total volume is recorded as urine volume in milliliters (cubic centimeters) per 24 hours.

Clinical Implications

1. Polyuria (increased urine output) with elevated blood urea nitrogen (BUN) and creatinine levels
   a. Diabetic ketoacidosis
   b. Partial obstruction of urinary tract
   c. Some types of tubular necrosis (aminoglycoside)
2. Polyuria with normal BUN and creatinine
   a. Diabetes mellitus and diabetes insipidus
   b. Neurotic states (compulsive water drinking)
   c. Certain tumors of brain and spinal cord
3. Oliguria (<200 mL in adults, or <15–20 mL/kg in children, per 24 hours)
a. Renal causes
1. Renal ischemia
2. Renal disease due to toxic agents (certain drugs are toxic to the renal system)
3. Glomerulonephritis
b. Dehydration caused by prolonged vomiting, diarrhea, excessive diaphoresis, or burns
c. Obstruction (mechanical) of some area of the urinary tract or system
d. Cardiac insufficiency
4. Anuria (<100 mL in 24 hours)
a. Complete urinary tract obstruction
b. Acute cortical necrosis (cortex of the kidney)
c. Glomerulonephritis (acute, necrotizing)
d. Acute tubular necrosis
e. Hemolytic transfusion reaction

Interfering Factors
1. Polyuria
   a. Intravenous glucose or saline
   b. Pharmacologic agents such as thiazides and other diuretics
   c. Coffee, alcohol, tea, caffeine
2. Oliguria
   a. Water deprivation, dehydration
   b. Excessive salt intake

Interventions

Pretest Patient Preparation
1. Explain the purpose and procedure of the test.
2. Withhold diuretics for 3 days before the test. Check with clinician.
3. Avoid excessive water (liquid) intake and excessive salt intake. Advise patients to avoid salty foods and added salt in the diet. Eliminate caffeine and alcohol. Determine the patient's usual liquid intake and request that intake not be increased beyond this daily amount during testing.
4. Follow guidelines in Chapter 1 for safe, effective, pretest care.

Posttest Patient Aftercare
1. Patient can resume normal fluid and dietary intake and medications, unless specifically ordered otherwise.
2. Interpret test outcomes and counsel appropriately.
3. Follow guidelines in Chapter 1 for safe, effective, posttest care.

Urine Specific Gravity (SG)

Specific gravity (SG) is a measurement of the kidneys' ability to concentrate urine. The test compares the density of urine against the density of distilled water, which has an SG of 1.000. Because urine is a solution of minerals, salts, and compounds dissolved in water, the SG is a measure of the density of the dissolved chemicals in the specimen. As a measurement of specimen density, SG is influenced by both the number of particles present and the size of the particles. Osmolality is a more exact measurement and may be needed in certain circumstances.

The range of urine SG depends on the state of hydration and varies with urine volume and the load of solids to be excreted under standardized conditions; when fluid intake is restricted or increased, SG measures the concentrating and diluting functions of the kidney. Loss of these functions is an indication of renal dysfunction.

Reference Values
Normal
Normal hydration and volume: 1.005–1.030 (usually between 1.010 and 1.025)
Concentrated urine: 1.025–1.030+
Dilute urine: 1.001–1.010
Infant < 2 years old: 1.001–1.018

Procedure
1. Test SG with the use of a multiple-test dipstick that has a separate reagent area for SG. An indicator changes color in relation to ionic concentration, and this result is translated into a value for SG.
2. Determine SG with a refractometer or total solids meter. The refractive index is the ratio of the velocity of light in air to the velocity of light in the test solution. A drop of urine is placed on a clear glass plate of the urinometer and another plate is pressed on top of the urine sample. The path of light is deviated when it enters the solution, and the degree of deviation (refraction) is directly proportional to the density of the solution.
3. The urinometer (hydrometer) is the least accurate method. It consists of a bulb-shaped instrument that contains a scale calibrated in SG readings. Urine (10–20 mL) is transferred into a small test tube–like cylinder, and the urinometer is floated in the urine. The SG is read off the urinometer at the meniscus level of the urine. This method is becoming obsolete owing to the ease of dipstick testing.
4. Specimen collection
   a. For regular UA testing, about 20 mL of a random sample is needed for testing (UA including SG).
   b. When a special evaluation of SG is ordered separately from the UA, the patient should fast for 12 hours before specimen collection.
Clinical Implications

1. **Normal SG**: SG values usually vary inversely with the amount of urine excreted (decreased urine volume = increased SG). However, this relationship is not valid in certain conditions, including:
   a. Diabetes—increased urine volume, increased SG
   b. Hypertension—normal volume, decreased SG
   c. Early chronic renal disease—increased volume, decreased SG

2. **Hyposthenuria** (low SG, 1.001–1.010) occurs in the following conditions:
   a. Diabetes insipidus (low SG with large urine volume). It is caused by absence or decrease of ADH, a hormone that triggers kidney absorption of water. Without ADH, the kidneys produce excessive amounts of urine that are not reabsorbed (sometimes 15–20 L/day).
   b. Glomerulonephritis (kidney inflammation without infection) and phlebitis (kidney inflammation with bacterial infection, but not in the acute type of this disease). SG can be low in glomerulonephritis, with decreased urine volume. Tubular damage affects the kidneys’ ability to concentrate urine.
   c. Severe renal damage with disturbance of both concentrating and diluting abilities of urine. The SG is low (1.010) and fixed (varying little from specimen to specimen); this is termed isosthenuria.

3. **Hypersthenuria** (increased SG, 1.025–1.035) occurs in the following conditions:
   a. Diabetes mellitus
   b. Nephrosis
   c. Excessive water loss (dehydration, fever, vomiting, diarrhea)
   d. Increased secretion of ADH and diuretic effects related to the stress of a surgical procedure
   e. Congestive heart failure
   f. Toxemia of pregnancy

Interfering Factors

1. Radiopaque x-ray contrast media, minerals, and dextran may cause falsely high SG readings on the refractometer. The reagent dipstick method is not affected by high-molecular-weight substances.
2. Temperature of urine specimens affects SG; cold specimens produce falsely high values using the hydrometer.
3. Highly buffered alkaline urine may also cause low readings (with dipsticks only).
4. Elevated readings may occur in the presence of moderate amounts of protein (100–750 mg/dl) or with patients receiving intravenous albumin.
5. Detergent residue (on specimen containers) can produce elevated SG results.
6. Diuretics and antibiotics cause high readings.
7. See Appendix J for drugs that affect test outcomes.

Intervention

**Pretest Patient Preparation**

1. Explain the purpose and procedure for urine collection.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes, counsel, and monitor appropriately for conditions associated with altered SG.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Urine Osmolality

Osmolality, a more exact measurement of urine concentration than SG, depends on the number of particles of solute in a unit of solution. More information concerning renal function can be obtained if serum and urine osmolality tests are run at the same time. The normal ratio between urine and serum osmolality is 3:1. A high urine-to-serum ratio is seen with concentrated urine. With poor concentrating ability, the ratio is low.

Whenever a precise measurement is indicated to evaluate the concentrating and diluting ability of the kidney, this test is done. Urine osmolality during water restriction is an accurate test of decreased kidney function. It is also used to monitor the course of renal disease; to monitor fluid and electrolyte therapy; to establish the differential diagnosis of hypernatremia, hyponatremia, and polyuria; and to evaluate the renal response to ADH.

**Reference Values**

**Normal**

24-hour specimen: 300–900 mOsm/kg of H₂O
Random specimen: 50–1200 mOsm/kg of H₂O

Urine-to-serum ratio: 1:1 to 3:1

**Procedure**

1. Tell patient that this is a 24-hour urine collection test.
2. For the 24-hour test, the patient voids at approximately 7:00 a.m. (0700). All of the urine voided is saved in a special 24-hour collection container kept on ice or refrigerated. A high-protein diet may be ordered.
3. At the end of the test, the specimen is labeled and sent to the laboratory.
4. Simultaneous determination of serum osmolality may be done. A high urine-to-serum ratio is seen with concentrated urine.

Clinical Implications
1. Osmolality is increased in:
   a. Prerenal azotemia
   b. Congestive heart failure
   c. Addison’s disease
   d. Syndrome of inappropriate ADH secretion (SIADH)
   e. Dehydration
   f. Amyloidosis
   g. Hyponatremia
2. Osmolality is decreased in:
   a. Acute renal failure
   b. Diabetes insipidus
   c. Hypokalemia
   d. Hypertension
   e. Primary polydipsia
   f. Hypercalcemia
   g. Compulsive water drinking (increased fluid intake)
3. Urine-to-serum ratio is:
   a. Increased in prerenal azotemia
   b. Decreased in acute tubular necrosis

Interfering Factors

1. Intravenous sodium administration
2. Intravenous dextrose and water administration

Interventions

Pretest Patient Preparation

1. Explain purpose and procedure of the test to the patient.
2. A normal diet is prescribed for 3 days before testing.
3. To increase sensitivity of the osmolality test, a high-protein diet may be ordered for 3 days before the test. No liquids with the evening meal and no food or liquids should be taken after the evening meal until collection. Check with your laboratory if the patient has diabetes.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Provide the patient with foods and fluids as soon as the last urine sample is obtained.
2. Interpret test outcomes and monitor appropriately.
3. Follow guidelines in Chapter 1 for safe, effective, posttest care.

Urine Appearance

The first observation made about a urine specimen is usually its appearance, which generally refers to the clarity of the specimen.

Cloudy urine signals a possible abnormal constituent, such as white blood cells (WBCs), RBCs, or bacteria. On the other hand, excretion of cloudy urine may not be abnormal because a change in urine pH can cause precipitation, within the bladder, of normal urinary components. Alkaline urine may appear cloudy because of phosphates; acid urine may appear cloudy because of urates.

Reference Values

Normal
Fresh urine is clear to slightly hazy.

Procedure

1. Observe the clarity of a fresh urine sample by visually examining a well-mixed specimen in front of a light source.
2. Use common terms to report appearance, including the following: clear, hazy, slightly cloudy, cloudy, turbid, and milky.
3. Document results. The degree of turbidity should correspond to the amount of material observed under the microscope.

Clinical Implications

1. Pathologic urines are often turbid or cloudy; however, normal urine can also appear cloudy.
2. Urine turbidity may result from urinary tract infections (UTIs).
3. Urine may be cloudy because of the presence of RBCs, WBCs, epithelial cells, or bacteria.

Interfering Factors

1. After ingestion of food, urates, carbonates, or phosphates may produce cloudiness in normal urine on standing.
2. Semen or vaginal discharges mixed with urine are common causes of turbidity.
3. Fecal contamination causes turbidity.
4. Extraneous contamination (e.g., talcum, vaginal creams, radiographic contrast media) can cause turbidity.
5. “Greasy” cloudiness may be caused by large amounts of fat.
6. Often, normal urine develops a haze or turbidity after refrigeration or standing at room temperature because of precipitation of crystals of calcium oxalate or uric acid.

Urine Color

The yellow color of urine is caused by the presence of the pigment urochrome, a product of metabolism that under normal conditions is produced at a constant rate. The actual amount of urochrome produced depends on the body's metabolic state, with increased amounts being produced in thyroid conditions and fasting states.

Urine specimens may vary in color from pale yellow to dark amber. Variations in the yellow color are related to the body's state of hydration. The darker amber color may be directly related to the urine concentration or SG.

Reference Values

Normal

The normal color of urine is pale yellow to amber.

Straw-colored urine is normal and indicates a low SG, usually < 1.010. (The exception may be a patient with an elevated blood glucose concentration, whose urine is very pale yellow but has a high SG.)

Amber-colored urine is normal and indicates a high SG and a small output of urine.

Procedure

Observe and record the color of freshly voided urine.

Clinical Implications

1. Almost colorless (straw-colored) urine:
   a. Large fluid intake
   b. Chronic interstitial nephritis
   c. Untreated diabetes mellitus
   d. Diabetes insipidus
   e. Alcohol and caffeine ingestion
   f. Diuretic therapy
   g. Nervousness

2. Orange-colored (amber) urine:
   a. Concentrated urine caused by fever, sweating reduced fluid intake, or first morning specimen
   b. Bilirubin (yellow foam when shaken)
   c. Carrots or vitamin A ingestion (large amounts)
   d. Certain urinary tract medications (e.g., phenazopyridine [Pyridium], nitrofurantoin)

3. Brownish-yellow or greenish-yellow urine may indicate bilirubin in the urine that has been oxidized to biliverdin (greenish foam when shaken).

4. Green urine:
   a. Pseudomonal infection
   b. Indican
   c. Chlorophyll

5. Pink to red urine:
   a. RBCs
   b. Hemoglobin, methemoglobin, oxyhemoglobin
   c. Myoglobin
   d. Porphyryns

6. Brown-black urine:
   a. RBCs oxidized to methemoglobin
   b. Methemoglobin
   c. Homogentisic acid (alkaptonuria)
   d. Melanin or melanogen
   e. Phenol poisoning (Lysol)

7. Smoky urine may be caused by RBCs.

8. Milky urine is associated with fat, cystinuria, many WBCs, or phosphates (not pathologic).

Interfering Factors

1. Normal urine color darkens on standing because of the oxidation of urobilinogen to urobilin. This decomposition process starts about 30 minutes after voiding.
2. Some foods cause changes in urine color:
   a. Beets turn the urine red.
   b. Rhubarb can cause brown urine.
3. Many drugs alter the color of urine:
   a. Cascara and senna laxatives in the presence of acid urine turn the urine reddish brown; in the presence of alkaline urine, they turn the urine red.
   b. Bright-yellow color in alkaline urine may be a result of riboflavin or phenazopyridine.
   c. Urine that darkens on standing may indicate antiparkinsonian agents such as levodopa (Sinemet).
   d. Black urine may be caused by cascara, chloroquine, iron salts (ferrous sulfate, ferrous fumarate, ferrous gluconate), metronidazole, nitrofurantoin, quinine, or senna.
   e. Blue urine may be caused by triamterene.
   f. Blue-green urine may be caused by amitriptyline, methylene blue, or mitoxantrone.
   g. Orange urine may be caused by heparin, phenazopyridine, rifampin, sulfasalazine, or warfarin.
   h. Red-pink urine may be caused by chloroxazone, daunorubicin, doxorubicin, heparin, ibuprofen, methyldopa, phenytoin, rifampin, or senna.
i. Pink to brown urine may be caused by laxatives.

j. Brown urine may be caused by chloroquine, furazolidone, or primaquine.

k. Green urine may be caused by indomethacin.

Interventions
Pretest Patient Preparation
Assess color of urine; instruct patient to monitor and to report abnormal urine colors.

<table>
<thead>
<tr>
<th>Clinical Alert</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. If the urine is a red color, do not assume drug causation. Check the urine for hemoglobin. Question the patient regarding hematuria and recent activity, injury, or infection. Sometimes, vigorous exercise can bring on hematuria.</td>
</tr>
<tr>
<td>2. Red urine that is negative for occult blood is an indication that porphyria may be present. Report at once and document test results.</td>
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<tr>
<td>3. Other grossly abnormal colors (eg, black, brown) should be documented and reported.</td>
</tr>
</tbody>
</table>

Posttest Patient Aftercare
1. Interpret abnormal urine colors and counsel appropriately.
2. Explain that follow-up testing may be needed.

Urine Odor
Normal, freshly voided urine has a faint odor owing to the presence of volatile acids. It is not generally offensive. Although not part of the routine UA, abnormal odors should be noted.

Reference Values
Normal
Fresh urine from most healthy persons has a characteristic aromatic odor.

Procedure
Smell the urine and record perceptions.

Clinical Implications
1. The urine of patients with diabetes mellitus may have a fruity (acetone) odor because of ketosis.
2. UTIs result in foul-smelling urine because of the presence of bacteria, which split urea to form ammonia.
3. The urine of infants with an inherited disorder of amino acid metabolism known as “maple syrup urine disease” smells like maple or burnt sugar.
4. Cystinuria and homocystinuria result in a sulfurous odor.
5. Oasthouse urine (Smith-Strang) disease causes an odor associated with the smell of a brewery (yeasts, hops).
6. In phenylketonuria, a musty, mousy smell may be evident.
7. Tyrosinemia is characterized by a cabbage-like or “fishy” urine odor.
8. Butyric/hexanoic acidemia produces a urine odor resembling that of sweaty feet.

Interfering Factors
1. Some foods, such as asparagus, produce characteristic urine odors.
2. Bacterial activity produces ammonia from the decomposition of urea, with its characteristic pungent odor.

Urine pH
The symbol $pH$ expresses the urine as a dilute acid or base solution and measures the free hydrogen ion ($H^+$) concentration in the urine; 7.0 is the point of neutrality on the pH scale. The lower the pH, the greater the acidity; the higher the pH, the greater the alkalinity. The pH is an indicator of the renal tubules’ ability to maintain normal hydrogen ion concentration in the plasma and extracellular fluid. The kidneys maintain normal acid-base balance primarily through reabsorption of sodium and tubular secretion of hydrogen and ammonium ions. Secretion of an acid or alkaline urine by the kidneys is one of the most important mechanisms the body has for maintaining a constant body pH.

Urine becomes increasingly acidic as the amount of sodium and excess acid retained by the body increases. Alkaline urine, usually containing bicarbonate-carbonic acid buffer, is normally excreted when there is an excess of base or alkali in the body.

The importance of urinary pH lies primarily in determining the existence of systemic acid-base disorders of metabolic or respiratory origin and in the management of urinary conditions that require the urine to be maintained at a specific pH.

Control of Urine pH
Control of urinary pH is important in the management of several diseases, including bacteriuria, renal calculi, and drug therapy in which streptomycin or methenamine mandelate is being administered.

1. Renal calculi
   a. Renal stone formation partially depends on the pH of urine. Patients being treated for renal calculi are frequently given diets or medication to change the pH of the urine so that kidney stones will not form.
   b. Calcium phosphate, calcium carbonate, and magnesium phosphate stones develop in alkaline urine. In such instances, the urine must be kept acidic (see Diet, number 4, below).
   c. Uric acid, cystine, and calcium oxalate stones precipitate in acid urines. Therefore, as part of treatment, the
urine should be kept alkaline (see Diet, number 4, below).

2. **Drug treatment**
   a. Streptomycin, neomycin, and kanamycin are effective for treating genitourinary tract infections, provided the urine is alkaline.
   b. During sulfa therapy, alkaline urine should help prevent formation of sulfonamide crystals.
   c. Urine should also be kept persistently alkaline in the presence of salicylate intoxication (to enhance excretion) and during blood transfusions.

3. **Clinical conditions**
   a. The urine should be kept acidic during treatment of UTI or persistent bacteriuria and during management of urinary calculi that develop in alkaline urine.
   b. An accurate measurement of urinary pH can be made only on a freshly voided specimen. If the urine must be kept for any length of time before analysis, it must be refrigerated.
   c. Highly concentrated urine, such as that formed in hot, dry environments, is strongly acidic and may produce irritation.
   d. During sleep, decreased pulmonary ventilation causes respiratory acidosis; as a result, urine becomes more acidic.
   e. Chlorothiazide diuretic administration causes acid urine to be excreted.
   f. Bacteria from a UTI or from bacterial contamination of the specimen produce alkaline urine. Urea is converted to ammonia.

4. **Diet**
   a. A vegetarian diet that emphasizes citrus fruits and most vegetables, particularly legumes, helps keep the urine alkaline. Alkaline urine after meals is a normal response to the secretions of hydrochloric acid in gastric juice ("alkaline tide").
   b. A diet high in meat and protein keeps the urine acidic.
   c. Cranberry juice is the only fruit that will maintain an acidic urine, and it has long been used as a remedy for minor UTIs.

**Reference Values**

**Normal**
The pH of normal urine can vary widely, from 4.6 to 8.0.
The average pH value is about 6.0 (acidic).

**Procedure**

1. Use reagent strips for a dipstick measurement. They produce a spectrum of color changes from orange to green-blue to identify pH ranges from 5.0 to 9.0.
2. Dip the reagent strip into a freshly voided urine specimen, and compare the color change with the standardized color chart on the bottle that correlates color results with pH values.
3. Maintenance of the urine at a consistent pH requires frequent urine pH testing.

**Clinical Implications** To be useful, the urine pH measurement must be used in conjunction with other diagnostic information. For example, in renal tubular necrosis, the kidney is not able to excrete a urine that is strongly acidic. Therefore, if the urine pH is 5.0, renal tubular necrosis is eliminated as a possible diagnosis.

1. **Acidic urine** (pH < 7.0) occurs in:
   a. Metabolic acidosis, diabetic ketosis, diarrhea, starvation, uremia
   b. UTIs caused by *Escherichia coli*
   c. Respiratory acidosis (carbon dioxide retention)
   d. Renal tuberculosis
   e. Pyrexia
2. **Alkaline urine** (pH > 7.0) occurs in:
   a. UTIs caused by urea-splitting bacteria (*Proteus and Pseudomonas*)
   b. Renal tubular acidosis, chronic renal failure
   c. Metabolic acidosis (vomiting)
   d. Respiratory alkalosis involving hyperventilation ("blowing off" carbon dioxide)
   e. Potassium depletion

**Interfering Factors**

1. With prolonged standing, the pH of a urine specimen becomes alkaline because bacteria split urea and produce ammonia.
2. Ammonium chloride and mandelic acid may produce acid urines.
3. Runover between the pH testing area and the highly acidic protein area on the dipsticks may cause alkaline urine to give an acidic reading.
4. Sodium bicarbonate, potassium citrate, and acetazolamide may produce alkaline urine.
5. Urine becomes alkaline after eating because of excretion of stomach acid; this is known as the "alkaline tide."

**Clinical Alert**
The pH of urine never reaches 9, either in normal or abnormal conditions. Therefore, a pH finding of 9 indicates that a fresh specimen should be obtained to ensure the validity of the UA.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and specimen collection procedure.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor patient appropriately (see Control of Urine pH).
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Urine Blood or Hemoglobin (Hb)

The presence of free hemoglobin in the urine is referred to as hemoglobinuria. Hemoglobinuria can be related to conditions outside the urinary tract and occurs when there is such extensive or rapid destruction (intravascular hemolysis) of circulating erythrocytes that the reticuloendothelial system cannot metabolize or store the excess free hemoglobin. The hemoglobin is then filtered through the glomerulus. Hemoglobinuria may also occur as a result of lysis of RBCs in the urinary tract.

When intact RBCs are present in the urine, the term hematuria is used. Hematuria is most closely related to disorders of the renal or genitourinary systems in which bleeding is the result of trauma or damage to these organs or systems.

This test detects RBCs, hemoglobin, and myoglobin in urine. Blood in urine is always an indicator of damage to the kidney or urinary tract.

The use of both a urine dipstick measurement and microscopic examination of urine provides a complete clinical evaluation of hemoglobinuria and hematuria. Newer forms of dipsticks contain a lysing reagent that reacts with occult blood and detects intact as well as lysed RBCs.

When urine sediment is positive for occult blood but no RBCs are seen microscopically, myoglobinuria can be suspected. Myoglobinuria is caused by excretion of myoglobin, a muscle protein, into the urine as a result of (1) traumatic muscle injury, such as may occur in automobile accidents, football injuries, or electric shock; (2) a muscle disorder, such as an arterial occlusion to a muscle or muscular dystrophy; (3) certain kinds of poisoning, such as carbon monoxide or fish poisoning; or (4) malignant hyperthermia related to administration of certain anesthetic agents. Myoglobin can be distinguished from free hemoglobin in the urine by chemical tests.

Reference Values

Normal
Negative/none

Procedure

1. Collect a fresh, random urine specimen.
   a. Hemoglobinuria (hemoglobin in urine)
      1. Dip reagent sticks into the urine; the color change on the dipstick correlates with a standardized color chart specifically used with that particular type of dipstick.
      2. The color chart indicates color gradients for negative, moderate, and large amounts of hemoglobin.
   b. Hematuria (RBCs in urine)
      1. This dipstick method allows detection of intact RBCs when the number is greater than 10 cells/µL of urine. The color change appears stippled on the dipstick.
      2. The degree of hematuria can be estimated by the intensity of the speckled pattern.

2. Centrifuge the urine sample and examine the sediment microscopically (see Microscopic Examination of Urine Sediment) to verify the presence of RBCs.
   a. Hemoglobinuria is suspected when no RBCs are seen or the number seen does not correspond to the degree of color on the dipstick.
   b. Myoglobinemia may be suspected if the urine is cherry-red, no RBCs are seen, and blood serum enzymes for muscle destruction are elevated.

Clinical Implications

1. Hematuria is found in:
   a. Acute UTI (cystitis)
   b. Lupus nephritis
   c. Urinary tract or renal tumors
   d. Urinary calculi (intermittent hematuria)
   e. Malignant hypertension
   f. Glomerulonephritis (acute or chronic)
   g. Pyelonephritis
   h. Trauma to kidneys
      i. Polycystic kidney disease
   j. Leukemia
   k. Thrombocytopenia
   l. Strenuous exercise
   m. Benign familial or recurrent hematuria (asymptomatic hematuria without proteinuria; other clinical and laboratory data are normal)
   n. Heavy smokers

2. Hemoglobinuria is found in:
   a. Extensive burns
   b. Transfusion reactions (incompatible blood products)
   c. Febrile intoxication
d. Certain chemical agents and alkaloids (poisonous mushrooms, snake venom)
e. Malaria
f. Bleeding resulting from operative procedures on the prostate (can be difficult to control, especially in the presence of malignancies)
g. Hemolytic disorders such as sickle cell anemia, thalassemia, and glucose-6-phosphate dehydrogenase deficiency
h. Paroxysmal hemoglobinuria (large quantities of hemoglobin appear in urine at irregular intervals)
i. Kidney infarction
j. Hemolysis occurring while the urine is in the urinary tract (RBC lysis from hypotonic urine or alkaline urine)
k. Fava bean sensitivity (causes severe hemolytic anemia)
l. Disseminated intravascular coagulation (DIC)
m. Strenuous exercise (“march hemoglobinuria”)

Clinical Alert

One of the early indicators of possible renal or urinary tract disease is the appearance of blood in the urine. This does not mean that blood will be present in every voided specimen, but in most cases of renal or urinary tract disease, occult blood will appear in the urine with a reasonable degree of frequency. Any positive test for blood should be rechecked on a new urine specimen. If blood still appears, the patient should be further evaluated.

Interfering Factors

1. Drugs causing a positive result for blood or hemoglobin include:
   a. Drugs toxic to the kidneys (eg, bacitracin, amphotericin)
   b. Drugs that alter blood clotting (warfarin [Coumadin])
   c. Drugs that cause hemolysis of RBCs (aspirin)
   d. Drugs that may give a false-positive result (eg, bromides, copper, iodides, oxidizing agents)
2. High doses of ascorbic acid or vitamin C may cause a false-negative result.
3. High SG or elevated protein reduces sensitivity.
4. Myoglobin produces a false-positive result.
5. Hypochlorites or bleach used to clean urine containers causes false-positive results.
6. Menstrual blood may contaminate the specimen and alter results.
7. Prostatic infections may cause false-positive results.
8. See Appendix J for a complete list of drugs that affect test outcomes.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure for urine specimen collection.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and explain possible need for follow-up testing.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Urine Protein (Albumin), Qualitative and 24-Hour

The presence of increased amounts of protein in the urine can be an important indicator of renal disease. It may be the first sign of a serious problem and may appear before any other clinical symptoms. However, there are other physiologic conditions (eg, exercise, fever) that can lead to increased protein excretion in urine. Also, there are some renal disorders in which proteinuria is absent.

In a healthy renal and urinary tract system, the urine contains no protein or only trace amounts. These consist of albumin (one third of normal urine protein is albumin) and globulins from the plasma. Because albumin is filtered more readily than the globulins, it is usually abundant in pathologic conditions. Therefore, the term albuminuria is often used synonymously with proteinuria.

Normally, the glomeruli prevent passage of protein from the blood to the glomerular filtrate. Therefore, the presence of protein in the urine is the single most important indication of renal disease. If more than a trace of protein is found persistently in the urine, a quantitative 24-hour evaluation of protein excretion is necessary.

Reference Values for 24-Hour Urine

Normal
Adult male: 10–140 mg/L or 1–14 mg/dL
Adult female: 30–100 mg/L or 3–10 mg/dL
Child: <10 years old: 10–100 mg/L or 1–10 mg/dL

Reference Values—Qualitative
Normal
Negative

Procedure
Qualitative Protein Collection

1. Collect a random urine sample in a clean container and test it as soon as possible.
2. Use a protein reagent dipstick and compare the test result color with the color comparison chart provided on the
reagent strip bottle. Protein can also be detected by turbidimetric methods using sulfosalicylic acid.

3. Test a new second specimen and investigate any interfering factors if one of these methods produces positive results. A 24-hour urine test may then be ordered for a quantitative measurement of protein.

24-Hour Urine Protein Collection

1. Label a 24-hour urine container with the name of the patient, the test, and the date and time the test is started.
2. Refrigerate the specimen as it is being collected.
3. See general instructions for 24-hour urine collection listed (see Long-Term, Timed Urine Specimen, page 171).
4. Record the exact starting and ending times for the 24-hour collection on the specimen container and on the patient's record. (The usual starting and ending times are 0700 to 0700.)

Clinical Implications

1. Proteinuria occurs by two main mechanisms: damage to the glomeruli or a defect in the reabsorption process that occurs in the tubules.
   a. Glomerular damage
      1. Glomerulonephritis, acute and chronic
      2. Systemic lupus erythematosus (SLE)
      3. Malignant hypertension
      4. Amyloidosis
      5. Diabetes mellitus
      6. Nephrotic syndrome
      7. Polycystic kidney disease
   b. Diminished tubular reabsorption
      1. Renal tubular disease
      2. Pyelonephritis, acute and chronic
      3. Cystinosis
      4. Wilson's disease
      5. Fanconi's syndrome (defect of proximal tubular function)
      6. Intersitial nephritis

2. In pathologic states, the level of proteinuria is rarely constant, so not every sample of urine is abnormal in patients with renal disease, and the concentration of protein in the urine is not necessarily indicative of the severity of renal disease.

3. Proteinuria may result from glomerular blood flow changes without the presence of a structural abnormality, as in congestive heart failure.

4. Proteinuria may be caused by increased serum protein levels.
   a. Multiple myeloma (Bence Jones protein)
   b. Waldenström's macroglobulinemia
   c. Malignant lymphoma

5. Proteinuria can occur in other nonrenal disease (“functional proteinuria”)
   a. Acute infection, septicemia
   b. Trauma, stress
   c. Leukemia, hematologic disorders
   d. Toxemia, preeclampsia of pregnancy
   e. Hyperthyroidism
   f. Vascular disease (hypertension), cardiac disease
   g. Renal transport rejection
   h. Central nervous system lesions
   i. Poisoning from turpentine, phosphorus, mercury, gold, lead, phenol, opiates, or other drugs
   j. Hereditary, sickle cell, oxalosis

6. Large numbers of leukocytes accompanying proteinuria usually indicate infection at some level in the urinary tract. Large numbers of both leukocytes and erythrocytes indicate a noninfectious inflammatory disease of the glomerulus. Proteinuria associated with pyelonephritis may have as many RBCs as WBCs.

7. Proteinuria does not always accompany renal disease.
   a. Pyelonephritis
   b. Urinary tract obstructions
   c. Nephrolithiasis
   d. Tumors
   e. Congenital malformations
   f. Renal artery stenosis

8. Proteinuria is often associated with the finding of casts on sediment examination because protein is necessary for cast formation.

9. Postural proteinuria results from the excretion of protein by some patients when they stand or move about. This type of proteinuria is intermittent and disappears when the patient lies down. Postural proteinuria occurs in 3% to 15% of healthy young adults. It is also known as orthostatic proteinuria.

Collecting the Specimen for Orthostatic Proteinuria

1. The patient is instructed to void at bedtime and to discard this urine.
2. The next morning, a urine specimen is collected immediately after the patient awakens and before the patient has been in an upright position for longer than 1 minute. This may involve the use of a bedpan or urinal.
3. A second specimen is collected after the patient has been standing or walking for at least 2 hours.
4. With postural proteinuria, the first specimen contains no protein, but the second one is positive for protein.
5. The urine looks microscopically normal; no RBCs or WBCs are apparent. Orthostatic proteinuria is considered a
benign condition and slowly disappears with time. Progressive renal impairment usually does not occur.

**Clinical Alert**

1. Proteinuria of > 2000 mg/24 hours in an adult or = 40 mg/24 hours in a child usually indicates a glomerular cause.
2. Proteinuria of > 3500 mg/24 hours points to a nephrotic syndrome.

**Interfering Factors for Qualitative Protein Test**

1. Because of renal vasoconstriction, the presence of a functional, mild, and transitory proteinuria is associated with:
   a. Strenuous exercise up to 300 mg/24 hours
   b. Severe emotional stress, seizures
   c. Cold baths, exposure to very cold temperatures
2. Increased protein in urine occurs in these benign states:
   a. Fever and dehydration (salt depletion)
   b. Non-immunoglobulin E food allergies
   c. Salicylate therapy
   d. In the premenstrual period and immediately after delivery
3. False or accidental proteinuria may occur because of a mixture of pus and RBCs in the urinary tract related to infections, menstrual or vaginal discharge, mucus, or semen.
4. False-positive results can occur from incorrect use and interpretation of the color reagent strip test.
5. Alkaline, highly buffered urine may produce false-positive results on the dipstick test.
6. Very dilute urine may give a falsely low protein value.
7. Certain drugs may cause false-positive or false-negative urine protein tests (see Appendix J).
8. Radiographic contrast agents may produce false-positive results with turbidimetric measurements.

**Interventions**

**Pretest Patient Preparation**

1. Instruct the patient about the purpose and procedure for collection of the 24-hour urine specimen. Emphasize the importance of compliance with the procedure.
2. Food and fluids are permitted; however, fluids should not be forced because very dilute urine can produce false-negative values.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and explain the possible need for follow-up testing (eg, urine differential/electrophoresis) and treatment (to prevent progression to renal failure).
2. See guidelines in Chapter 1 for safe, effective, informed posttest care.
3. See Chapter 8 for protein electrophoresis.

**Microalbuminuria/Albumin (24-Hour Urine)**

Microalbuminuria is an increase in urinary albumin that is below the detectable range of the standard protein dipstick test. It is not a different chemical form of albumin. Microalbuminuria occurs long before clinical proteinuria becomes evident.

This test allows for the routine detection of low concentrations of albumin in the urine. This test has become a standard for the screening, monitoring, and detection of deteriorating renal function in diabetic patients. Studies have shown that diabetic patients who progress to renal failure first excrete micro amounts of albumin and that, at this stage, intervening treatment can reverse the proteinuria and thus prevent progression to renal failure. This test is also used to monitor compliance of blood pressure control, glucose control, and protein restriction.

**Reference Values**

**Normal**

<30 mg/24 hours (<30 mg/day) or < 20 mg/L (10-hour collection)

**Procedure**

1. 24-hour: Same as for total urine protein
2. 10-hour: Overnight collection
   a. Last voiding before sleep (10:00 p.m.)
   b. Collect all urine at first morning voiding (8:00 a.m.)

These results approximate 24-hour collection.

**Clinical Implications**

Increased microalbuminuria is associated with:

1. Diabetes with early diabetic nephropathy
2. Hypertension—heart disease
3. Generalized vascular disease
4. Preeclampsia

**Interfering Factors**
1. Strenuous exercise
2. Hematuria (menses)
3. High-protein diet or high salt levels

Interventions

Pretest Patient Preparation The pretest care is the same as for 24-hour protein.

Posttest Patient Aftercare

1. Albumin excretion > 30 mg/24 hours or > 20 mg/L/10 hours indicates an abnormal excretion.
   a. Patient management should be reviewed.
   b. Patient compliance can be checked by glycosylated hemoglobin to determine further control.
2. Patients with borderline results should be assessed on more than one occasion before the significance of a given urine measurement is finally judged.
3. The posttest care is the same as for 24-hour total protein.

Urine β2-Microglobulin

β2-Microglobulin, an amino acid peptide component of the lymphatic human lymphocyte antibody (HLA) complex, is found on the outside of the plasma membrane. It is structurally related to the immunoglobulins.

This test measures β2-microglobulin, which is nonspecifically increased in inflammatory conditions and in active chronic lymphatic leukemia. It may be used to differentiate glomerular from tubular dysfunction. In glomerular disease, β2-microglobulin is increased in serum and decreased in urine, whereas in tubular disorders, it is decreased in serum and increased in urine. In aminoglycoside toxicity, β2-microglobulin levels become abnormal before creatinine levels begin to show abnormal values. Serum is also used to evaluate the prognosis of multiple myeloma.

Reference Values

Normal
Urine 24-hour specimen: <1 mg/day
Blood serum specimen: <2.7 µg/mL or <2.7 mg/L

Procedure

1. Collect a 24-hour urine specimen or a serum sample.
2. Keep the pH neutral or alkaline (pH > 6.0)
3. Freeze specimen if not analyzed immediately. Not stable at room temperature.

Clinical Implications

1. Increased urine β2-microglobulin occurs in:
   a. Renal tubular disorders (>50 mg/day)
   b. Heavy-metal poisoning (mercury, cadmium)
   c. Drug toxicity (aminoglycosides, cyclosporine)
   d. Fanconi's syndrome, Wilson's disease
   e. Pyelonephritis
   f. Renal allograft rejection
   g. Lymphoid malignancies associated with B-lymphocyte lineage
   h. Acquired immunodeficiency syndrome (AIDS) (can be used as a predictor of the progression to AIDS)
2. Increased serum β2-microglobulin occurs in:
   a. Multiple myeloma (associated with a poor survival prognosis)
   b. Renal dialysis
   c. Amyloidosis
   d. Viral infection

Interfering Factors

1. Acid urine—not stable, pH < 6.0
2. Certain antibiotics (eg, gentamicin, tobramycin)
3. Recent nuclear medicine scan
4. Increased synthesis in certain diseases (eg, Crohn's disease, hepatitis, sarcoidosis) decreases the usefulness of the blood serum test.
5. Random specimens are not recommended.

Interventions

Pretest Patient Preparation

1. Instruct patient regarding the purpose of and procedure for test.
2. See instructions for 24-hour urine collection (see Long-Term, Timed Urine Specimen, page 171).
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately.
2. See Chapter 1 guidelines for **safe, effective, informed posttest care**.

**Urine Glucose (Sugar)**

Glucose is present in glomerular filtrate and is reabsorbed by the proximal convoluted tubule. If the blood glucose level exceeds the reabsorption capacity of the tubules, glucose will appear in the urine. Tubular reabsorption of glucose is by active transport in response to the body’s need to maintain an adequate concentration of glucose. The blood level at which tubular reabsorption stops is termed the renal threshold, which for glucose is between 160 and 180 mg/dL (9–10 mmol/L).

**Types of Glucose Tests**

1. **Reduction tests** (Clinistest)
   a. These are based on reduction of cupric ions by glucose. When the compounds are added to urine, a heat reaction takes place. This results in precipitation and a change in the color of the urine if glucose is present.
   b. These tests are nonspecific for glucose because the reaction can also be caused by other reducing substances in the urine, including:
      1. Creatinine, uric acid, ascorbic acid
      2. Other sugars, such as galactose, lactose, fructose, pentose, and maltose
   c. These tests have a lower sensitivity than enzyme tests.

2. **Enzyme tests** (Clinistix, Diastix, Tes-Tape)
   a. These tests are based on interaction between glucose oxidase (an enzyme) and glucose. When dipped into urine, the enzyme-impregnated strip changes color according to the amount of glucose in the urine. The manufacturer's color chart provides a basis for comparison of colors between the sample and the manufacturer's control.
   b. These tests are specific for glucose only.

**Reference Values**

**Normal**
Random specimen: Negative
24-hour specimen: 1–15 mg/dL (60–830 µmol/L) or <0.5 g/24 hours (<2.8 mmol/day)

**Procedure**

1. Use a freshly voided specimen.
2. Follow directions on the test container exactly. Timing must be exact; the color reaction must be compared with the closest matching control color on the manufacturer's color chart to ascertain accurate results.
3. Record the results on the patient's record.
4. Refrigerate or ice the entire urine sample during collection if a 24-hour urine specimen is also ordered. See Table 3-3 for proper preservative.

**Clinical Alert**

1. Urine glucose > 1000 mg/dL (>55 mmol/L) (4+) is a critical value.
2. Determine exactly what drugs the patient is taking and whether the metabolites of these drugs can affect the urine glucose results. Frequent updating in regard to the effects of drugs on blood glucose levels is necessary in light of the many new drugs introduced and prescribed.
3. Test results may be reported as “plus” (+) or as percentages. Percentages are more accurate.
4. When screening for galactose (galactosuria) in infants, the reduction test must be used. The enzyme tests do not react with galactose.
5. Newborns should always be tested by both methods (reduction and enzymatic).

**Clinical Implications**

1. **Increased glucose** occurs in:
   a. Diabetes mellitus
   b. Endocrine disorders (thyrotoxicosis, Cushing's syndrome, acromegaly)
   c. Liver and pancreatic disease
   d. Central nervous system disorders (brain injury, stroke)
   e. Impaired tubular reabsorption
      1. Fanconi's syndrome
      2. Advanced renal tubular disease
   f. Pregnancy with possible latent diabetes (gestational diabetes)
2. **Increase of other sugars** (react only with reduction tests, not dipstick tests):
   a. Lactose—pregnancy, lactation, lactose intolerance
   b. Galactose—hereditary galactosuria (severe enzyme deficiency in infants; must be treated promptly)
   c. Xylose—excessive ingestion of fruit
   d. Fructose—hereditary fructose intolerance, hepatic disorders
   e. Pentose—certain drug therapies and rare hereditary conditions

**Interfering Factors**

1. Interfering factors for reduction test (false-positive results):
   a. Presence of non–sugar-reducing substances such as ascorbic acid, homogentisic acid, creatinine
   b. Tyrosine
   c. Nalidixic acid, cephalosporins, probenecid, and penicillin
d. Large amounts of urine protein (slows reaction)

2. Interfering factors for dipstick enzyme tests:
   a. Ascorbic acid (in large amounts) may cause a false-negative result
   b. Large amount of ketones may cause a false-negative result
   c. Peroxide or strong oxidizing agents may cause a false-positive result.

3. Stress, excitement, myocardial infarction, testing after a heavy meal, and testing soon after the administration of intravenous glucose may all cause false-positive results, most frequently trace reactions.

4. Contamination of the urine sample with bleach or hydrogen peroxide may invalidate results.

5. False-negative results may occur if urine is left to sit at room temperature for an extended period, owing to the rapid glycolysis of glucose.

6. High specific gravity depresses color development, low specific gravity intensifies it. See Appendix J for other drugs that affect test outcomes.

Interventions

Pretest Patient Preparation

1. Instruct the patient about the test purpose, the procedure, and the double-voiding technique.
   a. Discard the first voided morning specimen, then void 30 to 45 minutes later for the test specimen. This second specimen reflects the immediate state of glucosuria more accurately because the first morning specimen consists of urine that has been present in the bladder for several hours.
   b. Advise the patient not to drink liquids between the first and second voiding so as not to dilute the glucose present in the specimen.
   c. A urine glucose test combined with a blood glucose test gives a more complete assessment of diabetes.

2. Instruct the patient about the 24-hour urine collection procedure when applicable (see Long-Term, Timed Urine Specimen, page 171).

3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately.

2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

Urine glucose > 1000 mg/dL (>55 mmol/L)—test blood glucose, notify physician, and begin appropriate treatment.

Urine Ketones (Acetone; Ketone Bodies)

Ketones, which result from the metabolism of fatty acid and fat, consist mainly of three substances: acetone, β-hydroxybutyric acid, and acetoacetic acid. The last two substances readily convert to acetone, in essence making acetone the main substance being tested. However, some testing products measure only acetoacetic acid.

In healthy persons, ketones are formed in the liver and are completely metabolized so that only negligible amounts appear in the urine. However, when carbohydrate metabolism is altered, an excessive amount of ketones is formed (acidosis) because fat becomes the predominant body fuel instead of carbohydrates. When the metabolic pathways of carbohydrates are disturbed, carbon fragments from fat and protein are diverted to form abnormal amounts of ketone bodies. Increased ketones in the blood lead to electrolyte imbalance, dehydration, and, if not corrected, acidosis and eventual coma.

The excess presence of ketones in the urine (ketonuria) is associated with diabetes or altered carbohydrate metabolism. Some "fad" diets that are low in carbohydrates and high in fat and protein also produce ketones in the urine. Testing for urine ketones in patients with diabetes may provide the clue to early diagnosis of ketoadidosis and diabetic coma.

Indications for Ketone Testing

1. General: Screening for ketonuria is frequently done in hospitalized patients, presurgical patients, pregnant women, children, and persons with diabetes.

2. Glycosuria (diabetes):
   a. Testing for ketones is indicated in any patient showing elevated urine and blood sugars.
   b. When treatment is being switched from insulin to oral hypoglycemic agents, the development of ketonuria usually indicates a poor response to the oral hypoglycemic agents.
   c. The urine of diabetic patients treated with oral hypoglycemic agents should be tested regularly for glucose and ketones because oral hypoglycemic agents, unlike insulin, do not control diabetes when acute complications such as infection develop.
   d. Ketone testing is done to differentiate between diabetic coma–positive ketones and insulin shock–negative ketones.

3. Acidosis:
   a. Ketone testing is used to judge the severity of acidosis and to monitor the response to treatment.
   b. Urine ketone measurement frequently provides a more reliable indicator of acidosis than blood testing (it is especially useful in emergency situations).
   c. Ketones appear in the urine before there is any significant increase of ketones in the blood.

4. Pregnancy: During pregnancy, the early detection of ketones is essential because ketoacidosis is a prominent factor that contributes to intrauterine death.
Reference Values

Normal
Urine: Negative
Serum or plasma:
Acetone: <2.0 mg/dL or <0.34 mmol/L
Acetoacetate: <1 mg/dL or <0.1 mmol/L
β-hydroxybutyric acid: 0.21–2.81 mg/dL or 20–270 µmol/L

Procedure

1. Dip the ketone reagent strip in fresh urine, tap off excess urine, time the reaction accurately, and then compare the strip with the control color chart on the container.
2. Follow the manufacturer's directions exactly if procedure differs from the technique just described.
3. Do not use dipsticks to test for ketones in blood. Special testing products are designed for blood.

Clinical Implications

1. Ketosis and ketonuria may occur whenever increased amounts of fat are metabolized, carbohydrate intake is restricted, or the diet is rich in fats (either "hidden" or obvious). This state can occur in the following situations:
   a. Metabolic conditions
      1. Diabetes mellitus (diabetic acidosis)
      2. Renal glycosuria
      3. Glycogen storage disease (von Gierke's disease)
   b. Dietary conditions
      1. Starvation, fasting
      2. High-fat diets
      3. Prolonged vomiting, diarrhea
      4. Anorexia
      5. Low-carbohydrate diet
      6. Eclampsia
   c. Increased metabolic states caused by:
      1. Hyperthyroidism
      2. Fever
      3. Pregnancy or lactation
2. In nondiabetic persons, ketonuria occurs frequently during acute illness, severe stress, or strenuous exercise. Approximately 15% of hospitalized patients have ketones in their urine even though they do not have diabetes.
3. Children are particularly prone to developing ketonuria and ketosis.
4. Ketonuria occurs after anesthesia (ether or chloroform).

Interfering Factors

1. Drugs that may cause a false-positive result include:
   a. Levodopa
   b. Phenothiazines
   c. Ether
   d. Insulin
   e. Isopropyl alcohol
   f. Metformin
   g. Penicillamine
   h. Phenazopyridine (Pyridium)
   i. Captopril
2. False-negative results occur if urine stands too long, owing to loss of ketones into the air.
3. See Appendix J for other drugs that affect test outcomes.

Clinical Alert

Ketonuria signals a need for caution, rather than crisis intervention, in either a diabetic or a nondiabetic patient. However, this condition should not be taken lightly.

1. In the diabetic patient, ketone bodies in the urine suggest that the diabetes is not adequately controlled and that adjustments of either the medication or the diet should be made promptly.
2. In the nondiabetic patient, ketone bodies indicate a reduced carbohydrate metabolism and excessive fat metabolism.
3. Positive urine ketones in a child younger than 2 years of age is a critical alert.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.
Urine Nitrite (Bacteria)

This test is a rapid, indirect method for detecting bacteria in the urine. Significant UTI may be present in a patient who does not experience any symptoms. Common gram-negative organisms contain enzymes that reduce the nitrate in the urine to nitrite.

Clinicians frequently request the urine nitrate test to screen high-risk patients: pregnant women, school-aged children (especially girls), diabetic patients, elderly patients, and patients with a history of recurrent infections.

The majority of UTIs are believed to start in the bladder as a result of extreme contamination; if left untreated, they can progress upward all the way to the kidneys. Pyelonephritis is a frequent complication of untreated cystitis and can lead to renal damage. Detection of bacteria using the nitrate test and subsequent antibiotic therapy can prevent these serious complications. The nitrate test can also be used to evaluate the success of antibiotic therapy.

**Reference Values**

**Normal**

Negative for bacteria

**Procedure**

1. Obtain a first morning specimen because urine that has been in the bladder for several hours is more likely to yield a positive nitrate test than a random urine sample that may have been in the bladder for only a short time. A clean-catch (midstream) urine specimen is needed to minimize bacterial contamination from adjacent areas.
2. Follow the exact testing procedure according to prescribed guidelines for reliable test results. Any shade of pink is positive for nitrite-producing bacteria.
3. Compare the reacted reagent area on the dipstick with a white background to aid in the detection of a faint pink hue that might otherwise be missed.
4. Perform a microscopic examination to verify results, if at all possible.

**Clinical Implications**

1. Under the light microscope, the presence of > 20 bacteria per high-power field (hpf) may indicate a UTI. Untreated bacteriuria can lead to serious kidney disease.
2. The presence of a few bacteria suggests a UTI that cannot be confirmed or excluded until more definitive studies, such as culture and sensitivity tests, are performed. Again, this finding merits serious consideration for treatment.
3. A positive nitrate test is a reliable indicator of significant bacteriuria and is a cue for performing urine culture.
4. A negative result should never be interpreted as indicating absence of bacteriuria, for the following reasons:
   a. If an overnight urine sample is not used, there may not have been enough time for the nitrate to convert to nitrite in the bladder.
   b. Some UTIs are caused by organisms that do not convert nitrate to nitrite (eg, staphylococci, streptococci).
   c. Sufficient dietary nitrate may not be present for the nitrate-to-nitrite reaction to occur.

**Interfering Factors**

1. Azo dye metabolites and bilirubin can produce false-positive results.
2. Ascorbic acid can produce false-negative results.
3. False-positive results can be obtained if the urine sits too long at room temperature, allowing contaminant bacteria to multiply.
4. High specific gravity will reduce the sensitivity.

---

**Clinical Alert**

A negative urine nitrate test should never be interpreted as indicating the absence of bacteria.

**Interventions**

**Pretest Patient Preparation**

1. Explain the test purpose and urine specimen collection procedure. Instruct the patient in the procedure necessary for a clean-catch (midstream) specimen.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and monitor appropriately.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Urine Leukocyte Esterase**

Usually, the presence of leukocytes (WBCs) in the urine indicates a UTI. The leukocyte esterase test detects esterase released by the leukocytes into the urine. This is a standardized means for the detection of WBCs.

Microscopic examination and chemical testing are used to determine the presence of leukocytes in the urine. The chemical test is done with a leukocyte esterase dipstick. This test can also detect intact leukocytes, lysed leukocytes, and
WBC casts.

Reference Values

**Normal**

**Negative**

Procedures

1. Collect a fresh, random urine specimen with a clean-catch or midstream technique.
2. Follow directions for dipstick use exactly. Timing is critical for accurate results.
3. Note that a positive result causes a purple color on the dipstick. The test is not designed to measure the amount of leukocytes.

Clinical Implications

1. Positive results are clinically significant and indicate:
   - a. Cystitis (UTI)
   - b. Acute pyelonephritis
   - c. Acute Bright's disease
   - d. Bladder tumor
   - e. Systemic lupus erythematosus (SLE)
   - f. Tuberculosis infection
2. Urine with positive results from the dipstick should be examined microscopically for WBCs and bacteria.

Interfering Factors

1. False-positive results
   - a. Vaginal discharge, parasites, histocytes
   - b. Drug therapies (eg, ampicillin, kanamycin)
   - c. Salicylate toxicity
   - d. Strenuous exercise
2. False-negative results
   - a. Large amounts of glucose or protein
   - b. High specific gravity
   - c. Certain drugs (eg, tetracycline)

**Clinical Alert**

A urine sample that tests positive for both nitrite and leukocyte esterase should be cultured for pathogenic bacteria.

Interventions

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Urine Bilirubin

Bilirubin is formed in the reticuloendothelial cells of the spleen and bone marrow as a result of the breakdown of hemoglobin; it is then transported to the liver. Urinary bilirubin levels are increased to significant levels in the presence of any disease process that increases the amount of conjugated bilirubin in the bloodstream (see Chap. 6). Elevated amounts appear when the normal degradation cycle is disrupted by obstruction of the bile duct or when the integrity of the liver is damaged.

Urine bilirubin aids in the diagnosis and monitoring of treatment for hepatitis and liver damage. Urine bilirubin is an early sign of hepatocellular disease or intrahepatic or extrahepatic biliary obstruction. It should be a part of every UA because bilirubin often appears in the urine before other signs of liver dysfunction (eg, jaundice, weakness) become apparent. Not only does the detection of urinary bilirubin provide an early indication of liver disease, but also its presence or absence can be used in determining the cause of clinical jaundice.

Reference Values

**Normal**

**Negative** (0–0.02 mg/dL or 0–0.34 µmol/L)

Procedure

1. Examine the urine within 1 hour of collection because urine bilirubin is unstable, especially when exposed to light. If the urine is yellow-green to brown, shake the urine. If a yellow-green foam develops, bilirubin is probably present. Bilirubin alters the surface tension and allows foam to form. The yellow color is the bilirubin.
2. Chemical strip testing:
   - a. Dip a chemically reactive dipstick into the urine sample according to the manufacturer's directions.
b. Close comparison of color changes on the dipstick with control colors on the color chart is an absolute necessity. Failure to make a close approximation of color may result in failure to recognize urine bilirubin. Good lighting is required.

c. Interpret results as “negative” to “3+” or as “small,” “moderate,” or “large” amounts of bilirubin.

3. When it is crucial to detect even very small amounts of bilirubin in the urine, as in the earliest phase of viral hepatitis, Icotest tablets are preferred for testing because they are more sensitive to urine bilirubin. When elevated amounts of urine bilirubin are present, a blue to purple color forms on the absorptive mat. The intensity of the color and the rapidity of its development are directly proportional to the amount of bilirubin in the urine.

Clinical Implications

1. Even trace amounts of bilirubin are abnormal and warrant further investigation. Normally, there is no detectable bilirubin in the urine.

2. Increased bilirubin occurs in:
   a. Hepatitis and liver diseases caused by infections or exposure to toxic agents (cirrhosis)
   b. Obstructive biliary tract disease
   c. Liver or biliary tract tumors
   d. Septicemia
   e. Hyperthyroidism

NOTE

Urine bilirubin is negative in hemolytic disease.

Interfering Factors

1. Drugs may cause false-positive or false-negative results (see Appendix J).

2. Bilirubin rapidly decomposes when exposed to light; therefore, urine should be tested immediately.

3. High concentrations of ascorbic acid or nitrate cause decreased sensitivity.

Clinical Alert

Pyridium-like drugs or urochromes may give the urine an amber or reddish color and can mask the bilirubin reaction.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.

2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately for liver disease.

2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Urine Urobilinogen, Random and Timed

Bilirubin, which is formed from the degradation of hemoglobin, is transformed through the action of bacterial enzymes into urobilinogen after it enters the intestines. Some of the urobilinogen formed in the intestine is excreted as part of the feces, where it is oxidized to urobilin; another portion is absorbed into the portal bloodstream and carried to the liver, where it is metabolized and excreted in the bile. Traces of urobilinogen in the blood that escape removal by the liver are carried to the kidneys and excreted in the urine. This is the basis of the urine urobilinogen test. Unlike bilirubin, urobilinogen is colorless.

Urine urobilinogen is one of the most sensitive tests available to determine impaired liver function. Urinary urobilinogen is increased by any condition that causes an increase in the production of bilirubin and by any disease that prevents the liver from normally removing the reabsorbed urobilinogen from the portal circulation. An increased urobilinogen level is one of the earliest signs of liver disease and hemolytic disorders.

Although it cannot be determined by reagent strip, the absence of urobilinogen is also diagnostically significant and represents an obstruction of the bile duct.

Reference Values

Normal

Random specimen: 0.1–1 Ehrlich U/dL or <1 mg/dL

2-hour specimen: 0.1–1.0 Ehrlich U/2 hours or <1 mg/2 hours

24-hour specimen: 0.5–4.0 Ehrlich U/24 hours or 0.5–4.0 mg/day

Procedure

1. Follow instructions for collecting a timed 24-hour, 2-hour, or random specimen. Check with the laboratory for specific protocols.

2. Perform the 2-hour timed collection from 1:00 p.m. to 3:00 p.m. (1300 to 1500) or from 2:00 p.m. to 4:00 p.m. (1400 to 1600) for best results because peak excretion occurs during this time. No preservatives are necessary. Record the total amount of urine voided. Protect the collection receptacle from light. Test immediately after specimen
Urine urobilinogen is increased when there is:

- Increased destruction of RBCs
  1. Hemolytic anemia
  2. Pernicious anemia (megaloblastic)
  3. Malaria
- Hemorrhage into tissues
  1. Pulmonary infarction
  2. Excessive bruising
- Hepatic damage
  1. Biliary disease
  2. Cirrhosis (viral and chemical)
  3. Acute hepatitis
- Cholangitis

Urine urobilinogen is decreased or absent when normal amounts of bilirubin are not excreted into the intestinal tract. This usually indicates partial or complete obstruction of the bile ducts. The stool is pale in color. Decreased urinary urobilinogen is associated with:

- Cholelithiasis
- Severe inflammation of the biliary ducts
- Cancer of the head of the pancreas

During antibiotic therapy, suppression of normal gut flora may prevent the breakdown of bilirubin to urobilinogen; therefore, urine levels will be decreased or absent.

More comprehensive information is obtained when the tests for urobilinogen and bilirubin are correlated (see Table 3.5 for comparisons).

### Table 3.5 Comparison of Urine Urobilinogen and Urine Bilirubin Values

<table>
<thead>
<tr>
<th>Test</th>
<th>In Health</th>
<th>In Hemolytic Disease</th>
<th>In Hepatic Disease</th>
<th>In Biliary Obstruction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urine urobilinogen</td>
<td>Normal</td>
<td>Increased</td>
<td>Increased</td>
<td>Low or absent</td>
</tr>
<tr>
<td>Urine bilirubin</td>
<td>Negative</td>
<td>Negative</td>
<td>Positive or negative</td>
<td>Positive</td>
</tr>
</tbody>
</table>

### Clinical Alert

Urine urobilinogen rapidly decomposes at room temperature or when exposed to light.

### Interfering Factors

1. Drugs that may affect urobilinogen levels include those that cause cholestasis and those that reduce the bacterial flora in the gastrointestinal tract. Check with the pharmacist for specific drugs patient is taking.
2. Peak excretion is known to occur from noon to 4:00 p.m. The amount of urobilinogen in the urine is subject to diurnal variation.
3. Strongly alkaline urine shows a higher urobilinogen level, and strongly acidic urine shows a lower urobilinogen level.
4. Drugs that may cause increased urobilinogen include drugs that cause hemolysis. Check with the pharmacist for specific drugs the patient is taking.
5. If the urine is highly colored, the strip will be difficult to read.

### Interventions

#### Pretest Patient Preparation

1. Explain test purpose and urine collection procedures.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

#### Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately for anemia and gastrointestinal disorders. Advise concerning need for follow-up testing.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

### MICROSCOPIC EXAMINATION OF URINE SEDIMENT

In health, the urine contains small numbers of cells and other formed elements from the entire genitourinary tract: casts and epithelial cells from the nephron; epithelial cells from the kidney, pelvis, ureters, bladder, and urethra; mucus threads and spermatozoa from the prostate; and possibly RBCs or WBCs and an occasional cast. In renal parenchymal disease, the urine usually contains increased numbers of cells and casts discharged from an organ that is otherwise accessible only by biopsy or surgery (Table 3.6). Urinary sediment provides information useful for both diagnosis and prognosis. It provides a direct sampling or urinary tract morphology.
Table 3.6 Microscopic Examination of Urine Sediment

<table>
<thead>
<tr>
<th>Urine Sediment Component</th>
<th>Clinical Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacteria</td>
<td>Urinary tract infection</td>
</tr>
<tr>
<td>Casts</td>
<td>Tubular or glomerular disorders</td>
</tr>
<tr>
<td>Broad casts</td>
<td>Formation occurs in collecting tubules; serious kidney disorder, extreme stasis of flow</td>
</tr>
<tr>
<td>Epithelial (renal) casts</td>
<td>Tubular degeneration</td>
</tr>
<tr>
<td>Fatty casts</td>
<td>Nephrotic syndrome</td>
</tr>
<tr>
<td>Granular</td>
<td>Renal parenchymal disease</td>
</tr>
<tr>
<td>Waxy</td>
<td>Slasis of flow</td>
</tr>
<tr>
<td>Hyaline casts</td>
<td>Chronic renal failure, chronic renal disease, congestive heart failure; stress or exercise</td>
</tr>
<tr>
<td>Red blood cell casts</td>
<td>Acute glomerulonephritis</td>
</tr>
<tr>
<td>White blood cell casts</td>
<td>Pyelonephritis, acute interstitial nephritis</td>
</tr>
<tr>
<td>Epithelial cells</td>
<td>Damage to various parts of urinary tract</td>
</tr>
<tr>
<td>Renal cells</td>
<td>Tubular damage</td>
</tr>
<tr>
<td>Squamous cells</td>
<td>Normal or contamination</td>
</tr>
<tr>
<td>Erythrocytes</td>
<td>Most renal disorders, menstruation; strenuous exercise</td>
</tr>
<tr>
<td>Fat bodies (oval)</td>
<td>Nephrotic syndrome</td>
</tr>
<tr>
<td>Leukocytes</td>
<td>Most renal disorders; urinary tract infection; pyelonephritis</td>
</tr>
</tbody>
</table>

The urinary sediment is obtained by pouring 1 mL of fresh, well-mixed urine into a conical tube and centrifuging the sample at a specific speed for 10 minutes. The supernatant is poured off, and 1 mL of the sediment is resuspended. A small drop is placed on a slide, cover-slipped, and examined microscopically.

The urine sediment can be broken down into cellular elements (RBCs, WBCs, and epithelial cells), casts, crystals, and bacteria. These may originate anywhere in the urinary tract. When casts do occur in the urine, they may indicate tubular or glomerular disorders.

Casts are the only elements found in urinary sediment that are unique to the kidneys. They are formed primarily within the lumen of the distal convoluted tubule and collecting duct, providing a microscopic view of conditions within the nephron. Their shapes are representative of the tubular lumen.

Cast width is significant in determining the site of origin and may indicate the extent of renal damage. The width of the cast indicates the diameter of the tubule responsible for its formation. Cast width is described as narrow (as wide as 1 to 2 RBCs), medium-broad (3 to 4 RBCs), or broad (5 RBCs). The broad casts form in the collecting tubule and may be of any composition. Their presence usually indicates a marked reduction in the functional capacity of the nephron and suggests severe renal damage or end-stage renal disease.

The major constituent of casts is Tamm-Horsfall protein, a glycoprotein excreted by the renal tubular cells. It is found in normal and abnormal urine and is not detected by the urine dipstick method.

Clinical Alert

Microscopic examination of urine sediment can provide the following information:

1. Evidence of renal disease as opposed to infection of the lower urinary tract.
2. Type and status of a renal lesion or disease.

An 82-year-old female resident displayed the following signs and symptoms related to a urinary tract infection: high fever (101.0°F) for 24 hours; lethargy past 2 days; cloudy, foul-smelling urine; and dysuria. Urinalysis microscopic exam and culture sensitivity were ordered.

URINALYSIS

Interpretation of test results for routine urinalysis and urine culture with interventions.

Urinalysis Report:

<table>
<thead>
<tr>
<th>Macroscopic Analysis</th>
<th>Normal</th>
<th>Date: 06/16/03 Time: 2130</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color</td>
<td>Pale yellow-amber</td>
<td>Yellow</td>
</tr>
<tr>
<td>Clarity</td>
<td>Clear to slightly hazy</td>
<td>Cloudy *</td>
</tr>
</tbody>
</table>

Urine chemistries

<table>
<thead>
<tr>
<th></th>
<th>Normal</th>
<th>Date: 06/16/03 Time: 2130</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific gravity</td>
<td>1.005–1.030</td>
<td>1.0–2.0 *</td>
</tr>
<tr>
<td>Glucose</td>
<td>Negative</td>
<td>Negative</td>
</tr>
<tr>
<td>Ketones</td>
<td>Negative</td>
<td>Negative</td>
</tr>
<tr>
<td>pH</td>
<td>5.0–8.0</td>
<td>8.5 High</td>
</tr>
<tr>
<td>Protein</td>
<td>Negative</td>
<td>30 *</td>
</tr>
<tr>
<td>Blood</td>
<td>Negative</td>
<td>Small *</td>
</tr>
<tr>
<td>Bilirubin</td>
<td>Negative</td>
<td>Negative</td>
</tr>
<tr>
<td>Test</td>
<td>Value</td>
<td>Result</td>
</tr>
<tr>
<td>-----------------</td>
<td>-------------</td>
<td>--------------</td>
</tr>
<tr>
<td>Urobilinogen</td>
<td>0.2–1.0 EU/dL</td>
<td>0.2</td>
</tr>
<tr>
<td>Nitrite</td>
<td>Negative</td>
<td>Pos *</td>
</tr>
<tr>
<td>Leukocyte ester</td>
<td>Negative</td>
<td>Small</td>
</tr>
</tbody>
</table>

**Microscopic examination**

<table>
<thead>
<tr>
<th>Test</th>
<th>Value</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>BACT/hpf</td>
<td>None</td>
<td>4+ *</td>
</tr>
<tr>
<td>WBC/hpf</td>
<td>0–2</td>
<td>50–100</td>
</tr>
<tr>
<td>RBC/hpf</td>
<td>0–2</td>
<td>2–5 *</td>
</tr>
<tr>
<td>SQ EPITHT/lpf</td>
<td>0–2</td>
<td>10–20 *</td>
</tr>
<tr>
<td>Casts/lpf</td>
<td>None</td>
<td>Present *</td>
</tr>
<tr>
<td>Hyaline/lpf</td>
<td>Occasional</td>
<td>2–5 *</td>
</tr>
<tr>
<td>Triple Phos</td>
<td>None</td>
<td>Few *</td>
</tr>
</tbody>
</table>

* = abnormal, HPF = high-powered field, LPF = low-powered field, NEG = negative, BACT = bacteria, WBC = white blood cells, RBC = red blood cells, SPEC = specific (as in specific gravity), POS = positive, TRC = trace, ABN = abnormal, EU = Ehrlich units, MIC = minimum inhibitory concentration (the lowest concentration of the antibiotic that inhibits the organism's growth), S = sensitive or susceptible, R = resistant, TMP-SMX = trimethoprim sulfamethoxazole

---

**Urine Culture Antibiotic Drug Sensitivity and Organism Susceptibility**

**Collected:** 06/16/03—Time 2130  
**Received:** 06/17/03—Time 1006  
**Final Report—6/19/03 of antibiotic drug sensitivity and organism susceptibility**

**Susceptibility Testing**— *E. coli*

**S** = sensitivity or susceptibility; **R** = resistant; **TMP-SMX** = trimethoprim sulfamethoxazole  

<table>
<thead>
<tr>
<th>Antibiotic</th>
<th>Susceptibility</th>
<th>MIC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ampicillin</td>
<td>S</td>
<td>2</td>
</tr>
<tr>
<td>Piperacillin</td>
<td>S</td>
<td>&lt; 8</td>
</tr>
<tr>
<td>Ampicillin/Subbac</td>
<td>S</td>
<td>8/4</td>
</tr>
<tr>
<td>Cefazolin</td>
<td>S</td>
<td>&lt; 8</td>
</tr>
<tr>
<td>Gentamicin</td>
<td>S</td>
<td>1</td>
</tr>
<tr>
<td>Tobramycin</td>
<td>S</td>
<td>1</td>
</tr>
<tr>
<td>Tetracycline</td>
<td>S</td>
<td>2</td>
</tr>
<tr>
<td>Ciprofloxacin</td>
<td>S</td>
<td>1</td>
</tr>
<tr>
<td>Levofloxacin</td>
<td>S</td>
<td>&lt; 2</td>
</tr>
<tr>
<td>Nitrofurantoin</td>
<td>S</td>
<td>&lt; 32</td>
</tr>
<tr>
<td>TMP-SMX</td>
<td>S</td>
<td>5/9.5</td>
</tr>
</tbody>
</table>

Results of the tests were abnormal outcomes and the following interventions started on 06/19/03 with Bactrim DS (double strength) BID × 7 days; then Bactrim SS (single strength) every day until further orders; force fluids as appropriate. Repeat urinalysis and culture and sensitivity in 2 weeks. The rationale for Bactrim as drug of choice was because of both sensitivity and MIC (see legend).

---

**Urine Red Blood Cells and Red Blood Cell Casts**

In health, erythrocytes (RBCs) occasionally appear in the urine. However, persistent findings of even small numbers of RBCs should be thoroughly investigated because these cells come from the kidney and may signal serious renal disease. They are usually diagnostic of glomerular disease.

**Reference Values**

**Normal**

<table>
<thead>
<tr>
<th>Test</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>RBCs</td>
<td>0–3/hpf (high-power field)</td>
</tr>
<tr>
<td>RBC casts</td>
<td>0/lpf (low-power field)</td>
</tr>
</tbody>
</table>

**Procedure for Microscopic Urine Examination**

1. Collect a random urine specimen. Transport the specimen to the laboratory as soon as possible.
2. Urinary sediment is microscopically examined under both the low-power field (lpf) and the high-power field (hpf). Low power is used to find and count casts; RBCs, WBCs, and bacteria show up and are counted under high power. Amounts present are defined in the following terms: few, moderate, packed, and packed solid; or 1+, 2+, 3+, and 4+. Crystals and other elements are also noted.
3. Microscopic results should be correlated with the physical and chemical findings to ensure the accuracy of the report (Table 3.7).
Table 3.7 Common Correlations in Urinalysis

<table>
<thead>
<tr>
<th>Microscopic Elements</th>
<th>Physical Examination</th>
<th>Dipstick Measurement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red blood cells</td>
<td>Turbidity, red to brown color</td>
<td>Blood</td>
</tr>
<tr>
<td>White blood cells</td>
<td>Turbidity</td>
<td>Protein</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Nitrite</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Leukocytes</td>
</tr>
<tr>
<td>Epithelial cast cells</td>
<td>Turbidity</td>
<td>Protein</td>
</tr>
<tr>
<td>Bacteria</td>
<td>Turbidity, odor</td>
<td>pH</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Nitrite</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Leukocytes</td>
</tr>
<tr>
<td>Crystals</td>
<td>Turbidity, odor</td>
<td>pH</td>
</tr>
</tbody>
</table>

*Positive result.

Clinical Implications

1. **RBC casts** indicate hemorrhage in the nephron.
   a. RBC casts are found in three forms:
      1. Intact RBCs
      2. Degenerating cells within a protein matrix
      3. Homogenous blood casts ("hemoglobin casts")
   b. RBC casts indicate acute inflammatory or vascular disorders in the glomerulus and are found in:
      1. Glomerulonephritis (acute and chronic)
      2. Renal infarction
      3. Lupus nephritis
      4. Goodpasture's syndrome
      5. Severe pyelonephritis
      6. Congestive heart failure
      7. Renal vein thrombosis
      8. Acute bacterial endocarditis
      9. Malignant hypertension
     10. Periarteritis nodosa
   c. **RBCs should be present** if RBC casts are in the sediment.

2. **Red blood cells**
   a. The finding of more than 1 or 2 RBCs/hpf is abnormal and can indicate:
      1. Renal or systemic disease (glomerulonephritis)
      2. Trauma to the kidney (vascular injury)
   b. Increased numbers of RBCs occur in:
      1. Pyelonephritis
      2. Systemic lupus erythematosus (SLE)
      3. Renal stones
      4. Cystitis (acute or chronic)
      5. Prostatitis
      6. Tuberculosis (renal)
      7. Genitourinary tract malignancies
      8. Hemophilia, coagulation disorders
      9. Malaria
     10. Polyarteritis nodosa
     11. Malignant hypertension
     12. Acute febrile episodes
   c. Greater numbers of RBCs than WBCs indicate bleeding into the urinary tract, as may occur with:
      1. Trauma
      2. Tumors of rectum, colon, pelvis
      3. Aspirin overdose or other toxic drugs
      4. Anticoagulant therapy overdose
      5. Thrombocytopenia

Clinical Alert

1. In health, RBCs are occasionally found in the urine. However, persistent findings of even small numbers of RBCs should be thoroughly investigated, the first step being to request a fresh urine specimen for repeat testing.
2. Rule out the possible presence of menstrual blood, vaginal bleeding, or trauma to the perineal area in a female patient.

Interfering Factors

1. Increased numbers of RBCs may be found after a traumatic catheterization and after passage of urinary tract or kidney stones.
2. Alkaline urine hemolyzes RBCs and dissolves casts ("ghosts").
3. Some drugs can cause increased numbers of RBCs in the urine (see Appendix J).
4. RBC casts and RBCs may appear after very strenuous physical activity or participation in contact sports.
5. Heavy smokers show small numbers of RBCs in the urine.
6. Yeast or oil droplets may be mistaken for RBCs.
Interventions

Pretest Patient Preparation
1. Explain test purpose and procedure for random urine sample collection.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcomes and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Urine White Blood Cells and White Blood Cell Casts

Leukocytes (WBCs) may originate from anywhere in the genitourinary tract. They are also capable of amoeboid migration through the tissues to sites of infection or inflammation. An increase in urinary WBCs is called pyuria and indicates the presence of an infection or inflammation in the genitourinary system. However, WBC casts always come from the kidney tubules.

Reference Values
Normal
WBCs: 0–4/hpf
Normal women may have slightly more WBCs.
WBC casts: 0/lpf

Procedure
1. Collect a random urine specimen and transport it to the laboratory as soon as possible.
2. Urinary sediment is microscopically examined under high power for cells and under low power for casts.

Clinical Implications
1. White blood cells
   a. Large numbers of WBCs (>30/hpf) usually indicate acute bacterial infection within the urinary tract.
   b. Increased WBCs are seen in:
      1. All renal disease
      2. Urinary tract disease (e.g., cystitis, prostatitis, urethritis)
      3. Appendicitis, pancreatitis
      4. Strenuous exercise
      5. Chronic pyelonephritis
      6. Bladder tumors
      7. Tuberculosis
      8. Lupus erythematosus
      9. Interstitial nephritis
      10. Glomerulonephritis
   c. In bladder infections, WBCs tend to be associated with bacteria, epithelial cells, and relatively few RBCs.
   d. Large numbers of lymphocytes and plasma cells in the presence of a kidney transplant may indicate early tissue rejection (acute renal allograft rejection).
   e. Eosinophils are associated with tubulointerstitial disease and hypersensitivity to penicillin.
   f. WBC clumps suggest renal origin of WBCs and should be reported when present.

2. WBC casts
   a. WBC casts indicate renal parenchymal infection and may occur in:
      1. Pyelonephritis (most common cause)
      2. Acute glomerulonephritis
      3. Interstitial nephritis
      4. Lupus nephritis
   b. It can be very difficult to differentiate between WBC casts and epithelial cell casts.

Clinical Alert
A urine culture (see Chap. 7) should be done if elevated urine WBCs are found.

Interfering Factors
Vaginal discharge can contaminate a specimen with WBCs. Either a clean-catch urine specimen or a catheterized urine specimen should be obtained to rule out contamination as the cause for WBCs in the urine.

Clinical Alert
Pyelonephritis may remain completely asymptomatic even though renal tissue is being progressively destroyed. Therefore, careful examination (using low power) of urinary sediment for leukocyte casts is vital.

Interventions
Pretest Patient Preparation
The pretest care is the same as for the urine RBC test.

Posttest Patient Aftercare
The posttest care is the same as for the urine RBC test.
Urine Epithelial Cells and Epithelial Casts

Renal epithelial cell casts are formed from cast-off tubule cells that slowly degenerate, first into coarse and then into fine granular material. Epithelial casts are the most rare casts.

Urine epithelial cells are of three kinds:

1. **Renal tubule epithelial cells** are round and slightly larger than WBCs. Each cell contains a single large nucleus. These are the types of epithelial cells associated with renal disease. However, the presence of an occasional renal epithelial cell is not unusual because the renal tubules are continually sloughing old cells. In cases of acute tubular necrosis, renal tubular epithelial cells containing large nonlipid vacuoles may be seen. These are referred to as **bubble cells**. When lipids cross the glomerular membrane, the renal epithelial cells absorb the lipids and become highly refractive. These are called **oval fat bodies**. Both of these findings are significant and should be reported.

2. **Bladder epithelial cells** are larger than renal epithelial cells. They range from round to pear-shaped to columnar. Also known as “transitional” epithelial cells, they line the urinary tract from the renal pelvis to the proximal two thirds of the urethra.

3. **Squamous epithelial cells** are large, flat cells with irregular borders, a single small nucleus, and abundant cytoplasm. Most of these cells are urethral and vaginal in origin and do not have much diagnostic importance.

Reference Values

**Normal**

Renal tubule epithelial cells: 0–3/hpf

Squamous epithelial cells are common in normal urine sample.

Renal tubule epithelial casts: 0 (not seen)

Procedure

1. Collect a random urine specimen.
2. Examine the urine sediment microscopically.

Clinical Implications

1. Epithelial cell casts are found when they are also present in the urine after exposure to toxic agents or viruses.
2. Renal tubular epithelial cells are found in:
   a. Acute tubular necrosis
   b. Acute glomerulonephritis (secondary effects)
   c. Pyelonephritis
   d. Salicylate overdose (toxic reaction)
   e. Impending allograft rejection
   f. Viral infections (eg, cytomegalovirus)
   g. Poisoning from heavy metals or other toxins

Urine Hyaline Casts

Hyaline casts are clear, colorless casts that are formed when a renal protein within the tubules (Tamm-Horsfall protein) precipitates and gels. Tamm-Horsfall protein is excreted at a fairly constant rate by the tubule cells and provides immunologic protection from infection. Hyaline casts can be seen in physiologic states such as strenuous exercise and even in the mildest renal disease. They are not associated with any one particular disorder.

Reference Values

**Normal**

Occasional (0–2/lpf)

Procedure

1. Obtain a fresh urine sample.
2. Examine urinary sediment microscopically for casts under low power.
3. Examine casts when the light intensity is reduced because they are colorless and transparent.
4. Note that wrinkling and convoluting of the cast occurs as it ages.

Clinical Implications

1. Hyaline casts indicate possible damage to the glomerular capillary membrane. These casts appear in:
   a. Glomerulonephritis, pyelonephritis
   b. Malignant hypertension
   c. Chronic renal disease
   d. Congestive heart failure
   e. Diabetic nephropathy
2. Hyaline casts may be a temporary phenomenon in the presence of:
   a. Fever (dehydration)
   b. Postural orthostatic lordotic strain
   c. Emotional stress
   d. Strenuous exercise
   e. Heat exposure
3. Nephrotic syndrome may be suspected when large numbers of hyaline casts appear in the urine together with significant proteinuria, fine granular casts, fatty casts, oval bodies, or fat droplets.
4. In cylindroiduria, large numbers of hyaline casts may be present, but protein in the urine is absent. Cylindroids are hyaline casts that have been formed at the junction of the ascending loop of Henle and therefore have tapered ends.

**Clinical Alert**

Casts may not be found even when proteinuria is significant if the urine is dilute (1.010 SG) or alkaline. In these cases, the casts are dissolved as soon as they are formed.

**Interventions**
The pretest and posttest care are the same as for the urine RBC test.

**Urine Granular Casts**

Granular casts appear homogeneous, coarsely granular, colorless, and very dense. They then further degenerate into finely granular casts. It is not necessary to distinguish the different granular casts. Granular casts may result from degradation of cellular casts, or they may represent direct aggregation of serum proteins into a matrix of Tamm-Horsfall microprotein.

**Reference Values**

Normal
Occasional (0–2/lpf)

**Procedure**

1. Collect a random urine specimen and transport it to the laboratory as soon as possible.
2. Examine urinary sediment microscopically under low power.

**Clinical Implications**

1. Granular casts are found in:
   a. Acute tubular necrosis
   b. Advanced glomerulonephritis
   c. Pyelonephritis
   d. Malignant nephrosclerosis
2. Granular casts are found with hyaline casts after strenuous exercise or severe stress.

**Interventions**
The pretest and posttest care are the same as for the urine RBC test.

**Urine Waxy Casts or Broad Casts (Renal Failure Casts) and Fatty Casts**

Casts are formed in the collecting tubules under conditions of extreme renal stasis. Waxy casts form from the degeneration of granular casts.

*Broad, waxy casts* are 2 to 6 times the width of ordinary casts and appear waxy and granular. Casts may vary in size as disease distorts the tubular structure (they get wider because they are a mold of the tubules). Also, as urine flow from the tubules becomes compromised, casts are more likely to form. The finding of broad, waxy casts suggests a serious prognosis—hence, the term renal failure casts.

*Fatty casts* are formed from the attachment of fat droplets and degenerating oval fat bodies into a protein matrix. Fatty casts are highly refractile and contain yellow-brown fat droplets, or oval fat bodies.

**Reference Values**

Normal
Negative (not seen)

**Procedure**
Examine urine sediment microscopically under low power.

**Clinical Implications**

1. Broad and waxy casts occur in:
   a. Severe renal failure
   b. Tubular inflammation and degeneration (nephrotic syndrome)
   c. Localized nephron obstruction (extreme stasis of urine flow)
   d. Malignant hypertension
   e. Renal amyloidosis
   f. Diabetic nephropathy
   g. Renal allograft rejection
2. Fatty casts are found in:
   a. Disorders causing lipiduria, such as nephrotic syndrome and lipid nephrosis
   b. Chronic glomerulonephritis
   c. Kimmelstiel-Wilson syndrome
   d. Lupus
   e. Toxic renal poisoning
Clinical Alert

The presence of broad, waxy casts signals very serious renal disease.

Interventions

The pretest and posttest care are the same as for the urine RBC test.

Urine Crystals

A variety of crystals may appear in the urine. They can be identified by their specific appearance and solubility characteristics. Crystals in the urine may present no symptoms, or they may be associated with the formation of urinary tract calculi and give rise to clinical manifestations associated with partial or complete obstruction of urine flow.

The type and quantity of crystalline precipitate varies with the pH of the urine. Amorphous crystalline material has no significance and forms as normal urine cools.

Procedure

1. Collect a random urine specimen. Crystal identification should be done on freshly voided specimens.
2. Examine the urinary sediment microscopically under high power.
3. The pH of the urine is an important aid to identification of crystals and must be noted.
4. The problems associated with the identification of abnormal crystals can be resolved by a check on the medications the patient is receiving, saving considerable time and energy.

Clinical Implications

Table 3.8 describes the meaning of urine crystal findings.

Table 3.8 Urine Crystals

<table>
<thead>
<tr>
<th>Type of Crystal</th>
<th>Color</th>
<th>Shape</th>
<th>Clinical Implications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acid Urine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amorphous urates</td>
<td>Pink to brick red</td>
<td>Granules</td>
<td>Normal</td>
</tr>
<tr>
<td>Uric acid</td>
<td>Yellow-brown</td>
<td>Polymorphous—whetstones, rosettes or prisms, rhombohedral prisms, hexagonal plate</td>
<td>Normal; increased purine metabolism, gout, Lesch-Nyhan syndrome</td>
</tr>
<tr>
<td>Sodium urate</td>
<td>Colorless to yellow</td>
<td>Fan of slender prisms</td>
<td>No clinical significance</td>
</tr>
<tr>
<td>Cystine (rare)</td>
<td>Colorless, highly refractile</td>
<td>Flat hexagonal plates with well-defined edges, singly or in clusters</td>
<td>Cystinuria; cystinosis—cystine stones in kidney, crystals also in spleen and eyes</td>
</tr>
<tr>
<td>Cholesterol (rare)</td>
<td>Colorless</td>
<td>“Broken window panes” with notched corners</td>
<td>Nephritis, nephrotic syndrome, chyluria</td>
</tr>
<tr>
<td>Leucine (rare)</td>
<td>Yellow or brown, highly refractile</td>
<td>Spheroids with striations; pure form hexagonal</td>
<td>Protein breakdown, severe liver disease, Fanconi's syndrome</td>
</tr>
<tr>
<td>Tyrosine (rare)</td>
<td>Colorless or yellow</td>
<td>Fine, silky needles in sheaves or rosettes</td>
<td>Protein breakdown, severe liver disease, osteoharuse urine disease, tyrosinosis</td>
</tr>
<tr>
<td>Bilirubin</td>
<td>Reddish-brown</td>
<td>Cubes, rhombic plates, amorphous needles</td>
<td>Elevated bilirubin</td>
</tr>
<tr>
<td>Acid, Neutral, or Slightly Alkaline Urine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcium oxalate</td>
<td>Colorless</td>
<td>Octahedral dumbbells, often small—use high power</td>
<td>Normal; large amounts in fresh urine may indicate severe chronic renal disease, liver disease, ethylene glycol poisoning, diabetes mellitus, large doses of vitamin C</td>
</tr>
<tr>
<td>Hippuric acid (rare)</td>
<td>Colorless</td>
<td>Rhombic plates, four-sided prisms</td>
<td>No significance</td>
</tr>
<tr>
<td>Alkaline, Neutral, or Slightly Acid Urine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triple phosphate</td>
<td>Colorless</td>
<td>“Coffin lids,” 3–6 sided prism; occasionally fern-leaf</td>
<td>Urine stasis and chronic cystitis, chronic pyelitis and enlarged prostate</td>
</tr>
<tr>
<td>Alkaline Urine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcium carbonate</td>
<td>Colorless</td>
<td>Needles, spheres, dumbbells</td>
<td>Normal</td>
</tr>
<tr>
<td>Ammonium biurate</td>
<td>Yellow opaque brown</td>
<td>“Thorn apple” spheres, dumbbells, sheaves of needles</td>
<td>Normal</td>
</tr>
<tr>
<td>Calcium phosphate</td>
<td>Colorless</td>
<td>Prisms, plates, needles</td>
<td>Normal</td>
</tr>
<tr>
<td>Amorphous phosphates</td>
<td>White</td>
<td>Granules</td>
<td>Normal</td>
</tr>
</tbody>
</table>
Specific drugs (most commonly, ampicillin and sulfonamides) can cause increased levels of their own crystals, which could be a sign of improper hydration.

Interfering Factors

1. Refrigerated urine will precipitate out many crystals because the solubility properties of the compound are altered.
2. Urine left standing at room temperature will also cause precipitation of crystals or the dissolving of the crystals.
3. Radiographic dye can cause crystals in improperly hydrated patients. These resemble uric acid crystals and can be suspected in specimens that have an abnormally high specific gravity (>1.030).

Interventions

**Patient Preparation and Aftercare**
The pretest and posttest care are the same as for the urine RBC test.

**Urine Shreds**

Shreds consist of a mixture of mucus, pus, and epithelial (squamous) cells. They can be seen on gross examination.

**Procedure**

1. Examine a fresh urine specimen by visually checking for a hazy mass.
2. Centrifuge the specimen and examine the sediment microscopically to verify the presence of formed elements (Table 3.9).

### Table 3.9 Interpreting Urine Laboratory Findings

<table>
<thead>
<tr>
<th>Disease</th>
<th>Cause</th>
<th>Laboratory Findings</th>
<th>Signs</th>
<th>Chemical Findings</th>
<th>Microscopic Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute glomerulonephritis</td>
<td>Anti-basement membrane antibodies associated with strep infection, variety of infectious agents, toxins, allergens</td>
<td>Rapid appearance of hematuria, proteinuria, and casts</td>
<td>Gross hematuria, turbid, smoky</td>
<td>Protein &lt;1.0 g/dL Blood positive</td>
<td>Increased RBC, WBC, renal tubular epithelial Casts: RBC, granular, waxy, broad</td>
</tr>
<tr>
<td>Chronic glomerulonephritis</td>
<td>Represents end-stage result of persistent glomerular damage with continuing and irreversible loss of renal function</td>
<td>Symptoms include edema, hypertension, anemia, metabolic acidosis, oliguria progressing to anuria</td>
<td>Hematuria</td>
<td>Protein &gt;2.5 g/dL Blood, small amount SG low and fixed</td>
<td>Increased RBC, WBC, renal epithelial Casts: granular, waxy, broad</td>
</tr>
<tr>
<td>Nephrotic syndrome</td>
<td>Glomeruli whose basement membrane has become highly permeable to plasma proteins of large molecular weight and lipids, allowing them to pass in the tubules</td>
<td>Massive protein, edema, high levels of serum lipids, and low levels of serum albumin</td>
<td>Cloudy</td>
<td>Protein &gt;3.5 g/dL Blood, small amount</td>
<td>Increased RBC, oval fat bodies, free fat, renal epithelial Casts: fatty, waxy, renal</td>
</tr>
<tr>
<td>Acute tubular necrosis</td>
<td>Destruction of renal tubular epithelial cells Usually following a hypotensive event (shock), toxic element, or drugs and heavy metals</td>
<td>Oliguria and complete renal failure</td>
<td>Slightly cloudy</td>
<td>Protein &lt;1.0 g/dL Blood positive SG low</td>
<td>Increased RBC, WBC, renal epithelial Casts: renal, granular, waxy, broad</td>
</tr>
<tr>
<td>Cystitis (lower urinary tract)</td>
<td>Infection of the bladder most commonly caused by bacteria: <em>Escherichia coli</em> most common (85%)</td>
<td>Frequent and painful urination</td>
<td>Cloudy, foul smelling</td>
<td>Protein &lt;0.5 g/dL Blood, small amount Nitrite positive (usually) Leukocyte esterase positive (usually)</td>
<td>Increased WBC, bacteria, RBC, transitional epithelial</td>
</tr>
<tr>
<td>Urethritis (urethra in males)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Acute pyelonephritis (upper urinary tract)**

An infection of the kidney or renal pelvis
Caused by infectious organism that has traveled through the urinary tract and invaded the kidney tissue

More frequently in women with repeated urinary tract infections

<table>
<thead>
<tr>
<th>Protein &lt;1.0 g/dL</th>
<th>Blood positive</th>
<th>Nitrite positive</th>
<th>Leukocyte esterase positive</th>
</tr>
</thead>
<tbody>
<tr>
<td>Turbid, foul smelling</td>
<td>(usually)</td>
<td>(usually)</td>
<td>(usually)</td>
</tr>
</tbody>
</table>

Increased WBC (clumps), bacteria, renal epithelial
Casts: WBC, granular, renal occasionally waxy

**Chronic pyelonephritis**

Permanent scarring of the renal tissue

Polyuria and nocturia develop as tubular function is lost
With disease progression, there is hypertension and altered renal and glomerular flow

<table>
<thead>
<tr>
<th>Protein &lt;2.5 g/dL</th>
<th>Nitrite positive</th>
<th>Leukocyte esterase positive</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cloudy</td>
<td>(usually)</td>
<td>(usually)</td>
</tr>
</tbody>
</table>

Increased WBC, casts: granular, waxy, broad

**Acute interstitial nephritis**

Inflammation of the renal interstitium caused by drug toxicity or an allergic reaction

Fever, eosinophilia, skin rash

<table>
<thead>
<tr>
<th>Protein &lt;1 g/dL</th>
<th>Blood positive</th>
<th>Leukocyte esterase positive</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cloudy</td>
<td>(usually)</td>
<td>(usually)</td>
</tr>
</tbody>
</table>

Increased WBC, RBC, eosinophils, epithelial
Increased casts: granular, renal hyaline

SG, specific gravity.


**Clinical Implications**

1. When mucus predominates, the shreds float on the surface.
2. When epithelial cells predominate, the shreds occupy the middle zone.
3. When pus (WBCs) predominates, the shreds are drawn to the bottom of the specimen.
4. Other findings in urine caused by specimen contamination include microscopic yeast, *Trichomonas*, spermatozoa, vegetable fibers, parasites, and meat fibers. These should be reported because they have clinical significance.
   a. Yeast may indicate urinary moniliasis or vaginal moniliasis (*Candida albicans*)
   b. Parasites—usually from fecal or vaginal contamination
   c. Spermatozoa—seen after sexual intercourse, after nocturnal emissions, or in the presence of prostatic disease

**URINE CHEMISTRY**

**Urine Pregnancy Test; Human Chorionic Gonadotropin (hCG) Test**

From the earliest stage of development, the placenta produces hormones, either on its own or in conjunction with the fetus. The very young placental trophoblast produces appreciable amounts of the hormone human chorionic gonadotropin (hCG), which is excreted in the urine. This hormone is not found in the urine of men or of normal, young, nonpregnant women.

Increased urinary hCG levels form the basis of the tests for pregnancy; hCG is present in blood and urine whenever there is living chorionic/placental tissue. hCG is made up of α- and ß-subunits. The ß-subunit is the most sensitive and specific test for early pregnancy. hCG can be detected in the urine of pregnant women 26 to 36 days after the first day of the last menstrual period (ie, 5 to 7 days after conception). Pregnancy tests should return to negative 3 to 4 days after delivery.

**Reference Values**

- **Normal**
  - Positive: pregnancy exists
  - Negative: nonpregnant state

**Procedure**

1. Collect an early morning urine specimen. The first morning specimen generally contains the greatest concentration of hCG. A random specimen may be used, but the SG must be more than 1.005.
2. Do not use grossly bloody specimens. If necessary, a catheterized specimen should be used.

**Clinical Implications**

1. A positive result usually indicates pregnancy.
2. Positive results also occur in
a. Choriocarcinoma
b. Hydatidiform mole
c. Testicular and trophoblastic tumors in males
d. Chorioepithelioma
e. Chorioadenoma destruens
f. About 65% of ectopic pregnancies

3. Negative or decreased results occur in
a. Fetal demise
b. Abortion, threatened abortion (test remains positive for 1 week after procedure)

Interfering Factors

1. False-negative test results and falsely low levels of hCG may be caused by dilute urine (low SG) or by using a specimen obtained too early in pregnancy.
2. False-positive tests are associated with
   a. Proteinuria
   b. Hematuria
   c. The presence of excess pituitary gonadotropin
   d. Certain drugs (e.g., chlorpromazine, phenothiazines, methadone)

Urine Estrogen, Total and Fractions (Estradiol [E_2] and Estriol [E_3]), 24-Hour Urine and Total Estrogen—Blood

Estradiol is the most active of the endogenous estrogens. The test evaluates female menstrual and fertility problems. In men, estradiol is useful for evaluating estrogen-producing tumors. Estriol is the prominent urinary estrogen in pregnancy. Serial measurements reflect the integrity of the fetal-placental complex.

Total estrogens evaluate ovarian estrogen-producing tumors in premenarchal or postmenopausal females.

These measurements, together with the gonadotropin (follicle-stimulating hormone [FSH]) level (see Chap. 6), are useful in evaluating menstrual and fertility problems, male feminization characteristics, estrogen-producing tumors, and pregnancy. Estradiol (E_2) is the most active of the endogenous estrogens. Estriol (E_3) levels in both plasma and urine rise in pregnancy advances; significant amounts are produced in the third trimester. E_3 is no longer considered useful for detection of fetal distress. Total estrogens may be helpful to establish time of ovulation and the optimum time for conception.

Reference Values

Normal
Normal values vary widely between women and men and in the presence or pregnancy, the menopausal state, or the follicular, ovulatory, or luteal stage of the menstrual cycle.

Urine Estradiol (E_2)

Men: 0–6 µg/24 hours or 0–22 nmol/day
Women: Follicular phase, 0–3 µg/24 hours or 0–11 nmol/day
       Ovulatory peak, 4–14 µg/24 hours or 15–51 nmol/day
       Luteal phase, 4–10 µg/24 hours or 15–37 nmol/day
       Postmenopausal, 0–4 µg/24 hours or 0–15 nmol/day

Urine Estriol (E_3) (wide range of normal)

Men: 1–11 µg/24 hours or 4–40 nmol/day
Women: Follicular phase, 0–14 µg/24 hours or 0–51 nmol/day
        Ovulatory phase, 13–54 µg/24 hours or 48–198 nmol/day
        Luteal phase, 8–60 µg/24 hours or 29–220 nmol/day
        Postmenopausal, 0–11 µg/24 hours or 0–40 nmol/day
Pregnancy: 1st trimester, 0–800 µg/24 hours or 0–2,900 nmol/day
           2nd trimester, 800–12,000 µg/24 hours or 2,900–44,000 nmol/day
           3rd trimester, 5,000–50,000 µg/24 hours or 18,000–180,000 nmol/day

Urine Total Estrogens

Men: 15–40 µg/24 hours or 55–147 nmol/day
Women: Menstruating, 15–80 µg/24 hours or 55–294 nmol/day
        Postmenopausal, <20 µg/24 hours or <73 nmol/day
Pregnancy: 1st trimester, 0–800 µg/24 hours or 0–2,900 nmol/day
           2nd trimester, 800–5,000 µg/24 hours or 2,900–18,350 nmol/day
           3rd trimester, 5,000–50,000 µg/24 hours or 2,900–183,000 nmol/day

Blood Total Estrogens
Men: 20–80 pg/mL or 20–80 ng/L  
Women: 60–400 pg/mL or 60–400 ng/L  
Postmenopausal: <130 pg/mL or <130 ng/L  
Prepuberty: <25 pg/mL or <25 ng/L  
Puberty: 30–280 pg/mL or 30–280 ng/mL

**NOTE**

Total serum estrogen does not measure estriol (E₃) and should not be used in pregnancy or to assess fetal well-being.

**Procedure**

1. Obtain a venous blood sample if needed for total estrogen.
2. Collect a 24-hour urine specimen and use boric acid preservative for all estrogen tests. Keep the container refrigerated or on ice during collection.
3. Follow general collection procedures for a 24-hour urine specimen (see Long-Term, Timed Urine Specimen, page 171).
4. Record the age and sex of the patient.
5. Ensure that the number of gestation weeks is communicated if patient is pregnant.
6. Document the number of days into the menstrual cycle for the nonpregnant woman.

**Clinical Implications**

1. **Increased urine E₂** is found in the following conditions:
   a. Feminization in children (testicular feminization syndrome)
   b. Estrogen-producing tumors
   c. Precocious puberty related to adrenal tumors
   d. Hepatic cirrhosis
   e. Hyperthyroidism
   f. In women, estradiol increases during menstruation, before ovulation, and during the 23rd to 41st weeks of pregnancy.
2. **Decreased urine E₂** occurs in:
   a. Primary and secondary hypogonadism
   b. Kallmann's syndrome
   c. Hypofunction or dysfunction of the pituitary or adrenal glands
   d. Menopause
3. **Increased urine E₃** occurs in pregnancy; there is a sharp rise when delivery is imminent.
4. **Decreased urine E₃** is found in:
   a. Cases of placental insufficiency or fetal distress (abrupt drop of > 40% on 2 consecutive days). Serial monitoring of estriol for 4 consecutive days is recommended to evaluate fetal distress.
   b. Congenital heart disease
   c. Down syndrome
5. Blood and urine total estrogens are **increased in**:
   a. Malignant neoplasm of adrenal gland
   b. Malignant neoplasm of cell tumor of ovary
   c. Benign neoplasm of ovary
   d. Granulosa cell tumor of ovary
   e. Lutein cell tumor of ovary
   f. Theca cell tumor of ovary
   g. Testicular tumors
6. Blood and urine total estrogens are **decreased in**:
   a. Ovarian hypofunction (ovarian agenesis, primary ovarian malfunction)
   b. Intrauterine death
   c. Preeclampsia
   d. Hypopituitarism
   e. Hypofunction of adrenal cortex
   f. Menopause
   g. Anorexia nervosa

**Clinical Alert**

Estradiol may be used for Pergonal (menotropins, ie, combination of follicle-stimulating and luteinizing hormones used to promote ovarian follicular growth) monitoring. Serial measurements of E₂ during ovulation induction enable the physician to minimize high E₂ levels caused by ovarian overstimulation and thereby decrease side effects.

**Clinical Alert**

Normal values are guidelines and must be interpreted in conjunction with clinical findings.

**Interfering Factors**

1. Total estrogens
   a. Oral contraceptives
b. Estrogen therapy
c. Progesterone therapy
d. Pregnancy and after administration of acetazolamide during pregnancy

2. Estradiol
   a. Radioactive pharmaceuticals
   b. Oral contraceptives

3. Estriol
   a. Glucose and protein interfere with outcome.
   b. Day-to-day physiologic variation can be as much as 30%; therefore, single determinations are of limited use.
   c. Renal disease—in which case a serum assay would be more accurate.

Interventions

Pretest Patient Preparation

1. Explain the test purpose and procedure.
2. Stress test compliance. The patient must be able to adjust daily activities to accommodate urine collection protocols.
3. Do not administer radioisotopes for 48 hours before specimen collection.
4. Discontinue all medications for 48 hours before specimen collection (with physician's approval). Drugs deemed necessary must be documented and communicated.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume medications and activity.
2. Interpret test outcomes, monitor, and counsel appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

URINE DRUG INVESTIGATION SPECIMENS

When screening for unknown drugs, the most valuable samples are obtained from urine, gastric contents, and blood. Urine drug screening is preferred for several reasons:

1. Specimens are easily procured.
2. It is not an invasive procedure (unless bladder catheterization is involved).
3. Drug concentrations are more elevated in urine or may not be detectable in blood.
4. Drug metabolites are excreted for a longer period (days or weeks) through urine, indicating past drug use.
5. Urine test procedures are more easily done and are more economical.

Clinical Alert

Blood is the preferred medium for ethyl alcohol testing because the alcohol concentration is more elevated and therefore more reliably measured in a blood sample (see Chap. 6).

Toxicology screening should be performed:

1. To confirm clinical or postmortem diagnosis
2. To differentiate drug-induced disease from other causes, such as trauma or metabolic or infectious disease processes
3. To identify contributing diagnoses, such as ethanol abuse, trauma, presence of other drugs, or underlying psychosis
4. To seek a basis for high-risk interventions such as hemodialysis
5. To test for drug abuse in the workplace, especially when public safety is at risk or concern; also to test for doping in athletes
6. As part of preemployment screening for drug use or abuse
7. To test prisoners and parolees randomly to deter or detect drug use (Chart 3.2)
Chart 3.2 Common Urine Drug Tests

<table>
<thead>
<tr>
<th>Drug Type</th>
<th>Example Drugs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcohol</td>
<td></td>
</tr>
<tr>
<td>Amphetamines</td>
<td></td>
</tr>
<tr>
<td>Analgesics</td>
<td></td>
</tr>
<tr>
<td>Barbiturates</td>
<td></td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td></td>
</tr>
<tr>
<td>Cocaine, &quot;crack&quot;</td>
<td></td>
</tr>
<tr>
<td>Cyanide</td>
<td></td>
</tr>
<tr>
<td>Lysergic acid diethylamide (LSD)</td>
<td></td>
</tr>
<tr>
<td>Major tranquilizers †</td>
<td></td>
</tr>
<tr>
<td>Marijuana</td>
<td></td>
</tr>
<tr>
<td>Opiates</td>
<td></td>
</tr>
<tr>
<td>Phencyclidine (PCP)</td>
<td></td>
</tr>
<tr>
<td>Sedatives</td>
<td></td>
</tr>
<tr>
<td>Stimulants</td>
<td></td>
</tr>
<tr>
<td>Sympathomimetics</td>
<td></td>
</tr>
</tbody>
</table>

**Footnotes**

* Many of these drugs are detectable in urine but are not detectable in blood serum. However, all drugs detectable in blood serum are also detectable in urine, except for glutethimide. † Because minor tranquilizers are almost completely metabolized, they are not likely to be detected in urine unless an overdose is taken.

**Clinical Alert**

When reporting drug test results for substance abuse, health care workers and patients need to be aware of the psychological, social, economic, and legal implications and the potential liabilities associated with mismanaged or incorrectly reported results. Documented procedures should be established and followed to ensure that before a result is reported, corroborating evidence exists to support that result. Confirmation of all positive results must be done through an equally sensitive and specific method that uses a different chemical principle to cross-check the initial results. Keep in mind that problems associated with incorrect test results are directly proportional to the volume of drug abuse testing being done.

Urine screening is not a cure-all for preventing substance abuse in the workplace. When properly implemented, however, it can support a well thought-out substance abuse rehabilitation program. Screening can detect a problem that the employee may not admit to having. Sure knowledge that an employee abuses drugs allows an employer to move with confidence toward handling the problem.

**Witnessed Urine Sampling for Suspected Substance Abuse**

The following procedure is an example of the chain of custody. A chain-of-custody document is originated at the time the sample is collected. The donor and the individual who witnessed specimen collection must sign and date the document, as does every person who handles the sample thereafter. The sealed chain-of-custody specimen bag remains in the possession and control of the collector or is kept in a secured area until shipment to the testing facility. Sealed collections are placed in large shipping cartons or specially designated bags.

After initial and confirmatory testing, the sample is resealed in a labeled bag and securely stored for 30 days or longer. All records of tests done on the sample and the chain-of-custody report must be carefully maintained.

The test results should be released only to predesignated, authorized persons to lessen the risk for false or speculative information being communicated to inappropriate persons.

Several factors may interfere with accurate outcomes and could cause incorrect or false-positive or false-negative results: higher or lower pH than normal; presence of blood, sodium chloride, detergents, or other contaminants; or low specific gravity.

**Procedure**

1. Ensure that the tested patient's signed informed consent form and photo identification are available; they are required.
2. Instruct patient to remove extra outer garments and leave them outside the bathroom. Make provisions for personal privacy during specimen procurement.
3. Direct the donor to void a random sample of 60 to 100 mL of urine into the clean specimen cup. The toilet may not be flushed at any time.
4. Have the witness transfer the contents of the cup into the laboratory specimen bottle on receipt of the voided specimen from the donor. The donor is present for the entire transfer procedure (viewing this and the following procedure).
5. Check and record any visible signs of contamination (eg, sediment, discoloration). The entire procedure must be witnessed by a trained, designated individual who is legally responsible to ensure that the specimen has been obtained from the correct patient.
6. Affix a temperature-sensing strip to the specimen bottle, and read and record the temperature within 4 minutes of specimen collection. Temperature strips and collection containers must be at room temperature (urine temperature must be between 90° and 98°F).
7. Very firmly screw down the cap onto the laboratory specimen bottle to seal it. The rim of the specimen bottle should
be dry.

8. Affix one end of the tamper-evident tape to the side of specimen bottle. Record the date collected, and have the donor initial the evidence tape. Wrap the tamper-evident tape across the top of the bottle, and overlap the free end of the tape with the other end to discourage tampering with the specimen.

9. Seal the specimen bottle in a zip-lock bag with absorbent material.

10. After sealing, have the donor sign and date the Drug Screen Request Form in the space provided. The collector then signs, dates, and provides a telephone number on the Drug Screen Request Form, indicating that all of the above steps have been followed. Every person who handles the sample thereafter must also sign the form (ie, chain-of-custody procedure).

11. Put the original and the first copy of the Drug Screen Form and the sealed laboratory specimen bottle into the shipping container, and seal it. Place tamper-evident tape across the seal.

12. Retain the third copy of the form for agency records.

13. Give the fourth copy of the form to the donor, or send it to the company or place of employment, as required.

Clinical Alert

National Institute for Drug Abuse (NIDA)-approved laboratory standards have stringent requirements. At the collection site (eg, bathroom), place toilet bluing markers in the toilet tank, and use tamper-proof tape on water faucet and soap dispensers to prevent water access for specimen dilution.

Clinical Implications Certain drugs can be detected in the urine for hours to several days after ingestion (Table 3.10). (Check with agency laboratory for specific drugs and specific time intervals.)

Table 3.10 Screening Limits

<table>
<thead>
<tr>
<th>Drugs Tested</th>
<th>Screening Cut-off Levels</th>
<th>Length of Detection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcohol</td>
<td>20 ng/mL</td>
<td>12 h</td>
</tr>
<tr>
<td>Ethanol (all methods)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amphetamines</td>
<td>1,000 ng/mL</td>
<td>2–3 d</td>
</tr>
<tr>
<td>D-Amphetamine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methamphetamine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Barbiturates</td>
<td>200 ng/mL</td>
<td>Up to 30 d</td>
</tr>
<tr>
<td>Secobarbital</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>200 ng/mL</td>
<td>Up to 40 d</td>
</tr>
<tr>
<td>Nordiazepam</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Marijuana</td>
<td>50 ng/mL</td>
<td>30–60 d</td>
</tr>
<tr>
<td>11-nor-D9-THC-9 COOH</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cocaine metabolite</td>
<td>300 ng/mL</td>
<td>2–4 d</td>
</tr>
<tr>
<td>Benzoylcegonine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methadone</td>
<td>300 ng/mL</td>
<td>8–60 h</td>
</tr>
<tr>
<td>Methadone HCI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methaqualone</td>
<td>300 ng/mL</td>
<td>Up to 7 d</td>
</tr>
<tr>
<td>Methaqualone HCI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Opiates</td>
<td>300 ng/mL</td>
<td>2–4 d</td>
</tr>
<tr>
<td>Morphine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PCP</td>
<td>25 ng/mL</td>
<td>2–3 d</td>
</tr>
<tr>
<td>Phencyclidine HCI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Propoxyphene</td>
<td>300 ng/mL</td>
<td>1–3 d</td>
</tr>
<tr>
<td>D-Propoxyphene HCI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tricyclic antidepressants (TCAs)</td>
<td>1,000 ng/mL</td>
<td>1–3 d</td>
</tr>
<tr>
<td>Desipramine (triage plus TCA)</td>
<td>1,000 ng/mL</td>
<td></td>
</tr>
<tr>
<td>Heroin</td>
<td>2,000 ng/mL</td>
<td>1–2 h</td>
</tr>
<tr>
<td>Acetylmorphine</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Interfering Factors
Factors associated with incorrect test results for urine drug screens include the presence of:

1. Detergents
2. Sodium chloride (table salt) (NaCl)
3. Low SG (dilute urine)
4. High pH (acid urine)
5. Low pH (alkaline urine)
6. Blood in the urine

Interventions

Pretest Patient Preparation

1. Explain the test purpose and the procedure for specimen collection.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately regarding results and possible retesting.
See Chapter 1 guidelines for safe, effective, informed posttest care.

**Clinical Alert**

Screening tests will be positive for opiates if poppy seeds are ingested (eg, bagels), if the screening cut-off level is 300 ng/mL. Therefore, many screen labs have raised the cut-off level to 2000 ng/mL.

**TIMED URINE TESTS**

**Urine Chloride (Cl), Quantitative (24-Hour)**

Normally, the urinary chloride excretion approximates the dietary intake. The amount of chloride excreted in the urine in a 24-hour period is an indication of the state of the electrolyte balance. Chloride is most often associated with sodium balance and fluid change.

The urine chloride measurement may be used to diagnose dehydration or as a guide in adjusting fluid and electrolyte balance in postoperative patients. It also serves as a means of monitoring the effects of reduced-salt diets, which are of great therapeutic importance in patients with cardiovascular disease, hypertension, liver disease, and kidney ailments.

Urine chloride is often ordered along with sodium and potassium as a 24-hour urine test. The urinary anion gap (Na + K) - Cl is useful for initial evaluation of hyperchloremic metabolic acidosis. It is also used to determine whether a case of metabolic alkalosis is salt responsive.

**Reference Values**

- **Normal**
  - Adult: 140–250 mEq/24 hours or 140–250 mmol/day
  - Child <6 years old: 15–40 mEq/24 hours or 15–40 mmol/day
  - Child 10–14 years old: 64–176 mEq/24 hours or 64–176 mmol/day

Children's values are much lower than adult values. Values vary greatly with salt intake and perspiration. Different labs may have different values. It is difficult to talk about normal and abnormal ranges because the test findings have meaning only in relation to salt intake and output.

**Procedure**

1. Collect a 24-hour urine specimen.
2. Record the exact starting and ending times on the specimen container and in the patient's health care record.
3. The complete specimen should be sent to the laboratory for refrigeration until it can be analyzed.

**Clinical Alert**

Because the electrolytes and water balance are so closely related, evaluate the patient's state of hydration by checking daily weight, by recording accurate intake and output, and by observing and recording skin turgor, the appearance of the tongue, and the appearance of the urine sample.

**Clinical Implications**

1. **Decreased** urine chloride occurs in:
   a. Chloride-depleted patients (<10 mEq/L or <10 mmol/L); these patients have low serum chloride and are chloride responsive (they respond to chloride therapy so that serum and urine levels return to normal).
      1. Syndrome of inappropriate antiuretic hormone (SIADH) secretion
      2. Vomiting, diarrhea, excessive sweating
      3. Gastric suction
      4. Addison's disease
      5. Metabolic alkalosis
      6. Massive diuresis from any cause
      7. Villous tumors of the colon
   b. Chloride is decreased by endogenous or exogenous corticosteroids (>20 mEq/L or >20 mmol/L); this condition is not responsive to chloride administration. Diagnosis of a chloride-resistant metabolic alkalosis helps identify a corticotropin (ACTH)- or aldosterone-producing neoplasm, such as:
      1. Cushing's syndrome
      2. Conn's syndrome
      3. Mineralocorticoid therapy
      4. Postoperative chloride retention
2. **Increased** urine chloride occurs in:
   a. Increased salt intake
   b. Adrenocortical insufficiency
   c. Potassium depletion
   d. Bartter's syndrome
   e. Salt-losing nephritis

**Interfering Factors**

1. Decreased chloride is associated with:
a. Carbenicillin therapy  
b. Reduced dietary intake of chloride  
c. Ingestion of large amounts of licorice  
d. Alkali ingestion  
e. Dehydration  

2. Increased chloride is associated with:  
a. Ammonium chloride administration  
b. Excessive infusion of normal saline  
c. Ingestion of sulfides, cyanides, halogens, bromides, and sulfhydryl compounds  

**Interventions**

**Pretest Patient Preparation**

1. Instruct the patient about the test purpose and the method for collecting a 24-hour specimen.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and monitor appropriately for fluid imbalances.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Urine Sodium (Na), Quantitative (24-Hour)**

Sodium is a primary regulator for retaining or excreting water and maintaining acid-base balance. The body has a strong tendency to maintain a total base content; on a relative scale, only small shifts are found even under pathologic conditions. As the predominant base substance in the blood, sodium helps to regulate acid-base balance because of its ability to combine with chloride and bicarbonate. Sodium also promotes the normal balance of electrolytes in the intracellular and extracellular fluids by acting in conjunction with potassium under the effect of aldosterone. This hormone promotes the 1:1 exchange of sodium for potassium or the hydrogen ion.

This test measures one aspect of electrolyte balance by determining the amount of sodium excreted in a 24-hour period. It is done for diagnosis of renal, adrenal, water, and acid-base imbalances.

**Reference Values**

**Normal**

| Adult: 40–220 mEq/24 hours or 40–220 mmol/day | Child: 41–115 mEq/24 hours or 41–115 mmol/day |
| Values are diet dependent. |

**Procedure**

1. Properly label a 24-hour urine container.  
2. The urine container must be refrigerated or kept on ice.  
3. Follow general instructions for 24-hour urine collections (see Long-Term, Timed Urine Specimen, page 171).  
4. Record exact starting and ending times on the specimen container and in the patient’s health care record.  
5. Transfer the specimen to the laboratory for proper storage when the test is completed.

**Clinical Implications**

1. **Increased** urine sodium occurs in:  
a. Adrenal failure (Addison’s disease) (primary and secondary)  
b. Salt-losing nephritis  
c. Renal tubular acidosis  
d. SIADH  
e. Diabetic acidosis  
f. Aldosterone defect (AIDS-related hypoadrenalism)  
g. Tubulointerstitial disease  
h. Bartter's syndrome  

2. **Decreased** urine sodium occurs in:  
a. Excessive sweating, diarrhea  
b. Congestive heart failure  
c. Adrenocortical hyperfunction  
d. Nephrotic syndromes with acute oliguria  
e. Prerenal azotemia  
f. Cushing’s disease  
g. Primary aldosteronism  

**Interfering Factors**

1. Increased sodium levels are associated with caffeine intake, diuretic therapy, dehydration, dopamine, postmenstrual diuresis, increased sodium intake, and vomiting (see Appendix J).  
2. Decreased sodium levels are associated with intake of corticosteroids and propranolol; low sodium intake; premenstrual and water retention; overhydration and stress diuresis (see Appendix J).

**Interventions**

**Pretest Patient Preparation**
1. Instruct the patient about the purpose of the test, method of collection, and specimen refrigeration or icing. Written instructions can be helpful.
2. Encourage intake of food and fluids.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert

Because electrolytes and water balance are so closely related, determine the patient's state of hydration by checking and recording daily weights, accurate intake and output of fluids, and observations about skin turgor, the appearance of the tongue, and the appearance of the urine.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor as necessary for fluid and electrolyte state.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Urine Potassium (K), Quantitative (24-Hour) and Random

Potassium acts as a part of the body's buffer system and serves a vital function in the body's overall electrolyte balance. Because the kidneys cannot completely conserve potassium, this balance is regulated by the excretion of potassium through the urine. It takes the kidney 1 to 3 weeks to conserve potassium effectively.

This test provides insight into electrolyte balance by measuring the amount of potassium excreted in 24 hours. This measurement is useful in the study of renal and adrenal disorders and water and acid-base imbalances. An evaluation of urinary potassium can be helpful in determining the origin of abnormal potassium levels. Urine potassium values <20 mEq/L (or <20 mmol/L) are associated with nonrenal conditions, whereas values >20 mEq/L (or >20 mmol/L) are associated with renal causes.

Reference Values

Normal
Adult: 25–125 mEq/24 hours or 25–125 mmol/day
Child: 10–60 mEq/24 hours or 10–60 mmol/day

Values are diet dependent.

Procedure

1. Label a 24-hour urine container properly.
2. Refrigerate the urine container or keep it on ice during the collection.
3. Follow general instructions for 24-hour urine collection (see Long-Term, Timed Urine Specimen, page 171).
4. Record exact starting and ending times on the container and in the patient's health care record.
5. Transfer the specimen to the laboratory for proper storage.
6. A random urine potassium determination may be done.

Clinical Implications

1. Increased urine potassium occurs in:
   a. Primary renal diseases
   b. Diabetic and renal tubule acidosis
   c. Albright-type renal disease
   d. Starvation (onset)
   e. Primary and secondary aldosteronism
   f. Cushing's syndrome
   g. Onset of metabolic alkalosis
   h. Fanconi's syndrome
   i. Bartter's syndrome
2. Decreased urine potassium occurs in:
   a. Addison's disease
   b. Severe renal disease (eg, pyelonephritis, glomerulonephritis)
   c. In patients with potassium deficiency, regardless of the cause, the urine pH tends to fall.

This occurs because hydrogen ions are released in exchange for sodium ions, given that both potassium and hydrogen are excreted by the same mechanism.

Interfering Factors

1. Increased urinary potassium is associated with:
   a. Acetazolamide and other diuretics
   b. Cortisone
   c. Ethylenediaminetetraacetic acid (EDTA) anticoagulant
   d. Penicillin, carbenicillin
   e. Thiazides
   f. Licorice
   g. Sulfates (see Appendix J)
2. Decreased urinary potassium is associated with:
   a. Amiloride
   b. Diazoxide
Intravenous glucose infusion (see Appendix J)

Clinical Alert
In the presence of excessive vomiting or gastric suctioning, the resulting alkalosis maintains urinary potassium excretion at levels inappropriately high for the degree of actual potassium depletion that occurs.

Interventions
Pretest Patient Preparation
1. Instruct the patient about the purpose of the test, the collection procedure, and the need for refrigeration or icing of the 24-hour urine specimen. Written instructions can be helpful.
2. Food and fluids are permitted and encouraged.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert
1. Because electrolytes and water balance are so closely related, determine the patient's state of hydration by checking and recording daily weights, accurate intake and output of fluids, and observations about skin turgor, the appearance of the tongue, and the appearance of the urine.
2. Observe for signs of muscle weakness, tremors, changes in electrocardiographic tracings, and dysrhythmias. The degree of hypokalemia or hyperkalemia at which these symptoms occur varies with each person.

Posttest Patient Aftercare
1. Interpret test outcomes and monitor appropriately for signs and symptoms of electrolyte imbalances and kidney disorders.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Urine Uric Acid, Quantitative (24-Hour)
Uric acid is formed from the metabolic breakdown of nucleic acids composed of purines. Excessive uric acid relates to excessive dietary intake of purines or to endogenous uric acid production in certain disorders. Normally, one third of the uric acid formed is degraded by bacteria in the intestines.

This test evaluates uric acid metabolism in gout and renal calculus formation. Evaluation of excess uric acid excretion is important to aid in evaluating stone formation and nephrolithiasis. It also reflects the effects of treatment with uricosuric agents by measuring the total amount of uric acid excreted within a 24-hour period.

Reference Values
Normal
With normal diet: 250–750 mg/24 hours or 1.48–4.43 mmol/day
With purine-free diet: <400 mg/24 hours or <2.48 mmol/day
With high-purine diet: <1000 mg/24 hours or <5.90 mmol/day

Procedure
1. Properly label a 24-hour urine container to which the appropriate preservative has been added.
2. Follow general instructions for 24-hour urine collection (see Long Term, Timed Urine Specimen, page 171).
3. Record exact starting and ending times on the specimen container and in the patient's health care record.
4. When collection is completed, send the specimen to the laboratory.

Clinical Implications
1. Increased urine uric acid (uricosuria) occurs in:
   a. Nephrolithiasis (primary gout)
   b. Chronic myelogenous leukemia (secondary nephrolithiasis)
   c. Polycythemia vera
   d. Lesch-Nyhan syndrome
   e. Wilson's disease
   f. Viral hepatitis
   g. Sickle cell anemia
   h. High uric acid concentration in urine with low urine pH may produce uric acid stones in the urinary tract. (These patients do not have gout.)
2. Decreased urine uric acid is found in:
   a. Chronic kidney disease
   b. Xanthinuria
   c. Folic acid deficiency
   d. Lead toxicity

Interfering Factors
1. Many drugs increase uric acid levels, including:
   a. Salicylates (aspirin) and other anti-inflammatory drugs
   b. Diuretics
   c. Vitamin C (ascorbic acid)
d. Warfarin
e. Cytotoxic drugs used to treat lymphoma and leukemia (see Appendix J)

2. Other factors increasing uric acid urine levels include:
   a. X-ray contrast media
   b. Strenuous exercise
c. Diet high in purines (eg, kidney, sweetbreads) (see Chap. 6)
3. Allopurinol decreases uric acid levels (see Appendix J)

Interventions

**Pretest Patient Preparation**

1. Instruct the patient about the test purpose, interfering factors, collection process, and refrigeration or icing of the 24-hour urine specimen. A written reminder may be helpful.
2. Encourage food and fluids. In some situations, a diet high or low in purines may be ordered during and before specimen collection.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Resume usual diet.
2. Interpret test outcomes and counsel appropriately regarding prescribed treatment and possible need for further testing.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Urine Calcium (Ca), Quantitative (24-Hour)**

Calcium hemostasis is maintained by the parathyroid hormone. The bulk of calcium excreted is eliminated in the stool. However, a small quantity of calcium is normally excreted in the urine. This amount varies with the quantity of dietary calcium ingested. Increased calcium in urine results from an increase in intestinal calcium absorption, a lack of renal tubule reabsorption of calcium, resorption or loss of calcium from bone, or a combination of these mechanisms. Values in both healthy and sick persons have a wide range.

The urine calcium test is used for evaluation of calcium intake and/or the rate of intestinal absorption, bone resorption, and renal loss. Urine calcium is high in 30% to 80% of cases of primary hyperparathyroidism but does not reliably diagnose this disease. Urine calcium test does not have much value in a differential diagnosis.

**Reference Values**

**Normal**

- Normal diet: 100–300 mg/24 hours or 2.50–7.50 mmol/day
- Low-calcium diet: 50–150 mg/24 hours or 1.25–3.75 mmol/day

**Procedure**

1. Label properly a 24-hour urine container.
2. Procure an acid-washed bottle. See Table 3-3 regarding 24-hour urine collection data.
3. Follow general instructions for 24-hour urine collection (see Long-Term, Timed Urine Specimen, page 171). Refrigerate during collection.
4. Record exact starting and ending times of the collection on the specimen container and in the patient's health care record.
5. Send the specimen to the laboratory when collection is completed.
6. Perform a random (Sulkowitch) test in an emergency. Follow directions for random urine collection in first part of the chapter.

**Clinical Implications**

1. Increased urine calcium is found in:
   a. Hyperparathyroidism (30% to 80% of cases)
   b. Sarcoïdosis
c. Primary cancers of breast and bladder
d. Osteolytic bone metastases (carcinoma, sarcoma)
e. Multiple myeloma
f. Paget's disease
g. Renal tubular acidosis
h. Fanconi's syndrome
i. Vitamin D intoxication
j. Idiopathic hypercalcuria
k. Osteoporosis (especially after immobilization)
l. Osteitis deformis
m. Thyrotoxicosis
2. Increased urinary calcium almost always accompanies increased blood calcium levels.
3. Calcium excretion levels greater than calcium intake levels are always excessive; urine excretion values > 400–500 mg/24 hours (>10–12.5 mmol/d) are reliably abnormal.
4. Increased calcium excretion occurs whenever calcium is mobilized from the bone, as in metastatic cancer or prolonged skeletal immobilization.
5. When calcium is excreted in increasing amounts, the situation creates the potential for nephrolithiasis or nephrocalcinosis, especially with high protein intake.
6. Decreased urine calcium is found in:  
   a. Hypoparathyroidism  
   b. Familial hypocalcuria hypercalcemia  
   c. Vitamin D deficiency  
   d. Preeclampsia  
   e. Acute nephrosis, nephritis, renal failure  
   f. Renal osteodystrophy  
   g. Vitamin D–resistant rickets  
   h. Metastatic carcinoma of prostate  
   i. Malabsorption syndrome—celiac-sprue disease, steatorrhea  
7. Urine calcium decreases late in normal pregnancy.

Interfering Factors

1. Falsely elevated levels may be caused by:  
   a. Some drugs (eg, calcitonin; vitamins A, K, and C, and corticosteroids) (see Appendix J)  
   b. Urine procured immediately after meals in which high calcium intake has occurred (eg, milk)  
   c. Increased exposure to sunlight  
   d. Immobilization (especially in children)  
2. Falsely decreased levels may be found with:  
   a. Increased ingestion of phosphate, bicarbonate, antacids  
   b. Alkaline urine  
   c. Thiazide diuretics (can be used to lower calcium levels therapeutically)  
   d. Oral contraceptives, estrogens  
   e. Lithium (see Appendix J)

Interventions

Pretest Patient Preparation

1. Instruct the patient about the test purpose and procedure. Written instructions may be helpful.  
2. Encourage food and fluids.  
3. If the urine calcium test is done because of a metabolic disorder, the patient should eat a low-calcium diet, and calcium medications should be restricted for 1 to 3 days before specimen collection.  
4. For a patient with a history of renal stone formation, urinary calcium results will be more meaningful if the patient's usual diet is followed for 3 days before specimen collection. Do not stop medications.  
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes, monitor and counsel accordingly regarding calcium imbalances.  
2. Follow Chapter 1 guidelines regarding safe, effective, informed posttest care.

Clinical Alert

1. Observe patients with very low urine calcium levels for signs and symptoms of tetany (muscle spasms, twitching, hyperirritable nervous system).  
2. The first sign of calcium imbalance may be pathologic fracture that can be related to calcium excess.  
3. The Sulkowitch test (random urine sample) can be used in an emergency, especially when hypercalcemia is suspected, because hypercalcemia is life-threatening.

Urine Magnesium (Mg), Quantitative (24-Hour)

Magnesium excretion controls serum magnesium balance. Urinary magnesium excretion is diet dependent. With normal dietary intake of 200–500 mg/day, urine excretion is normally 75–150 mg/24 hours (3–6 mmol/d). The remainder of the dietary intake is excreted in the stool.  

This test evaluates magnesium metabolism, investigates electrolyte status, and is a component of a workup for nephrolithiasis. It is useful for assessing the cause of abnormal serum magnesium.

Reference Values

Normal  
75–150 mg/24 hours or 3.0–6.0 mEq/24 hours or 3.00–6.00 mmol/day

Procedure

1. Collect a 24-hour urine specimen in a metal-free and acid-rinsed container. The pH must be < 2.  
2. Record exact starting and ending times.  
3. See Long-Term, Timed Urine Specimen (page 171) for 24-hour urine collection guidelines.

Clinical Implications

1. Increased urine magnesium is associated with:  
   a. Increased blood alcohol  
   b. Bartter's syndrome  
   c. Chronic glomerulonephritis
2. **Decreased** urine magnesium is associated with:
   a. Malabsorption
   b. Long-term chronic alcoholism (poor diet)
   c. Long-term parenteral therapy
   d. Magnesium deficiency
   e. Chronic renal disease
   f. Hypoparathyroidism
   g. Hypercalcuria
   h. Decreased renal function (eg, Addison's disease)

**Interfering Factors**

1. Increased magnesium levels are associated with:
   a. Corticosteroids
   b. Cisplatin therapy
   c. Thiazide diuretics
   d. Amphotericin (see [Appendix J](#))
   e. Blood in urine

2. Decreased magnesium levels; many drugs affect test outcomes (see [Appendix J](#))

**Interventions**

**Pretest Patient Preparation**

1. Explain purpose of test and collection procedures.
2. Instruct that the specimen will be unacceptable if it comes in contact with any type of metal.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and monitor appropriately for abnormal magnesium excretion.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Urine Oxalate, Quantitative (24-Hour)**

Oxalate is an end product of metabolism. Normal oxalate is derived from dietary oxalic acid (10%) and from the metabolism of ascorbic acid (35%–50%) and glycine (40%). Patients who form calcium oxalate kidney stones appear to absorb and excrete a higher proportion of dietary oxalate in the urine.

The 24-hour urine collection for oxalate is indicated in patients with surgical loss of distal small intestine, especially those with Crohn’s disease. The incidence of nephrolithiasis in patients who have inflammatory bowel disease is 2.6% to 10%. Hyperoxaluria is regularly present after jejunoileal bypass for morbid obesity; such patients may develop nephrolithiasis.

Oxaluria is also a characteristic of ethylene glycol intoxication. Additionally, vitamin C increases oxalate excretion and in some people may be a risk factor for calcium oxalate nephrolithiasis. Such ingestion can usually be determined through the patient's history. If oxalate excretion becomes normal after reduction of vitamin C intake, additional therapy to prevent stones may not be required.

**Reference Values**

**Normal**
- Men: <55 mg/24 hours or <611 µmol/day
- Women: <50 mg/24 hours or <555 µmol/day

**Procedure**

1. Collect and refrigerate or place on ice a 24-hour urine specimen according to protocols. Do not acidify.
2. See [Long-Term, Timed Urine Specimen](#) (page 171) for directions for a 24-hour urine collection.

**Clinical Implications**

1. **Increased** urine oxalate is associated with:
   a. Ethylene glycol poisoning (>150 mg/24 hours or >1700 µmol/day)
   b. Primary hyperoxaluria, a rare genetic disorder (100–600 mg/24 hours or 1100–6700 µmol/day [nephrocalcinosis])
   c. Pancreatic disorders (diabetes, steatorrhea)
   d. Cirrhosis, biliary diversion
   e. Vitamin B₆ deficiency (pyridoxine)
   f. Sarcoidosis
   g. Crohn's disease (inflammatory bowel disease)
   h. Celiac disease (sprue)
   i. Jejunoileal bypass for treatment of morbid obesity

2. **Decreased** urine oxalate occurs in renal failure

**Interfering Factors**

1. Foods containing oxalates, such as rhubarb, strawberries, beans, beets, spinach, tomatoes, gelatin, chocolate,
cocoa, and tea, cause increased levels.
2. Ethylene glycol and methoxyflurane cause increased levels (see Appendix J).
3. Calcium causes decreased levels (see Appendix J).
4. Ascorbic acid (vitamin C) increases levels.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Advise the patient to avoid foods that promote oxalate excretion before the test. A list of such foods is helpful.
   Normal fluid intake should be continued.
3. Vitamin C should not be taken within 24 hours before the beginning of the test nor during the test.
4. The patient should be ambulatory and preferably at home.
5. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume normal diet and exercise.
2. Interpret test outcomes and counsel appropriately about abnormal levels.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Urine Pregnanediol (24-Hour)

Pregnanediol levels in normally menstruating women are constant during the follicular phase. Levels increase sharply during the luteal phase. During pregnancy, levels gradually increase, falling sharply before the onset of labor and delivery.

This test measures ovarian and placental function. Specifically, it measures a part of the hormone progesterone and its principal excreted metabolite, pregnanediol. Progesterone exerts its main effect on the endometrium by causing the endometrium to enter the secretory phase and to become ready for the implantation of the blastocyte should fertilization take place.

Pregnanediol excretion is elevated in pregnancy and decreased in luteal deficiency or placental failure.

NOTE

A serum progesterone test is more informational and is now used as an index of progesterone production.

Reference Values

Normal

This test is difficult to standardize; it varies with age, sex, and length of existing pregnancy.

<table>
<thead>
<tr>
<th>Category</th>
<th>Value</th>
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</thead>
<tbody>
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<td>Child</td>
<td>&lt;0.1 mg/24 hours or &lt;0.312 µmol/day</td>
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<tr>
<td>Men</td>
<td>0–1.9 mg/24 hours or 0–5.9 µmol/day</td>
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<tr>
<td>Women</td>
<td>Follicular phase, 0–2.6 mg/24 hours or 0–8.1 µmol/day</td>
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<tr>
<td></td>
<td>Luteal, 2.6–10.6 mg/24 hours or 8.1–33.1 µmol/day</td>
</tr>
<tr>
<td>Pregnancy</td>
<td>1st trimester, 10–35 mg/24 hours or 31–109 µmol/day</td>
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<tr>
<td></td>
<td>2nd trimester, 35–70 mg/24 hours or 109–218 µmol/day</td>
</tr>
<tr>
<td></td>
<td>3rd trimester, 70–100 mg/24 hours or 218–312 µmol/day</td>
</tr>
</tbody>
</table>

Procedure

1. Label a 24-hour urine container properly.
2. Refrigerate the specimen or use a boric acid preservative. Check laboratory policy. Protect the specimen from light.
3. Follow general instructions for 24-hour urine collection (see Long-Term, Timed Urine Specimen, page 171).
4. Record exact starting and ending times on the specimen container and in the patient's health care record.
5. Send the completed specimen to the laboratory.

Clinical Implications

1. Increased urine pregnanediol is associated with:
   a. Luteal cysts of ovary (ovarian cyst)
   b. Arrhenoblastoma of the ovary
   c. Congenital hyperplasia of adrenal gland
   d. Granulosa theca cell tumor of ovary
2. Decreased urine pregnanediol is associated with:
   a. Amenorrhea (ovarian hypofunction)
   b. Threatened abortion (if <5.0 mg/24 hours or <15.6 µmol/day, abortion is imminent)
   c. Fetal death, intrauterine death, placental insufficiency
   d. Toxemia, eclampsia
   e. Ovarian failure
   f. Chronic nephritis in pregnancy
Interfering Factors
Decreased values occur with estrogen or progesterone therapy and with the usage of oral contraceptives.

Interventions

Pretest Patient Preparation

1. Instruct the patient about the test purpose and the 24-hour urine specimen collection procedure. A written reminder may be helpful.
2. Allow food and fluids.
3. See Chapter 1 guidelines regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately about abnormal ovarian and placental function.
2. Follow Chapter 1 guidelines regarding safe, effective, informed posttest care.

Urine Pregnanetriol (24-Hour)

Pregnanetriol is a ketogenic steroid reflecting one segment of adrenocorticol activity. Pregnanetriol should not be confused with pregnanediol, despite the similarity of name. This test has been largely replaced with the serum test 17-hydroxyprogesterone.

This 24-hour urine test is done to diagnose congenital adrenal hyperplasia, adrenogenital syndrome, owing to a defect in 21-hydroxilation. The diagnosis of adrenogenital syndrome is indicated in:

1. Adult women who show signs and symptoms of excessive androgen production with or without hypertension.
2. Craving for salt
3. Sexual precocity in boys
4. Infants who exhibit signs of failure to thrive
5. Presence of external genitalia in females (pseudohermaphroditism). In males, differentiation must be made between a virilizing tumor of the adrenal gland, neurogenic and constitutional types of sexual precocity, and interstitial cell tumor of the testes.

Reference Values

Normal
Adult female: 0–1.4 mg/24 hours or 0–4.2 µmol/day
Adult male: 0.2–2.2 mg/24 hours or 0.6–6.5 µmol/day
Child (<9 years old): <0.3 mg/24 hours or <0.9 µmol/day
Child (10–16 years old): 0.1–0.6 mg/24 hours or 0.3–1.8 µmol/day

Procedure

1. Label a 24-hour urine container properly.
2. Refrigerate the specimen if necessary; some laboratories may require a boric acid preservative in the collection receptacle.
3. Follow general instructions for 24-hour urine collection (see Long-Term, Timed Urine Specimen, page 171).
4. Record exact starting and ending times on the specimen container and in the patient's health care record.
5. Send the completed specimen to the laboratory.

Clinical Implications

1. Elevated urine pregnanetriol occurs in:
   a. Congenital adrenocortical hyperplasia
   b. Stein-Leventhal syndrome
   c. Ovarian and adrenal tumors
2. Decreased urine pregnanetriol occurs in:
   a. Hydroxylase deficiency (rare)
   b. Ovarian failure

Interventions

Pretest Patient Preparation

1. Instruct the patient about the test purpose and procedure for collection of a 24-hour urine specimen. A written reminder may be helpful.
2. Allow food and fluids.
3. Avoid muscular exercise before and during specimen collection.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately about adrenogenital syndrome.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Urine 5-Hydroxyindoleacetic Acid (5-HIAA) (24-Hour)

Serotonin is a vasoconstricting hormone normally produced by the argentaffin cells of the gastrointestinal tract. The
The principal function of the cells is to regulate smooth muscle contraction and peristalsis. 5-hydroxyindoleacetic acid (5-HIAA) is the major urinary metabolite of serotonin. 5-HIAA assays are more useful than the parent hormone serotonin.

This urine test is conducted to diagnose the presence of a functioning carcinoid tumor, which can be shown by significant elevations of 5-HIAA. Excess amounts of 5-HIAA are produced by most carcinoid tumors. Carcinoid tumors produce symptoms of flushing, hepatomegaly, diarrhea, bronchospasm, and heart disease.

**Reference Values**

**Normal**

Qualitative: Negative
Quantitative: 2–7 mg/24 hours or 11–37 µmol/day

**Procedure**

1. Do not allow the patient to eat any bananas, pineapple, tomatoes, eggplants, plums, or avocados for 48 hours before or during the 24-hour test because these foods contain serotonin.
2. Properly label a 24-hour urine container that contains the preservative (acid).
3. Discontinue the following drugs 48 hours before sample collection: acetaminophen, salicylates, phenacetin, naproxen, imipramine, and monoamine oxidase inhibitors.
4. Follow general directions for 24-hour urine collection (see Long-Term, Timed Urine Specimen, page 171).
5. Record exact starting and ending times of the collection on the specimen container and in the patient's health care record.
6. Send the completed specimen to the laboratory.

**Clinical Implications**

1. Levels > 25 mg/24 hours or > 131 µmol/day indicate large carcinoid tumors, especially when metastatic:
   a. Ileal tumors
   b. Pancreatic tumors
   c. Duodenal tumors
   d. Biliary tumors
2. **Increased** urine 5-HIAA is found in:
   a. Ovarian carcinoid tumor
   b. Nontropical sprue
   c. Bronchial adenoma (carcinoid type)
   d. Malabsorption
   e. Celiac disease
   f. Whipple's disease
   g. Oat cell cancer of respiratory system
3. **Decreased** urine 5-HIAA is found in:
   a. Depressive illness
   b. Small intestine resection
   c. Phenylketonuria (PKU)
   d. Hartnup's disease
   e. Mastocytosis

**Interfering Factors**

1. False-positive results occur with:
   a. Ingestion of banana, pineapple, plum, walnut, eggplant, tomato, chocolate, and avocado, because of their serotonin content
   b. Many drugs (see Appendix J)
   c. After surgery (surgical stress)
2. False-negative results can be caused by specific drugs that depress 5-HIAA production.

**Interventions**

**Pretest Patient Preparation**

1. Instruct the patient about test purpose and procedure for collection of the 24-hour urine specimen. Written instructions may be helpful.
2. Encourage intake of food and water. Foods high in serotonin content must not be eaten for 48 hours before or during the test.
3. If possible, no drugs should be taken for 72 hours before the test nor during the test (especially aforementioned drugs), including over-the-counter drugs.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Resume normal diet and medications when test is completed.
2. Interpret test outcome and counsel appropriately about abnormal 5-HIAA levels.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Clinical Alert**

A serum serotonin assay may detect some carcinoids missed by the urine 5-HIAA assay.
Urine Vanillylmandelic Acid (VMA); Catecholamines (24-Hour)

The principal substances formed by the adrenal medulla and excreted in urine are VMA, epinephrine, norepinephrine, metanephrine, and normetanephrine. These substances contain a catechol nucleus together with an amine group and therefore are referred to as catecholamines. Most of these hormones are changed into metabolites, the principal one being 3-methoxy-4-hydroxymandelic acid, known as vanillylmandelic acid, or VMA.

VMA is the primary urinary metabolite of the catecholamine group. It has a urine concentration 10 to 100 times greater than the concentrations of the other amines. It is also fairly simple to detect; methods used for catecholamine determination are much more complex.

This 24-hour urine test of adrenomedullary function is done primarily when pheochromocytoma, a tumor of the chromaffin cells of the adrenal medulla, is suspected in a patient with hypertension.

The assay for pheochromocytoma is most valuable when a urine specimen is collected during a hypertensive episode. Because a 24-hour urine collection represents a longer sampling time than a symptom-directed serum sample, the 24-hour urine test may detect a pheochromocytoma missed by a single blood level determination.

Reference Values

<table>
<thead>
<tr>
<th>Normal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adults</td>
</tr>
<tr>
<td>VMA: up to 9 mg/24 hours or up to 45 µmol/day</td>
</tr>
<tr>
<td>Catecholamines (total): &lt;100 µg/day or &lt;591 nmol/day</td>
</tr>
<tr>
<td>Epinephrine: 0–20 µg/24 hours or 0–109 nmol/day</td>
</tr>
<tr>
<td>Metanephrine: 74–297 µg/24 hours or 375–1506 nmol/day</td>
</tr>
<tr>
<td>Norepinephrine: 15–80 µg/24 hours or 89–473 nmol/day</td>
</tr>
<tr>
<td>Normetanephrine: 105–354 µg/24 hours or 573–1933 nmol/day</td>
</tr>
<tr>
<td>Dopamine: 65–400 µg/24 hours or 420–2612 nmol/day</td>
</tr>
</tbody>
</table>

Children's levels are different from those of adults. Check with your laboratory for values in children.

NOTE

Different laboratories report values in different units—this should be kept in mind when analyzing results.

Procedure

1. Properly label a 24-hour container with acid preservative and refrigerate the container or keep it on ice.
2. Follow general instructions for 24-hour urine collection (see Long-Term, Timed Urine Specimen, page 171).
3. Record exact starting and ending times of the collection on the specimen container and in the patient's health care record.
4. Send the specimen to the laboratory.

Clinical Implications

1. Increased urine VMA occurs as follows:
   a. High levels in pheochromocytoma
   b. Slight to moderate elevations in
      1. Neuroblastoma
      2. Ganglioneuroma
      3. Ganglioblastoma
      4. Carcinoid tumor (some cases)
2. Increased urine catecholamines are found in:
   a. Pheochromocytoma
      1. Norepinephrine, >170 mg/24 hours or >170 mg/day
      2. Epinephrine, >35 mg/24 hours or >35 mg/day
   b. Neuroblastomas
   c. Ganglioneuromas
   d. Myocardial infarction (acute)
   e. Hypothyroidism
   f. Diabetic acidosis
   g. Long-term manic-depressive states
3. Decreased urine catecholamines are found in:
   a. Diabetic neuropathy
   b. Parkinson's disease

Interfering Factors

1. Increased urine VMA and catecholamines are caused by:
   a. Hypoglycemia—for this reason, the test should not be scheduled while the patient is receiving nothing by mouth.
   b. Many foods, such as the following:
      1. Caffeine-containing products (eg, tea, coffee, cocoa, carbonated drinks)
      2. Vanilla
      3. Fruit, especially bananas
      4. Licorice
   c. Many drugs cause increased VMA levels, especially reserpine, a-methyldopa, levodopa, monoamine oxidase
inhibitors, sinus and cough medicines, bronchodilators, and appetite suppressants.

d. Exercise, stress, smoking, and pain cause physiologic increases of catecholamines.

e. Heavy alcohol intake increases catecholamine levels.

2. Falsely decreased levels of VMA and catecholamines are caused by:

a. Alkaline urine  
b. Uremia (causes toxicity and impaired excretion of VMA)

c. Radiographic contrast agents—for this reason, an intravenous pyelogram should not be scheduled before a VMA test.

d. Certain drugs (see Appendix J)

Interventions

Pretest Patient Preparation

1. Instruct the patient about the test purpose and the procedure for collection of the 24-hour urine specimen. A written reminder may be helpful, especially regarding restricted foods.

2. Explain diet and drug restrictions. Diet restrictions vary among laboratories, but coffee, tea, bananas, cocoa products, vanilla products, and aspirin are always excluded for 3 days (2 days before and 1 day during specimen collection).

3. Many laboratories require that all drugs be discontinued for 1 week before testing.

4. Encourage adequate rest, food, and fluids.

5. Stress, strenuous exercise, and smoking should be avoided during the test.

6. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. The patient may resume pretest diet, drugs, and activity when the test is completed.

2. Interpret test outcomes and counsel appropriately.

3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Urine Porphyrins and Porphobilinogens (24-Hour and Random); \(\beta\)-Aminolevulinic Acid (ALA, \(\beta\)-ALA)

Porphyrins are cyclic compounds formed from \(\beta\)-aminolevulinic acid (\(\beta\)-ALA), which plays a role in the formation of hemoglobin and other hemoproteins that function as carriers of oxygen in the blood and tissues. In health, insignificant amounts of porphyrin are excreted in the urine. However, in certain conditions, such as porphyria (disturbance in metabolism of porphyrin), liver disease, and lead poisoning, increased levels of porphyrins and \(\beta\)-ALA are found in the urine. Disorders in porphyrin metabolism also result in increased amounts of porphobilinogen in urine. The most common signs and symptoms of acute intermittent porphyria are abdominal pain, photosensitivity, sensory neuropathy, or psychosis. Patients with the porphyrias may pass urine that is pink, port wine, or burgundy colored.

When urine is tested for the presence of porphyrins, porphobilinogen, and/or ALA, it is also given the black-light screening test (Wood's light test). Porphyrins are fluorescent when exposed to black or ultraviolet light. See Chapter 2 for other tests for porphyria.

This test is used to diagnose porphyrias and lead poisoning in children. The following is a summary of laboratory findings for various porphyrias.

**Congenital Erythropoietic Porphyria.** Elevations of urine uroporphyrin and coproporphyrin occur, with the former exceeding the latter. Lesser amounts of hepta, hexa, and penta carboxyporphyrins are secreted. ALA and porphobilinogen levels are normal.

**Acute Intermittent Porphyria.** Porphobilinogen and \(\beta\)-ALA are elevated in acute attacks, and small increases of urine uroporphyrin and coproporphyrin may be found. During periods of latency, the values are normal.

**Porphyria Cutanea Tarda.** A more common form of porphyria—increased uroporphyrins, uroporphyrinogen, and hepta carboxyporphyrins.

**Protoporphyria.** Mild disease, which mainly has the clinical symptoms of solar urticaria and solar eczema (exposure to sunshine). Increased fecal protoporphyrin.

**Hereditary Coproporphyria.** Urine coproporphyrin and porphobilinogen are markedly increased during acute attacks; increases of urine uroporphyrin may also be found.

**Variegate Porphyria.** In acute attacks, results are similar to those seen in acute intermittent porphyria. Porphobilinogen and \(\beta\)-ALA usually return to normal between attacks. Urine coproporphyrin exceeds uroporphyrin excretion during acute attacks.

**Chemical Porphyrias.** ("Intoxication porphyria.") Porphyrinogenic chemicals include certain halogenated hydrocarbons, which cause increased uroporphyrin levels in the urine. Also increased are ALA, coproporphyrin, and porphobilinogen.

**Lead Poisoning.** \(\beta\)-ALA levels exceed those of porphobilinogen, which may remain normal. In children, ALA secretion in urine is more sensitive than blood lead levels.

Reference Values

*Normal*
Porphobilinogens
Random specimen: 0–2.0 mg/L or negative or 0–8.8 µmol/L
24-hour specimen: 0–1.5 mg/24 hours or 0–6.6 mg/day
?- ALA Random specimen: 0–4.5 mg/L or 0–34 µmol/L
24-hour specimen: 1.5–7.5 mg/24 hours or 11.4–57.2 µmol/day (Table 3.11)

Table 3.11 Specimen Values

<table>
<thead>
<tr>
<th>Porphyrins *</th>
<th>Male</th>
<th>Female</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(µg/24 h) (nmol/d)</td>
<td>(µg/24 h) (nmol/d)</td>
</tr>
<tr>
<td>Random specimen</td>
<td>Negative</td>
<td>Negative</td>
</tr>
<tr>
<td>24-h Specimens:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Uroporphyrin</td>
<td>8–44</td>
<td>10–53</td>
</tr>
<tr>
<td>Coproporphyrin</td>
<td>10–109</td>
<td>15–167</td>
</tr>
<tr>
<td>Heptacarboxypherin</td>
<td>0–12</td>
<td>0–9</td>
</tr>
<tr>
<td>Pentacarboxypherin</td>
<td>0–4</td>
<td>0–3</td>
</tr>
<tr>
<td>Hexacarboxypherin</td>
<td>0–5</td>
<td>0–5</td>
</tr>
</tbody>
</table>

*Total porphyrins: 20–121 µg/L or 24–146 nmol/L.

Procedure

1. Properly label a 24-hour clean-catch urine container.
2. Provide refrigeration or icing. The specimen must be kept protected from exposure to light. Check with your laboratory regarding the need for preservatives.
3. Follow general instructions for 24-hour urine collection (see Long-Term, Timed Urine Specimen, page 171).
4. Record exact starting and ending times on the specimen container and in the patient’s health care record.
5. Send the specimen to the laboratory.
6. Obtain midmorning or midafternoon specimens for random tests because it is more likely that the patient will excrete porphyrins at those times. Transport the specimen to the laboratory immediately. Protect the specimen from light.
7. Observe and record the urine color. If porphyrins are present, the urine may appear amber-red or burgundy in color, or it may vary from pale pink to almost black. Some patients excrete urine of normal color that turns dark after standing in the light.

Clinical Implications

1. Increased urine porphobilinogen occurs in:
   a. Porphyria (acute intermittent type)
   b. Variegate porphyria
   c. Hereditary coproporphyria
   d. See pages 246–247 for list of other porphyrias.
2. Increased fractionated porphyrins occur in:
   a. Acute intermittent porphyria
   b. Congenital erythropoietic porphyria
   c. Hereditary porphyria
   d. Variegate porphyria
   e. Chemical porphyria caused by heavy-metal poisoning or carbon tetrachloride
   f. Lead poisoning
   g. Viral hepatitis
   h. Cirrhosis (alcoholism)
   i. Newborn of mother with porphyria
   j. Congenital hepatic porphyria
3. Increased urine ?- ALA can occur in:
   a. Acute intermittent porphyria (acute phase)
   b. Variegate porphyria (during crisis)
   c. Hereditary coproporphyria
   d. Lead poisoning does not increase urine ?-ALA until serum lead levels reach > 40 µg/dL; urine ?-ALA may remain elevated for several months after control of lead exposure.
   e. Congenital hepatic porphyria
   f. Slight increase in pregnancy, diabetic acidosis
4. Decreased urine ?- ALA is found in alcoholic liver disease

Clinical Alert

Porphobilinogen is not increased in lead poisoning.

Interfering Factors

1. Oral contraceptives and diazepam can cause acute porphyria attacks in susceptible patients.
2. Alcohol ingestion interferes with the test.
3. Many other drugs, especially phenazopyridine, procaine, sulfamethoxazole, and the tetracyclines, interfere with the
Interventions

**Pretest Patient Preparation**

1. Instruct the patient about the purpose and procedure of collection a 24-hour urine specimen. A written reminder may be helpful.
2. Allow food and fluids, but alcohol and excessive fluid intake should be avoided during the 24-hour collection.
3. If possible, discontinue all drugs for 2 to 4 weeks before the test so that results will be accurate.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. The patient may resume normal activities and medications.
2. Interpret test outcomes and counsel appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

This test should not be ordered for patients receiving Donnatal or other barbiturate preparations. However, if intermittent porphyria is suspected, the patient should take those medications according to prescribed protocols because these drugs may provoke an attack of porphyria.

Urine Amylase Excretion and Clearance (Random, Timed Urine, and Blood)

Amylase is an enzyme that changes starch to sugar. It is produced in the salivary glands, pancreas, liver, and fallopian tubes and is normally excreted in small amounts in the urine. If the pancreas or salivary glands are inflamed, much more of the enzyme enters the blood and, consequently, more amylase is excreted in the urine.

This test of blood and urine indicates pancreatic function and is done to differentiate acute pancreatitis from other causes of abdominal pain, epigastric discomfort, or nausea and vomiting.

In patients with acute pancreatitis, the urine often shows a prolonged elevation of amylase, compared with a short-lived peak in the blood. Moreover, urine amylase may be elevated when blood amylase is within normal range, and, conversely, the blood amylase may be elevated when the urine amylase is within normal range. The advantage of the amylase-creatinine clearance test is that it can be done on a single random urine specimen and a single serum sample instead of having to wait for a 2- or 24-hour urine collection. The ratio is increased in certain conditions other than acute pancreatitis, such as diabetic acidosis and renal insufficiency. Although the usefulness of this test in pancreatic disease has been questioned, it can be helpful to screen for macroamylasia.

**Reference Values**

**Normal**

Amylase/creatinine clearance ratio: 1%–4% or 0.01–0.04 clearance fraction

This is a ratio calculated as follows:

\[
\text{Urine amylase (U) × Serum creatinine (mg/dL) × 100}
\]

\[
\text{Serum amylase (U) × Urine creatinine (mg/dL)}
\]

**Urine Amylase**

- 2-hour specimen: 2–34 U or 16–283 nkat/hour
- 24-hour specimen: 24–408 U or 400–6800 nkat/day

Values vary according to laboratory methods used. Check with your lab.

**NOTE**

kat = katal, which is a measure of enzyme activity.

**Procedure**

For the amylase clearance test, a venous blood sample of 4 mL must be collected at the same time the random urine specimen is obtained.

1. Order a random, 2-hour, or 24-hour timed urine specimen. A 2-hour specimen is usually collected.
2. Refrigerate the urine specimen. Amylase is unstable in acidic urine. The pH must be adjusted to pH > 7.0.
3. Follow general instructions for the appropriate urine collection.
4. Record exact starting and ending times on the specimen container and on the health care record. This is very important for calculation of results.
5. Send the specimen to the laboratory.

**Clinical Implications**

1. Amylase/creatinine clearance is increased in:
   a. Pancreatitis, pancreatic cancer
   b. Diabetic ketoacidosis (some patients)
   c. Toxemia of pregnancy, hyperemesis of pregnancy
   d. Renal insufficiency
2. Amylase/creatinine clearance is decreased in macroamylasia.
3. Urine amylase is increased in:
   a. Pancreatitis
   b. Parotitis
   c. Intestinal obstruction
   d. Diabetic ketoacidosis
   e. Strangulated bowel
   f. Pancreatic cyst
   g. Peritonitis
   h. Biliary tract disease
   i. Some lung and ovarian tumors
4. Urine amylase is decreased in:
   a. Pancreatic insufficiency
   b. Advanced cystic fibrosis
   c. Severe liver disease
   d. Renal failure
   e. Macroamylasemia

Interfering Factor

1. Acid pH—decreases urine amylase.
2. Some drugs produce increased amylase and possible pancreatitis.

Interventions

Pretest Patient Preparation

1. Instruct the patient about the test purpose and procedure for urine specimen collection. A written instruction sheet may be helpful.
2. Encourage fluids, if they are not restricted.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

Follow-up calcium levels should be checked in fulminating pancreatitis because extremely low calcium levels can occur.

Phenyketonuria (PKU); Urine Phenylalanine (Random Urine and Blood)

Routine blood and urine tests are done on newborns to detect phenylketonuria (PKU), an inherited disease that can lead to mental retardation and brain damage if untreated. This disease is characterized by a lack of the enzyme that converts phenylalanine, an amino acid, to tyrosine, which is necessary for normal metabolic function. Because dietary phenylalanine is not converted to tyrosine, phenylalanine, phenylpyruvic acid, and other metabolites accumulate in blood and urine. Tyrosine and the derivative catecholamines are deficient, which results in mental retardation. Both sexes are affected equally, with most cases occurring in persons of northern European ancestry.

This test is used for newborns to detect the metabolic disorder hyperphenylalaninemia. If untreated, this disorder can lead to mental retardation. Dietary restrictions of phenylalanine have shown good results.

Reference Values

**Normal**

Blood: <2 mg/dL (2–5 days after birth) or <121 µmol/L

Urine: Negative dipstick (detects phenylalanine in range of 5–10 mg/dL or 302–605 µmol/L)

24-hour urine: 1.2–1.7 mg/24 hours (10 days to 7 weeks after birth) or 7.2–10.3 µmol/day

Adults: <16.5 mg/24 hours or <100 µmol/day

Children (3–12 years old): 4.0–17.5 mg/24 hours or 24–106 µmol/day

**Procedure**

Collecting the Blood Sample

1. Cleanse the skin with an antiseptic and pierce the infant's heel with a sterile disposable lancet.
2. Support the infant, if bleeding is slow, so that the blood flows by means of gravity while spotting the blood with filter paper.
3. Fill the circles on the filter paper completely. This can best be done by placing one side of the filter paper against the infant's heel and watching for the blood to appear on the other side of the paper until it completely fills the circle.
4. Do not touch blood circles until they are completely dry. Keep in cool, dry area.
5. Transport samples to testing site within 12 to 24 hours.
6. Confirm all positive filter paper tests with a quantitative blood or urine test.

Collecting the Urine Sample in Nursery or at Home
1. Dip the reagent strip into a fresh sample of urine or press it against a wet diaper (phenylalanines and phenylpyruvic acid may not appear in urine until the infant is 2 to 3 weeks of age).
2. After exactly 30 seconds, compare the strip with a color chart according to manufacturer's directions.
3. Salicylates and phenothiazine may cause abnormal color reactions.
4. All positive tests must be confirmed with a quantitative chemical test.

**Clinical Implications** Increased phenylalanine is found in:

1. Hyperphenylalaninemia. In a positive test for PKU, the blood phenylalanine is > 15 mg/dL or > 907 µmol/L. Blood tyrosine is < 5 mg/dL or < 276 µmol/L; it is never increased in PKU.
2. Obesity
3. In low-birth-weight or premature infants, transient hyperphenylalaninemia, along with transient hypertyrosinemia, may occur.

**Interfering Factors**

1. Premature infants, those weighing < 2.3 kg (<5 pounds), may have elevated phenylalanine and tyrosine levels without having the genetic disease. This is a result of delayed development of appropriate enzyme activity in the liver (liver immaturity).
2. Antibiotics interfere with the blood assay.
3. Cord blood cannot be used for analysis.
4. Two days of protein feeding must be done before blood is taken.

**Instructions to Mothers**

1. Inform the mother about the purpose of the test and the methods of collecting the specimens.
2. Most parents are interested to know that PKU (a genetic disease in which a defective gene is passed on from each parent) was first recognized by a young mother of two mentally retarded children. She was aware that the urine of these children had a peculiar odor and, on the basis of this, was able to have a biochemist study the urine and identify phenylpyruvic acid. Her discovery led to the first successful dietary treatment, restriction of phenylalanine (eg, in milk) for those newborn babies identified as having PKU. This resulted in normal mental development of these children.
3. Interpret test outcomes and counsel regarding diet if results are positive.

**Clinical Alert**

The established standard is that all newborn infants should be tested for PKU and congenital hypothyroidism before discharge.

1. The blood test must be performed at least 3 days after birth or after the child has ingested protein (milk) for at least 24 to 48 hours.
2. Urine testing is usually done at the 4- or 6-week checkup if a blood test was not done.
3. PKU studies should be done on all infants who weigh = 2.3 kg (=5 pounds) before they leave the hospital.
4. Sick or premature infants should be tested within 7 days after birth regardless of protein intake, weight, or antibiotic therapy.

**D-Xylose Absorption (Timed Urine and Blood)**

The D-xylose test is a diagnostic measure for evaluating malabsorptive conditions and intestinal absorption of D-xylose, a pentose not normally present in the blood in significant amounts. It is passively absorbed in the proximal small bowel, passes unchanged in the liver, and is excreted by the kidneys.

This test directly measures intestinal absorption. When D-xylose (which is not metabolized by the body) is administered orally, blood and urine levels are checked for absorption rates. Absorption is normal in pancreatic insufficiency but is impaired in intestinal malabsorption. It is a reliable index of the functional integrity of the jejunum in pediatric patients.

**Reference Values**

**Normal Blood**

- 1-hour absorption of 5-g dose—infant: >15 mg/dL or >1.0 mmol/L
- 1-hour absorption of 5-g dose—child: >20 mg/dL or >1.3 mmol/L
- 2-hour absorption of 5-g dose—adult: >20 mg/dL or >1.3 mmol/L
- 2-hour absorption of 25-g dose—adult: >25 mg/dL or >1.6 mmol/L

**Urine Xylose 5-Hour Reference Range for 25-g dose**

- Child: 16%–33% of 5-g dose
- Adult: >16% of 5-g dose or >4.0 g of max (0.5 g/kg to a maximum of 25 g)
- Adult, 65 years of age and older: >14% of dose or >3.5 g of maximum

**Procedure**

1. Have the patient refrain from foods containing pentose for 24 hours before test.
2. Do not allow food or liquids by mouth for at least 8 hours before the start of the test. Pediatric patients should fast only 4 hours.
3. Have the patient void at the beginning of the test. Discard this urine.
4. Administer the oral dose of D-xylose after it has been dissolved in 100 mL of water. Adult dosage is 25 g; for
children younger than 12 years of age, a 5-g oral dose is recommended. For adults, additional water up to 250 mL should be taken at this time and another 250 mL in 1 hour. Record these times on the patient's health care record. Give no further fluids (except water) or food until the test is completed.

5. Draw a 3-mL sample of venous blood within 60 to 120 minutes later.
6. Have the patient rest quietly in one place until the test is completed.
7. Have the patient void 5 hours from the start of the test. Save all urine voided during the test.

Clinical Implications

1. Urine D-xylose is decreased in:
   a. Intestinal malabsorption
   b. Impaired renal function
   c. Small bowel ischemia
   d. Whipple's disease
   e. Viral gastroenteritis (vomiting)
   f. Bacterial overgrowth in small intestine.
2. The D-xylose test is normal in the following conditions:
   a. Malabsorption due to pancreatic insufficiency
   b. Postgastrectomy
   c. Malnutrition

Interfering Factors

1. Many drugs and antibiotics (see Appendix J)
2. Nonfasting state, treatment with hyperalimentation
3. Foods rich in pentose (fruits and preserves)
4. Vomiting of the xylose test meal (25-g dose may cause gastrointestinal distress),
5. Impaired renal function—use serum test only
6. In adults, the serum test has little value—use 5-hour urine test.

Interventions
Pretest Patient Preparation

1. Explain purpose and procedure of the test and the urine collection process. The entire 5-hour specimen must be collected.
2. The patient must fast at least 8 hours before the start of the test; children younger than 9 years of age should fast for only 4 hours.
3. Water may be taken at any time.
4. Weigh the patient to determine the proper dose of D-xylose.
5. The patient must not ingest contraindicated drugs for 1 week before the test.
6. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Normal food, fluids, and activities can be resumed.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

Nausea, vomiting, and diarrhea may result from ingestion of the D-xylose. If vomiting occurs, the test is invalid and must be repeated. A 5-gram dose is more tolerated but is less sensitive.

Urine Creatinine; Creatinine Clearance (Timed Urine and Blood)

Creatinine is a substance that, in health, is easily excreted by the kidney. It is the byproduct of muscle energy metabolism and is produced at a constant rate according to the muscle mass of the individual. Endogenous creatinine production is constant as long as the muscle mass remains constant. Because all creatinine filtered by the kidneys in a given time interval is excreted into the urine, creatinine levels are equivalent to the glomerular filtration rate (GFR). Disorders of kidney function prevent maximum excretion of creatinine. The creatinine clearance test is part of most batteries of quantitative urine tests. Creatinine clearance is measured together with other urinary components in order to interpret the overall excretion rate of the various urinary components.

The creatinine clearance test is a specific measurement of kidney function, primarily glomerular filtration. It measures the rate at which the kidneys clear creatinine from the blood. In a broad sense, clearance of a substance may be defined as the imaginary volume (in milliliters) of plasma from which the substance would have to be completely extracted in order for the kidney to excrete that amount in 1 minute. In addition to estimating the GFR, this test is used to evaluate renal function in patients.

Because the excretion of creatinine in a given person is relatively constant, the 24-hour urine creatinine level is used as a check on the completeness of a 24-hour urine collection. It is of no help in the evaluation of renal function unless it is done as part of a creatinine clearance test.

Reference Values

Normal
Urine creatinine, men: 14–26 mg/kg/24 hours or 124–230 µmol/kg/day
Urine creatinine, women: 11–20 mg/kg/24 hours or 97–177 µmol/kg/day
Blood creatinine: 0.8–1.2 mg/dL or 71–106 µmol/L (Table 3.12)

Table 3.12 Mean Creatinine Clearance (mL/min/1.73 m²) *

<table>
<thead>
<tr>
<th>Age (y)</th>
<th>Men</th>
<th>Women</th>
</tr>
</thead>
<tbody>
<tr>
<td>20–30</td>
<td>90–140 or 0.8–1.3 mL/sec/m²</td>
<td>72–110 or 0.69–1.06 mL/sec/m²</td>
</tr>
<tr>
<td>30–40</td>
<td>59–137 or 0.5–1.3 mL/sec/m²</td>
<td>71–121 or 0.68–1.17 mL/sec/m²</td>
</tr>
</tbody>
</table>

*Values slowly increase to adult levels, then slowly decrease each decade thereafter (the decrease per decade is approximately 6.5 mL/min/1.73 m² or 0.06 mL/sec/m²).

Procedure
1. Properly label a 12-hour or 24-hour urine container.
2. Refrigerate or ice the specimen.
3. Follow general instructions for 24-hour urine collection (see Long-Term, Timed Urine Specimen, page 171).
4. Record exact starting and ending times on the specimen container and in the patient's health care record.
5. Send the entire specimen to the laboratory.
6. Obtain a 5-mL venous blood sample for creatinine when the test begins.
7. Record the patient's height and weight on the container and in the patient's health care record. Creatinine clearance values are based on the body surface area, and these values are needed to calculate the surface area.
8. Ensure that the patient is adequately hydrated throughout the test to provide proper urine flow.

Clinical Implications
1. Decreased creatinine clearance is found in any condition that decreases renal blood flow:
   a. Impaired kidney function, intrinsic renal disease, glomerulonephritis, pyelonephritis, nephrotic syndrome, acute tubular dysfunction, amyloidosis, interstitial nephritis
   b. Shock, dehydration
   c. Hemorrhage
   d. Chronic obstructive lung disease
   e. Congestive heart failure
2. Increased creatinine clearance is found in:
   a. State of high cardiac output
   b. Pregnancy
   c. Burns
   d. Carbon monoxide poisoning
3. Increased urine creatinine is found in:
   a. Acromegaly
   b. Gigantism
   c. Diabetes mellitus
   d. Hypothyroidism
4. Decreased urine creatinine is found in:
   a. Hyperthyroidism
   b. Anemia
   c. Muscular dystrophy
   d. Polymyositis, neurogenic atrophy
   e. Inflammatory muscle disease
   f. Advanced renal disease, renal stenosis
   g. Leukemia

Interfering Factors
1. Exercise may increase creatinine clearance and urine creatinine.
2. Pregnancy substantially increases creatinine clearance.
3. Many drugs decrease creatinine clearance (see Appendix J).
4. The creatinine clearance overestimates the GFR when there is severe renal impairment. The serum creatinine is more indicative of the GFR in this situation.
5. A diet high in meat may elevate the urine creatinine concentration.
6. Proteinuria and advanced renal failure make creatinine clearance an unreliable method for determining GFR.

Clinical Alert
Determination of urine creatinine is of little value for evaluating renal function unless it is done as part of a creatinine clearance test.

Interventions
Pretest Patient Preparation
1. Instruct the patient about the purpose and procedure of the test and urine specimen collection. A written reminder may be helpful.
2. Allow food and encourage fluids for good hydration. Large urine volumes ensure optimal test results. Avoid tea and
coffee (diuretics).
3. Avoid vigorous exercise during the test.
4. Drugs affecting the results should be stopped beforehand (especially adrenocorticotropic hormone [ACTH], cortisone, or typoxine). Check with physician.
5. Avoid eating large amounts of meat. Check with physician.
6. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. The patient may resume normal food, fluids, and activity.
2. Interpret test outcomes and monitor appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Urine Cystine (Random and 24-Hour)

Cystinuria is a condition characterized by increased amounts of the amino acid cystine in the urine. The presence of increased urinary cystine is caused not by a defect in the metabolism of cystine but rather by the inability of the renal tubules to reabsorb cystine filtered by the glomeruli. The tubules fail to reabsorb not only cystine but also lysine, ornithine, and arginine; this rules out the possibility of an error in metabolism, even though the condition is inherited.

These urine tests are useful for the differential diagnosis of cystinuria, an inherited disease characterized by bladder calculi (cystine has low solubility). Patients with cystine stones face recurrent urolithiasis and repeated urinary infections.

Reference Values

Normal
Random specimen: Negative
24-hour specimen, adult: <38 mg/24 hours or <316 µmol/day
24-hour specimen, child: 5–31 mg/24 hours or 42–258 µmol/day

Procedure

1. Obtain a random 20-mL urine specimen for a qualitative screening test.
2. When collecting a 24-hour urine specimen, the container needs a preservative (toluene). Follow general procedures for a 24-hour urine specimen (see Long-Term, Timed Urine Specimen, page 171).

Clinical Implications

1. Urine cystine is increased in cystinuria (up to 20 times normal).
2. Urine cystine is decreased in burn patients.

Clinical Alert

1. Cystinosis, a different entity from cystinuria, is not detected by cystine studies. Most patients with infantile nephropathic cystinosis have neurologic defects that become apparent in infancy. Failure to thrive and renal dysfunction are evidence of this disease.
2. Patients with cystinosis have a defect in renal tubular reabsorption that develops into Fanconi’s syndrome, which leads to a generalized amino aciduria. Cystine is elevated in the urine in the same proportion as all amino acids; the concentration is not high enough to form cystine stones. Plasma cystine is normal, but cystine is elevated in kidneys, eyes, spleen, and bone marrow; for purposes of diagnosis, it is usually measured in WBCs.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure for timed urine collection.
2. See Chapter 1 guidelines regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Urine Hydroxyproline (Timed Urine and Blood)

Hydroxyproline is an amino acid found only in collagen. It increases during periods of rapid growth, in bone diseases, and in some endocrine disorders. Urine hydroxyproline is almost entirely peptide bound, and only 10% is in the free form.

Total hydroxyproline is considered to be a marker for bone resorption because 50% of human collagen resides in bone. This test indicates the presence of reabsorption of bone collagen in various disorders and evaluates the degree of destruction from primary or secondary bone tumors. Free hydroxyproline is used as an aid to diagnose hydroxyprolinemia, a rare genetic disorder characterized by mental retardation and thrombocytopenia.

NOTE

During periods of rapid growth in early childhood and in puberty, total hydroxyproline is greatly increased.
Reference Values

**Normal Urine:**
- **Total hydroxyproline (24-hour):** 15–45 mg/24 hours or 115–345 µmol/day
- **Adult females:** 0.4–2.9 mg/2-hour specimen or 3–22 µmol/2 hours
- **Adult males:** 0.4–5.0 mg/2-hour specimen or 3–38 µmol/2 hours
- **Children < 5 years old:** 100–400 µg/mg creatinine or 86–345 mmol/day
- **Children 5–12 years:** 100–150 µg/mg creatinine or 86–129 mmol/day

**Blood (Plasma)—Free Hydroxyproline:**
- **Newborn:** 0.52 ± 0.52 mg/dL or 40 ± 40 µmol/L
- **Child (male):** <0.66 mg/dL or <50 µmol/L
- **Child (female):** <0.58 mg/dL or <44 µmol/L
- **Adult (male):** <0.55 mg/dL or <42 µmol/L
- **Adult (female):** <0.45 mg/dL or <34 µmol/L

**Procedure**

1. Obtain a 2-hour specimen after the patient has fasted overnight (preferred method).
2. Notify the laboratory of the patient's age and sex.
3. If ordered, collect a 24-hour urine specimen. No preservative is required, but the specimen must be refrigerated or placed on ice.
4. Follow 24-hour urine collection procedures. The laboratory will record the total 24-hour volume.
5. Note that the preferred method of testing in the first few months of life is blood sampling (free hydroxyproline only for genetic screening).

**Clinical Implications**

1. **Free hydroxyproline is increased in:**
   - a. Hydroxyprolinemia, a hereditary autosomal recessive condition (very rare)
   - b. Familial iminoglycinuria, also inherited and rare
2. **Total hydroxyproline is increased in:**
   - a. Hyperparathyroidism, hyperthyroidism
   - b. Paget's disease—measures the severity and the response to treatment
   - c. Marfan's syndrome, acromegaly
   - d. Osteoporosis
   - e. Myeloma
   - f. Severe burns
3. **Total hydroxyproline is increased in:**
   - a. Hypopituitarism
   - b. Hypothyroidism
   - c. Hypoparathyroidism

**Interfering Factors**

1. Gelatin may affect test results (false-positive test). For best results, the patient should be on a nonprotein diet.
2. Bed rest increases values
3. Pregnancy increases values

**Interventions**

**Pretest Patient Preparation**

1. Explain the test purpose and procedure for a timed urine collection. Fasting and special fluid requirements before testing are often required for a 24-hour timed procedure. Check with laboratory.
2. Avoid gelatin foods for several days before the test.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. The patient may resume normal diet and activity.
2. Interpret test outcomes and counsel appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Urine Lysozyme (Random, 24-Hour Urine, and Blood)**

Lysozyme (muramidase) in blood or urine is a bacteriolytic enzyme that comes from degradation of granulocytes and monocytes, but not lymphocytes. It is increased in leukemia owing to degradation of granulocytic or monocytic cells.

This blood and urine test differentiates acute myelogenous or monocytic leukemia from acute lymphatic leukemia. It is useful to monitor the response to treatment of acute myelogenous and active monocytic leukemia.

**Reference Values**

**Normal**

- **Blood plasma:** 0.4–1.3 mg/dL or 4–13 mg/L
- **Urine, 24-hour specimen:** 0–3 mg/24 hours

Reference values are not established for random urine specimens.

**Procedure**
1. Collect a 5-mL EDTA-anticoagulated blood sample or urine specimen.
2. Follow general instructions for 24-hour urine collections. 
   Transport the sample to the laboratory immediately after collection.

**Clinical Implications**

1. *Lysozyme levels are increased in:*
   a. Acute myelogenous leukemia (granulocytic)
   b. Acute monocytic leukemia
   c. Malignant histiocytosis
2. *Lysozyme levels may be increased in:*
   a. Renal disorders and transplant rejection
   b. Tuberculosis
   c. Sarcoidosis (sarcoid lymph nodes)
   d. Crohn's disease
   e. Polycythemia vera
3. *Lysozyme levels are normal* in acute lymphatic leukemia.
4. *Lysozyme levels are decreased* in neutropenia with hypoplasia of bone marrow.

**Interventions**

**Pretest Patient Preparation**

1. Explain the test purpose and procedure for urine or blood collection.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Urine Amino Acids, Total and Fractions (Random, 24-Hour Urine, and Blood)**

Many abnormalities in amino acid transport or metabolism can be detected by physiologic fluid analysis (urine, plasma, or cerebrospinal fluid). Free amino acids are found in urine and in acid filtrates of protein-containing fluids. Urine is used for initial screening of inborn metabolic errors. Both transport errors and metabolic errors can be detected by changes in observed amino acid patterns. In many cases, metabolic errors are detected when amino acid or metabolite exceeds its renal threshold.

This test is useful for the diagnosis and monitoring of inborn errors of metabolism and transport in cases of suspected genetic abnormalities in patients with mental retardation, reduced growth, or other unexplained symptoms. More than 50 aminoacidopathies are now recognized.

**Reference Values**

**Normal**
Urine and blood amino acid values are age dependent.

**Procedure**

1. Obtain a fasting blood specimen.
2. Collect a random 24-hour timed urine specimen. Keep the specimen refrigerated or on ice.

**Clinical Implications**

1. *Total plasma amino acids are increased in:*
   a. Specific aminoacidopathies (Table 3.13)

<table>
<thead>
<tr>
<th>Aminoacidurias</th>
<th>Amino Acids Increased in Urine and Blood</th>
<th>Presence of Abnormal Enzymes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenylketonuria</td>
<td>Phenylketonuria</td>
<td>Phenylamine hydroxylase</td>
</tr>
<tr>
<td>Tyrosinosis</td>
<td>Tyrosine</td>
<td>p-Hydroxyphenyl-pyruvic acid oxidase</td>
</tr>
<tr>
<td>Histidinemia</td>
<td>Histidine</td>
<td>Histidase</td>
</tr>
<tr>
<td>Maple syrup urine disease</td>
<td>Valine, leucine, and isoleucine</td>
<td>Branched chain ketoacid decarboxylase</td>
</tr>
<tr>
<td>Hypervalinemia</td>
<td>Valine</td>
<td>Probably valine transaminase</td>
</tr>
<tr>
<td>Hyperglycinemia</td>
<td>Glycine (lysine on high-protein diet)</td>
<td>Increased glycine and propionic acid</td>
</tr>
<tr>
<td>Hyperprolinemia</td>
<td>Proline</td>
<td>Proline oxidase</td>
</tr>
<tr>
<td>Type I</td>
<td></td>
<td>Pyrroline-5-carboxylate dehydrogenase</td>
</tr>
<tr>
<td>Type II</td>
<td></td>
<td>Hydroxyproline oxidase</td>
</tr>
<tr>
<td>Hydroxyprolinemia</td>
<td>Hydroxyproline</td>
<td></td>
</tr>
</tbody>
</table>
Homocystinuria  Methionine, homocystine  Cystathionine synthetase
Hyperlysinemia  Lysine  Lysine-a-ketoglutarate reductase
Citrullinemia  Citrulline  Argininosuccinic acid synthetase
Alkaptonuria  Homogentisic acid (2,5-dihydroxyphenylacetic acid); no abnormal amino acid  Homogentisic acid oxidase
Oasthouse urine disease  Methionine, phenylalanine, valine, leucine, isoleucine, and tyrosine, and also a-hydroxybutyric acid in urine  Possibly methionine malabsorption syndrome

b. Secondary causes
1. Diabetes with ketosis
2. Malabsorption
3. Hereditary fructose intolerance
4. Conditions with severe brain damage
5. Reye's syndrome
6. Acute and chronic renal failure
7. Eclampsia
8. Specific aminoacidopathies

2. Total plasma amino acids are decreased in:
   a. Adrenocortical hyperfunction
   b. Huntington's chorea
   c. Phlebotomus fever
   d. Nephritic syndrome
   e. Rheumatoid arthritis
   f. Hartnup's disease

3. Total urine amino acids are increased in specific aminoacidurias (see Table 3-13).
4. Absence of amino acids occurs as listed in Table 3-14.

<table>
<thead>
<tr>
<th>Disease</th>
<th>Amino Acids in Urine</th>
<th>Presence of Abnormal Enzyme</th>
</tr>
</thead>
<tbody>
<tr>
<td>Argininosuccinic aciduria</td>
<td>Argininosuccinic acid (also citrulline)</td>
<td>Argininosuccinase</td>
</tr>
<tr>
<td>Cystathioninuria</td>
<td>Cystathionine</td>
<td>Cystathioninases</td>
</tr>
<tr>
<td>Homocystinuria</td>
<td>Homocystine</td>
<td>Cystathionine synthetase</td>
</tr>
<tr>
<td>Hypophosphatasia</td>
<td>Phosphoethanolamine</td>
<td>Serum alkaline phosphate</td>
</tr>
</tbody>
</table>

5. Renal transport aminoacidurias include the elements listed in Table 3-15.

<table>
<thead>
<tr>
<th>Disease</th>
<th>Amino Acids in Urine</th>
<th>Abnormality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cystinuria</td>
<td>Cystine; lysine; arginine, ornithine (basic amino acids)</td>
<td>Incomplete absorption of cystine, lysine, arginine, ornithine</td>
</tr>
<tr>
<td>Hartnup's disease</td>
<td>Monoaminomonocarboxylic (neutral) amino acids (proline, glycine, hydroxyproline, and methionine not increased)</td>
<td>Incomplete absorption of monoaminomonocarboxyamino acids</td>
</tr>
<tr>
<td>Glycinuria, renal type</td>
<td>Glycine—proline, hydroxyproline</td>
<td>Membrane transport defect</td>
</tr>
<tr>
<td>Familial iminoglycinuria</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

6. Secondary aminoacidurias occur in the following:
   a. Viral hepatitis
   b. Multiple myeloma
   c. Hyperparathyroidism
   d. Rickets (vitamin D resistant)
   e. Osteomalacia
   f. Hereditary fructose intolerance
   g. Galactosemia
   h. Liver disease or necrosis
   i. Renal failure, renal disease
   j. Cystinosis
   k. Muscular dystrophy (progressive)

Interfering Factors

1. Amino acid concentration displays a marked circadian rhythm—30% variation, highest in midafternoon and lowest in morning.
2. Hyperalimentation and intravenous therapy affect outcome.
3. Drugs such as amphetamines, norepinephrine, levodopa, and all antibiotics affect results.
4. Age is a significant factor, especially in newborns and infants.
5. Pregnancy decreases values.

**Interventions**

**Pretest Patient Preparation**

1. Genetic counseling is recommended before specimen collection.
2. Instruct the patient regarding the test purpose, collection procedure, and need for refrigeration. A written reminder may be helpful.
3. Allow foods and moderate amounts of fluids (do not overhydrate).
4. It may be necessary to consume proteins or carbohydrates for a challenge load to produce certain amino acid metabolites.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel appropriately. Genetic counseling may be necessary.
2. Follow Chapter 1 guidelines regarding safe, effective, informed posttest care.

**BIBLIOGRAPHY**

Belsey R, Baer DM: Specimen collection for diagnosing UTI. Medical Laboratory Observer 28: 29, 1996
Cooper C: What color is that urine specimen. Am J Nurs 93(8): 37, 1990
Finnegan K: Correlating routine urinalysis with selected kidney disorders. Advances for Laboratory Professionals 2(12), June 1999
Thompson WG: Things that go red in the urine; and others that don't. Lancet 347: 5, 1996
OVERVIEW OF STOOL STUDIES

Stool examination is often done for evaluation of gastrointestinal (GI) disorders. These studies are helpful in detecting GI bleeding, GI obstruction, obstructive jaundice, parasitic disease, dysentery, ulcerative colitis, and increased fat excretion. (See Stool Analysis on p. 266.)

An adult excretes 100 to 200 g of fecal matter a day, of which as much as 75% may be water. The feces are what remain of the 8 to 10 L of digested fluid-like material that enters the intestinal tract each day, and oral food and fluids, saliva, gastric secretions, pancreatic juice, and bile add to the formation of feces.

Feces are composed of the following materials:

1. Waste residue of indigestible material (eg, cellulose) from food eaten during the previous 4 days
2. Bile (pigments and salts): stool color is normally due to bile pigments that have been altered by bacterial action.
3. Intestinal secretions
4. Water and electrolytes
5. Epithelial cells that have been shed
6. Large numbers of bacteria
7. Inorganic material (10%–20%), chiefly calcium and phosphates
8. Undigested or unabsorbed food (normally present in very small quantities)

The output of feces depends on a complex series of absorptive, secretory, and fermentative processes. Normal function of the colon involves three physiologic processes: (1) absorption of fluid and electrolytes; (2) contractions that churn and expose the contents to the GI tract mucosa and transport the contents to the rectum; and (3) defecation.

The small intestine is approximately 23 feet (7 m) long, and the large intestine is 4 to 5 feet (1.2–1.5 m) long. The small intestine degrades ingested fats, proteins, and carbohydrates to absorbable units and then absorbs them. Pancreatic, gastric, and biliary secretions exert their effects on the GI contents to prepare this material for active mucosal transport. Other active substances absorbed in the small intestine include fat-soluble vitamins, iron, and calcium. Vitamin B\textsubscript{12}, after combining with intrinsic factors, is absorbed in the ileum. The small intestine also absorbs as much as 9.5 L of water and electrolytes for return to the bloodstream. Small intestine contents (ie, chyme) begin to enter the rectum as soon as 2 to 3 hours after a meal, but the process is not complete until 6 to 9 hours after eating.

The large intestine performs less complex functions than the small intestine. The proximal or right colon absorbs most of the water remaining after the GI contents have passed through the small intestine. Colonic absorption of water, sodium, and chloride is a passive process. Fecal water excretion is only about 100 mL/day. The colon mainly moves the luminal contents to and fro by seemingly random contractions of circular smooth muscle. Increased propulsive activity (ie, peristalsis) occurs after eating. Peristaltic waves are caused by the gastrocolic and duodenocolic reflexes, which are initiated after meals and stimulated by the emptying of the stomach into the duodenum. The muscles of the colon are innervated by the autonomic nervous system. Additionally, the parasympathetic nervous system stimulates movement, and the sympathetic system inhibits movement. Massive peristalsis usually occurs several times a day. Resultant distention of the rectum initiates the urge to defecate. In persons with normal motility and a mixed dietary intake, normal
colon transit time is 24 to 48 hours.

**STOOL ANALYSIS**

Stool analysis determines the various properties of the stool for diagnostic purposes. Some of the more frequently ordered tests on feces include tests for leukocytes, blood, fat, ova, parasites, and pathogens (Table 4.1). Stool culture is explained in Chap. 7, Microbiologic Studies. Stool is also examined by chromatographic analysis for the presence of gallstones. The discovery of a gallstone from feces provides the only proof that a common bile duct stone has been dislodged and excreted. Stool testing also screens for colon cancer and asymptomatic ulcerations or other masses of the GI tract and evaluates GI diseases in the presence of diarrhea and/or constipation. Stool testing is done in immunocompromised persons for parasitic diseases. Fat analysis is used as the gold standard to diagnose malabsorption syndrome.

Table 4.1 Stool Testing for Infections

<table>
<thead>
<tr>
<th>Source of Stool Infection</th>
<th>Clinical Signs or Symptoms</th>
<th>Laboratory Test (Sequence or Follow-up)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Community Acquired from intermediate hosts, ie:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Home—pets, dogs, contaminated water</td>
<td>Diarrhea, bloody, purulent Steatorrhea</td>
<td>Screen stool for ova and parasites Microscopic exam of stool for ova and parasites</td>
</tr>
<tr>
<td>Occupational Fishing—snails and worms</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Meat cutters—from contaminated animals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Health care workers—from patients</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Farm workers—animals (cows, pigs), garden, flies, mosquitoes, insects, fleas, bugs</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Recreational—backpacking, poor sanitation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Travel—Third World—contaminated water supply</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Contact with fair animals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hookworm infection—not fecal or oral; it is direct penetration of skin by larva in contaminated soil or in animal droppings</td>
<td>Diarrhea, bloody, purulent Steatorrhea</td>
<td>Screen stool for ova and parasites Microscopic exam of stool for ova and parasites</td>
</tr>
</tbody>
</table>

| Nosocomial Acquired from institutions such as hospitals or nursing homes | | |
| | Diarrhea Medication history of antibiotic use | Eosinophilia in blood sample Exam for worms in stool or around anus Fecal smear for leukocytes and yeast |

| Personal Contact with an infected host when patient is compromised (weakened immune system, ie, HIV) or debilitated as in frail children or elderly | | |
| | | Screen stool for ova and parasites Microscopic exam of stool for ova and parasites |

Patients and health care personnel may dislike collecting and examining fecal material; however, this natural aversion must be overcome in light of the value of a stool examination for diagnosing disturbances and diseases of the GI tract, the liver, and the pancreas.

**Random Collection and Transport of Stool Specimens**

1. Observe standard/universal precautions (see Appendix A) when procuring and handling specimens to avoid infectious pathogens (eg, hepatitis A, *Salmonella*, and *Shigella*).
2. Collect feces in a dry, clean, urine-free container that has a properly fitting cover.
3. The specimen should be uncontaminated with urine or other bodily secretions such as menstrual blood. Stool can be collected from the diaper of an infant or incontinent adult. Samples can be collected from temporary ostomy bags.
4. While wearing gloves, collect the entire stool specimen and transfer it to a container using a clean tongue blade or similar object. A sample 2.5-cm (1-inch) long or 64.7 mg (1 oz) of liquid stool may be sufficient for some tests.
5. For best results, cover specimens and deliver to the laboratory immediately after collection. Depending on the examination to be performed, the specimen should be either refrigerated or kept warm. If you are unsure of how to handle the specimen, contact the laboratory for detailed instructions concerning the disposition of the fecal specimen before collection is begun.
6. Post signs in bathrooms that say “DO NOT DISCARD STOOL” or “SAVE STOOL” to serve as reminders that fecal
Collection and Transport of Specimens for Ova and Parasites

1. Wear gloves. Observe standard precautions (see Appendix A). Collect feces in a dry, clean, urine-free container. If unsure of how to collect specimen, contact the laboratory before collection is begun.
2. Warm stools are best for detection of ova and parasites. Do not refrigerate specimens for ova and parasites.
3. Special vials that contain 10% formalin and polyvinyl alcohol (PVA) fixative may be used for collecting stool samples to test for ova and parasites. In this case, specimen storage temperature is not critical.
4. Because of the cyclic life cycle of parasites, three separate random stool specimens for analysis are recommended.
5. Place the specimen in a biohazard bag.

Collection and Transport of Specimens for Enteric Pathogens

1. While wearing gloves, collect feces in a dry, clean, urine-free container. If unsure of how to collect the specimen, contact the laboratory before collection is begun. Observe standard precautions.
2. Some coliform bacilli produce antibiotic substances that destroy enteric pathogens. Refrigerate the specimen immediately to prevent this from happening in the sample.
3. A diarrheal stool will usually give accurate results.
4. A freshly passed stool is the specimen of choice.
5. Collect stool specimens before antibiotic therapy is initiated and as early in the course of the disease as possible.
6. If mucus or blood is present, it definitely should be included with the specimen because pathogens are more likely to be found in these substances. If only a small amount of stool is available, a walnut-sized specimen is usually adequate.
7. Accurately label all stool specimens with the patient's name, date, and tests ordered on the specimen. Keep the outside of the container free from contamination and immediately send the sealed container to the laboratory.
8. For best preservation and transport of pathogens, a Cary-Blair solution vial with indicator should be used.

Interfering Factors for All Types of Stool Collection

1. Stool specimens from patients receiving tetracyclines, anti-diarrheal medications, barium, bismuth, oil, iron, or magnesium may not yield accurate results.
2. Bismuth found in paper towels and toilet tissue interferes with accurate results.
3. Do not collect or retrieve stool from the toilet bowl or use a specimen that has been contaminated with urine, water, or toilet bowl cleaner. A clean, dry bedpan may be the best receptacle for defeication.
4. Inaccurate test results may result if the sample is not representative of the entire stool evacuation.
5. Lifestyle, personal habits, travel, home and work environments, and bathroom accessibility are some of the factors that may interfere with proper sample procurement.
6. Specimen not transported promptly. Trophozites in liquid stool disintegrate rapidly after defeication; therefore, the specimen needs to be examined 30 minutes from start of collection of specimen, not 30 minutes from end of collection. Semi-formed stool should be examined within 60 minutes after defeication. No trophozites are seen in formed stool.

Interventions

Pretest Patient Preparation

1. Explain the collection purpose, procedure, and interfering factors in language the patient understands. Because the specimen cannot be obtained on demand, it is important to provide detailed instructions before the test so that the specimen is collected when the opportunity presents itself. Provide written instructions if necessary.
2. Provide proper containers and other collection supplies. Instruct the patient to defeicate in a large-mouthed plastic container, bag, or clean bedpan. Provide for and respect the patient's privacy.
3. Instruct the patient not to urinate into the collecting container or bedpan.
4. Do not place toilet paper in the container or bedpan because it interferes with testing.
5. If the patient has diarrhea, a large plastic bag attached by adhesive tape to the toilet seat may be helpful in the collection process. After defeication, the bag can be placed into a gallon container.
6. Specimens for most tests can be produced by a warm saline enema or Fleet Phospho-Soda enema.
7. Tests for both ova and parasites and cultures for enteric pathogens may be ordered together. In this case, the specimen should be divided into two samples, with one portion refrigerated for culture testing and one portion kept at room temperature for ova and parasite testing. There are commercial collection kits that require the stool to be divided and placed into separate vials for better recovery of ova and parasites and enteric pathogens. (See Chap. 7, Microbiologic Studies.)
8. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Provide patient privacy and the opportunity to cleanse perineal area and hands. Assist as necessary.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

1. Any stool collected may harbor highly infective pathogens. Use extreme caution and proper handling techniques at all times.
2. Instruct patients in proper hand-washing techniques after each use of the bathroom.
STOOL STUDIES

Stool Consistency, Shape, Form, Amount, and Odor

Inspection of the feces is an important diagnostic tool. The quantity, form, consistency, and color of the stool should be noted. When diarrhea is present, the stool is watery. Large amounts of mushy, frothy, foul-smelling stool are characteristic of steatorrhea. Constipation is associated with firm, spherical masses of stool. Feces have a characteristic odor that varies with diet and the pH of the stool.

Normally, evacuated feces reflect the shape and caliber of the colonic lumen as well as the colonic motility. The normal consistency is somewhat plastic and neither fluid, mushy, nor hard. Consistency can also be described as formed, soft, mushy, frothy, or watery. The odor of normal stool is caused by indole and skatole, formed by bacterial fermentation and putrefaction.

Reference Values

Normal

1. 100–200 g/d
2. Characteristic odor present; plastic, soft, formed; soft and bulky on a high-fiber diet; small and dry on a high-protein diet; seeds and small amounts of vegetable fiber present (as opposed to muscle fiber) (Table 4.2)

<table>
<thead>
<tr>
<th>Table 4.2 Normal Values in Stool Analysis</th>
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<tr>
<td>Macroscopic Examination</td>
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<tr>
<td>Amount</td>
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<td>Color</td>
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<td>Odor</td>
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<td>Pus</td>
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<td>Parasites</td>
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| Microscopic Examination                   | Normal Values                                     |
| Fat                                       | Colorless, neutral fat (18%) and fatty acid crystals and soaps |
| Undigested food, meat fibers, starch, trypsin | None to small amount                              |
| Eggs and segments of parasites            | None                                              |
| Bacteria and viruses                      | None                                              |
| Yeasts                                    | None                                              |
| Leukocytes                                | None                                              |

| Chemical Examination                      | Normal Values                                     |
| Water                                     | Up to 75%                                         |
| pH                                        | Neutral to weakly alkaline (pH 7.0–7.5)            |
| Occult blood                              | Negative                                          |
| Urobilinogen                              | 50–300 mg/24 h                                    |
| Porphyrins                                | Coproporphyrins: 400–1200 µg/24 h (611–1832 nmol/d) |
|                                           | Uroporphyrins: 10–40 µg/24 h (12–48 nmol/d)       |
| Nitrogen                                  | <2.5 g/24 h (<178 mmol/d)                         |
| Apt test for swallowed blood              | Negative in adults; positive in newborns          |
| Trypsin                                   | 20–95 U/g                                         |
| Osmolality, used with stool               | 200–250 mOsm                                      |
| Na + K to calculate osmotic gap           | 5.8–9.8 mEq/24 h (5.8–9.8 mmol/d)                 |
| Sodium                                    | 2.5–3.9 mEq/24 h (2.5–3.9 mmol/d)                 |
| Chloride                                  | 15.7–20.7 mEq/24 h (15.7–20.7 mmol/d)             |
| Potassium                                 | 0–6 g/24 h (0–21 mmol/d)                          |
| Lipids (fatty acids)                      | <0.25 g/dL                                        |
| Carbohydrates (as reducing substances)    |                                                   |

Note: Reference values for electrolytes differ greatly from laboratory to laboratory.

Procedure Collect a random stool specimen in a plastic container.

Clinical Implications
1. Fecal consistency alterations
   a. Diarrhea due to the following:
      1. Infection—Salmonella, Shigella, Yersinia, human immunodeficiency virus (HIV) enteropathy, Campylobacter
      2. Inflammatory disorder—Crohn's disease, ulcerative colitis
      3. Steatorrhea—sprue, celiac disease
      4. Carbohydrate malabsorption—lactose or sucrose deficiency
      5. Endocrine abnormalities—diabetes mellitus, hyperthyroid or hypothyroid, adrenal insufficiency
      6. Hormone-producing tumors—Zollinger-Ellison syndrome, gastrinoma, medullary thyroid carcinoma, villous adenoma
      7. Colon carcinoma
      8. Infiltration of lesions due to lymphoma, scleroderma of bowel
      9. Drugs, antibiotics, cardiac medications, chemotherapy
      10. Osmotically active dietary items—sorbitol, psyllium fiber, caffeine, ethanol
      11. GI surgery—gastrectomy, stomach stapling, intestinal resection
      12. Factitious—self-induced laxative abuse associated with psychiatric disorders
   b. "Pasty" stool associated with high-fat content can be caused by the following:
      1. Common bile duct obstruction
      2. Celiac disease (sprue and steatorrhea); stool resembles aluminum paint
      3. Cystic fibrosis—greasy "butter" stool appearance due to pancreatic involvement
   c. Bulky or frothy stool is usually due to steatorrhea and celiac disease.

2. Alterations in stool size or shape indicate altered motility or colon wall abnormalities.
   a. A narrow, ribbon-like stool suggests the possibility of spastic bowel, rectal narrowing or stricture, decreased elasticity, or a partial obstruction.
   b. Excessively hard stools are usually due to increased fluid absorption because of prolonged contact of luminal contents with colon mucosa during delayed transit time through the colon.
   c. A large-circumference stool indicates dilatation of the viscus.
   d. Small, round, hard stools (ie, scybala) accompany habitual, moderate constipation.
   e. Severe fecal retention can produce huge, firm, impacted stool masses with a small amount of liquid stool as overflow. These must be removed manually, occasionally under light anesthesia.

3. Fecal odor should be assessed whenever a stool specimen is collected.
   a. A foul odor is caused by dehydration of undigested protein and is produced by excessive carbohydrate ingestion.
   b. A sickly sweet odor is produced by volatile fatty acids and undigested lactose.

4. Mucus in stool occurs in constipation, malignancy, and colitis (see p. 278).

Interventions

Pretest Patient Preparation

1. Ensure that the patient avoids barium procedures and laxative preparations for 1 week before stool specimen collection.
2. Advise the patient of the purpose of test and instruct him or her in collection techniques and refrigeration of specimen. Provide collection container.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Assessment of Diarrhea and Constipation

1. When performing a workup for the differential diagnosis of diarrhea or constipation, a patient history is most important. The following factors should be charted:
   a. An estimate of volume and frequency of fecal output
   b. Stool consistency and presence of blood, pus, mucus, oiliness, or bad odor in specimen; evaluate through direct observation
   c. Decrease or increase in frequency of defecation
   d. Sensations of rectal fullness with incomplete stool evacuation
   e. Painful defecation
2. Assess dietary habits and food allergies.
3. Assess emotional state of patient—psychological stress may be major cause of altered bowel habits.
4. Be alert for signs of laxative abuse.

Posttest Patient Aftercare

1. Evaluate outcome; interpret, report, and record findings. If abnormalities are detected, counsel patient appropriately. If patient has watery diarrhea, note history of contact with affected family members, travel to a developing country, involvement in vacation or resort backpacking, community and municipal water supply, or contact with farm animals. Explain that additional testing (eg, colonoscopy) may be necessary.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Stool Color

The brown color of normal feces is probably due to stercobilin (urobilin), a bile pigment derivative, which results from the action of reducing bacteria in bilirubin and other undetermined factors.

The first indication of GI disturbances is often a change in the normal brown color of the feces. A change in color can provide information about pathologic conditions, organic dysfunction, or intake of drugs. Color abnormalities may aid the
clinician in selection of appropriate diagnostic chemical and microbiologic stool tests.

Reference Values

**Normal** Brown

**Procedure** Collect a random stool specimen. Observe standard precautions.

**Clinical Implications** The color of feces changes in some disease states is as follows:

1. Yellow, yellow-green, or green: severe diarrhea
2. Black, with a tarry consistency: usually the result of bleeding in the upper GI tract (>100 mL blood)
3. Maroon, red, or pink: possibly the result of bleeding of the lower GI tract from tumors, hemorrhoids, fissures, or an inflammatory process
5. Pale, with a greasy consistency: pancreatic deficiency causing malabsorption of fat

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Grossly visible blood always indicates abnormal state.

1. Blood streaked on the outer surface of stool usually indicates hemorrhoids or anal abnormalities.
2. Blood present in stool can also be caused by abnormalities higher in the colon. If transit time is sufficiently rapid, blood from the stomach or duodenum can appear as bright red, dark red, or maroon in stool.

**Interfering Factors**

1. Stool darkens on standing.
2. The color of stool is influenced by diet (certain foods), food dyes, and drugs (see Appendix J).
   a. Yellow-rhubarb; yellow to yellow-green color occurs in the stool of breast-fed infants who lack normal intestinal flora.
   b. Pale yellow, white, or gray stools are due to barium intake.
   c. Green color occurs with diets high in chlorophyll-rich green vegetables such as spinach or with some drugs (see Appendix J).
   d. Black color may be due to foods such as cherries, an unusually high proportion of dietary meat, artificially colored foods such as black jelly beans, or drugs and supplements such as charcoal, bismuth, or iron.
   e. Light-colored stool with little odor may be due to diets high in milk and low in meat.
   f. Clay-like color may be due to a diet with excessive fat intake or barium intake.
   g. Red color may be due to a diet high in beets or tomatoes, red food coloring, or peridium compound.
   h. Certain color changes may result from specific drugs (see Appendix J).

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A complete dietary and drug history will help to differentiate significant abnormalities from interfering factors.

**Interventions**

**Pretest Patient Preparation**

1. Advise patient of purpose of test. Ask patient to notify clinician about stool color changes. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
2. Record dietary and drug history.
3. Ensure that the patient avoids laxatives and barium procedures for 1 week before collection.

**Posttest Patient Aftercare**

1. Interpret and document abnormal appearance and colors of stool; counsel patient appropriately regarding the meaning of color changes and explain need for further testing (eg, GI studies).
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Blood in Stool; Occult Blood**

The most frequently performed fecal analysis is chemical screening for the detection of occult (ie, hidden) blood. Bleeding in the upper GI tract may produce a black, tarry stool. Bleeding in the lower GI tract may result in an overtly bloody stool. However, no visible signs of bleeding may be present with smaller amounts of blood found in early stages of GI diseases; thus, the chemical detection of occult blood is necessary to identify and treat disease early in its course. Occult blood testing is also controversial owing to many false-positive and false-negative results. If the patient preparation and collection of specimen is followed explicitly, the results are more accurate.

An average, healthy person passes up to 2.0 mL of blood per 150 g of stool into the GI tract daily. Passage of more than 2.0 mL of blood in the stool in 24 hours is pathologically significant. Detection of occult blood in the stool is very useful in detecting early disease of the GI tract. This test demonstrates the presence of blood produced by upper GI bleeding, as in the presence of gastric ulcer; it also screens for colonic carcinomas while they are still in the localized stages. With proper medical follow-up, an 84% survival rate has been demonstrated for treatment of colonic carcinoma.

Reference Values

**Normal** Negative for blood

**Procedure**
1. Obtain a random stool specimen. Observe standard precautions. Tests for detecting fecal blood use the pseudoperoxidase activity of hemoglobin reacting with hydrogen peroxide to oxidize a colorless compound to a colored one (usually blue). Hemoccult II (Smith-Kline) is the most widely used commercial test with the lowest percentage of false-positive results (1%–12%). This test system uses guaiac-impregnated filter paper as the chromogen that produces the blue color in a positive reaction.

2. Apply a thin smear of stool inside the indicated circle using a wood applicator stick and allow it to dry. If stool is bloody, the collector may be at risk for hepatitis B, hepatitis C, or HIV infection.

3. Protect the Hemoccult slide from light, heat, and humidity. Do not refrigerate.

4. Do not allow the delay between smearing the stool and testing to exceed 14 days. Do not refrigerate sample before testing.

Clinical Implications

1. Stool that appears dark red to tarry black indicates a loss of 50.0 to 75.0 mL of blood from the upper GI tract. Smaller quantities of blood in the GI tract can produce similar-appearing stools or appear as bright red blood.

2. A stool sample should be considered grossly bloody only after a chemical testing for presence of blood. This will eliminate the possibility that abnormal coloring caused by diet or drugs may be mistaken for bleeding in the GI tract.

3. Positive testing for occult blood may be caused by the following conditions:
   a. Carcinoma of colon
   b. Ulcerative colitis and other inflammatory lesions
   c. Adenoma
   d. Diaphragmatic hernia
   e. Gastric carcinoma
   f. Rectal carcinoma
   g. Peptic ulcer
   h. Gastritis
   i. Vasculitis
   j. Amyloidosis
   k. Kaposi's sarcoma

Clinical Alert

1. To be accurate, the test employed must be repeated three to six times on different stool samples; some bowel lesions may bleed intermittently.

2. The patient's diet should be free of meat and vegetable sources of peroxidase activity (eg, turnips, horseradish, red or rare meat, cauliflower, broccoli, cantaloupe, parsnips). Only after following this regimen can a positive series of tests be considered an indication for further patient evaluation and testing.

Interfering Factors

1. Drugs such as salicylates (aspirin), steroids, indomethacin, nonsteroidal anti-inflammatory drugs (NSAIDs), anticoagulants, colchicine, and antimetabolites are associated with increased GI blood loss in average, healthy persons and with more pronounced bleeding when disease is present. GI bleeding can also follow parenteral administration of the above-mentioned drugs and should be avoided 7 days before testing.

2. Drugs that may cause false-positive results for occult blood testing include the following:
   a. Boric acid
   b. Bromides
   c. Colchicine
   d. Iodine, povidone-iodine (Betadine)
   e. See Appendix J for other drugs.

3. Foods that may cause false-positive results for occult blood testing include the following:
   a. Meats, including processed meats and liver, which in the diet contain hemoglobin, myoglobin, and certain enzymes that can give false-positive test results for up to 4 days after consumption.
   b. Vegetables and fruits with peroxidase activity (eg, turnips, horseradish, mushrooms, broccoli, apples, radishes, bananas, cantaloupe)

4. Substances that cause false-negative results for occult blood testing include the following:
   a. Ascorbic acid (vitamin C) in excess of 250 mg/day
   b. Vitamin C–enriched foods and juices
   c. Iron supplements that contain vitamin C in excess of 250 mg
   d. See Appendix J for other drugs.

5. Other factors affecting test results include the following:
   a. Bleeding hemorrhoids may produce erroneous results; take samples from center of stool to avoid this error.
   b. Collection of specimen during menstrual period
   c. Hematuria (ie, blood in urine)
   d. Some long-distance runners (23%) have positive outcomes for occult blood.
   e. Toilet bowl cleansers may interfere with the chemical reaction of the test; remove bowl cleaners and flush twice before proceeding with test.

Interventions

Pretest Patient Preparation

1. Explain the purpose, procedure, and interfering factors of the test as well as the need to follow appropriate stool collection protocols for using special kit for fecal occult blood or a plastic container with a lid.

2. Recommend that the patient consume a high-residue diet, starting 72 hours before and continuing throughout the collection period. Roughage in diet can increase test accuracy by helping to uncover silent lesions that bleed
intermittently. The diet may include the following:

a. Meats: only small amounts of chicken, turkey, and tuna
b. Vegetables: generous amounts of both raw and cooked vegetables, including lettuce, corn, spinach, carrots, and celery; avoid vegetables with high peroxidase activity (see \textsuperscript{3b} above)
c. Fruits: plenty of fruits, especially prunes
d. Cereals: bran and bran-containing cereals
e. Moderate amounts of peanuts and popcorn daily. If any of the above foods are known to cause discomfort, the patient should consult the physician.

3. Ensure that the patient receives no barium enemas 72 hours before or during testing.

\textbf{Education Alert}

- Do not collect samples during or until 3 days after your menstrual period, or while you have bleeding hemorrhoids or blood in your urine.
- Do not consume the following medications, vitamins, and foods: for 7 days before and during the test period, avoid aspirin or other NSAIDs; for 72 hours before and during the test period, avoid vitamin C in excess of 250 mg/d (from all sources, dietary and supplementary), red meat (eg, beef, lamb), including processed meats and liver, and raw fruits and vegetables (especially melons, radishes, turnips, and horseradish).
- Remove toilet bowl cleaners from toilet tank and flush twice before proceeding to defecate.
- Collect samples from three consecutive bowel movements or three bowel movements closely spaced in time and spread a small stool sample (minimum, 1 mL) on each of the three slides or card provided.
- Protect card or slides from heat, light, and volatile chemicals (eg, iodine, bleach). Keep cover flap of slides closed when not in use.

\textbf{Posttest Patient Aftercare}

1. Patient may resume normal diet after testing is complete.
2. Interpret occult blood test results and record findings. Counsel the patient regarding abnormal findings and monitor as necessary. Advise that further testing (eg, barium enema, defecography) and follow-up may be required.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

\textbf{Clinical Alert}

Blood in the stool is abnormal and should be reported and recorded.

\textbf{Apt Test for Swallowed Blood}

Dr. L. Apt developed the test for identifying the swallowed blood syndrome. The swallowed blood syndrome refers to bloody stools usually passed on the second or third day of life. The blood may be swallowed during delivery or may be from a fissure of the mother's nipple in breast-fed infants. This condition must be differentiated from GI hemorrhage of the newborn. The test is based on the fact that the infant's blood contains largely fetal hemoglobin (Hb F), which is alkali resistant. This blood can be differentiated from the mother's blood using laboratory methods.

The Apt test is used to differentiate swallowed blood syndrome from infant GI hemorrhage. The test can be done on feces or vomitus. In the laboratory, the blood is dissolved and treated with NaOH for alkali denaturation. Fetal hemoglobin is alkali resistant, and the solution of blood remains pink. Swallowed blood of maternal origin contains adult hemoglobin, which is converted to brownish hematin when the alkali is added.

\textbf{Reference Values}

\textit{Normal} Test result will indicate whether blood present in newborn feces or vomitus is of maternal or fetal origin.

\textbf{Procedure}

1. Collect a random stool specimen from a newborn infant; observe standard precautions.
2. The following are acceptable specimens:
   a. Blood-stained diaper
   b. Grossly bloody stool
   c. Bloody vomitus or gastric aspiration
3. Place specimen or specimens in a biohazard bag and deliver to the laboratory as soon as possible. Refrigerate the specimen or specimens if there is any delay.

\textbf{Clinical Implications}

1. Fetal hemoglobin, which is pink in color, is present in gastric hemorrhage of the newborn.
2. Adult hemoglobin, which is brownish in color, is present in swallowed blood syndrome in the infant.

\textbf{Interfering Factors}

1. The test is invalid with black, tarry stools because the blood has already been converted to hematin.
2. The test is invalid if there is insufficient blood present; grossly visible blood must be present in the specimen.
3. Vomitus with pH < 3.9 produces an invalid test result.
4. The presence of maternal thalassemia major produces a false-positive test result because of increased maternal hemoglobin F.

\textbf{Interventions}
**Pretest Patient Preparation**

1. Advise parent or parents of the purpose of the test.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Review test results and counsel the parent or parents regarding test outcome, further testing, and possible treatment for infant GI hemorrhage.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Mucus in Stool**

The mucosa of the colon secretes mucus in response to parasympathetic stimulation. Recognizable mucus in a stool specimen is abnormal and should be reported and recorded.

**Reference Values**

**Normal** Negative for mucus

**Procedure** Collect a random stool specimen. Observe and report findings of mucus.

**Clinical Implications**

1. Translucent gelatinous mucus clinging to the surface of formed stool occurs in the following conditions:
   a. Spastic constipation
   b. Mucous colitis
   c. Emotionally disturbed patients
   d. Excessive straining at stool
2. Bloody mucus clinging to the feces suggests the following conditions:
   a. Neoplasm
   b. Inflammation of the rectal canal
3. In villous adenoma of the colon, copious quantities of mucus may be passed (up to 3–4 L in 24 hours).
4. Mucus and diarrhea with white and red blood cells is associated with the following conditions:
   a. Ulcerative colitis
   b. Bacillary dysentery (<i>Salmonella</i>)
   c. Ucerating cancer of colon
   d. Acute diverticulitis
   e. Intestinal tuberculosis
   f. Regional enteritis
   g. Amebiasis

**Interventions**

**Pretest Patient Preparation**

1. Advise patient of purpose of observing for stool mucus.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
3. Ensure that the patient avoids laxatives and barium procedures for 1 week before test.

**Posttest Patient Aftercare**

1. Report and record presence, type, and amount of mucus.
2. Counsel patient appropriately. Monitor bowel habits. Explain that further testing and follow-up monitoring may be necessary.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Stool pH**

Stool pH is diet dependent and is based on bacterial fermentation in the small intestine. Carbohydrate fermentation changes the pH to acid; protein breakdown changes the pH to alkaline.

Stool pH testing is done to evaluate carbohydrate and fat malabsorption and assess disaccharidase deficiency. Breast-fed infants have slightly acid stool; bottle-fed infants have slightly alkaline stools.

**Reference Values**

**Normal**

1. Neutral to slightly acid or alkaline: pH 7.0–7.5 depending on diet
2. Newborns: pH 5.0–7.5

**Procedure**

1. Collect a fresh, random stool specimen in a plastic container with a tight-fitting lid (see p. 266).
2. Refrigerate specimen.

**Clinical Implications**
1. Increased pH (alkaline)
   a. Secretory diarrhea without food intake
   b. Colitis
   c. Villous adenoma
   d. Antibiotic use (impairment of colonic fermentation)
2. Decreased pH (acid)
   a. Carbohydrate malabsorption
   b. Fat malabsorption
   c. Disaccharidase deficiency (intestinal)

Interfering Factors
1. Barium procedures and laxatives affect test outcomes. They should be avoided for 1 week before stool sample collection.
2. Specimens contaminated with urine will invalidate the test.

Interventions
Pretest Patient Preparation
1. Explain the purpose and procedure of the test, following general guidelines in Chapter 1 for safe, effective, informed pretest care.
2. Advise patient that laxatives and barium procedures should be avoided for 1 week before stool sampling.

Posttest Patient Aftercare
1. Interpret pH outcome and record findings. If abnormal pH is found, assess dietary patterns and antibiotic use.
2. Monitor as appropriate for malabsorption syndrome.
3. Order a stool reducing substance test if disaccharidase deficiency is suspected (see Stool Reducing Substances Test below).
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Stool Reducing Substances Test
Normally, sugars are rapidly absorbed in the upper small intestine. However, if this is not the case, they remain in the intestine and cause osmotic diarrhea due to osmotic pressure of the unabsorbed sugar in the intestine, drawing fluid and electrolytes into the gut. The unabsorbed sugars are measured as reducing substances. Reducing substances that can be detected in the stool include glucose, fructose, lactose, galactose, and pentose. Carbohydrate malabsorption is a major cause of watery diarrhea and electrolyte imbalance seen in patients with the short bowel syndrome. Idiopathic lactase deficiency is common, occurring in 70% to 75% of Southern European Greeks and Italians, 70% of Black adults, >90% of Asian adults, and 5% to 20% of Caucasian American adults.

The finding of elevated levels of reducing substances in the stool is abnormal and suggests carbohydrate malabsorption. A presumptive diagnosis of disaccharide intolerance can be made with an elevated reducing substance level along with an acid (ie, low) pH.

Reference Values
Normal
1. Normal: <0.25 g/dL (or <13.9 mmol/L) reducing substances in stool
2. Questionable: 0.25–0.50 g/dL (or 13.9–27.8 mmol/L) reducing substances in stool

Abnormal >0.5 g/dL (or >27.8 mmol/L) reducing substances in stool

Procedure Collect a fresh, random stool specimen and immediately deliver it to the laboratory (see p. 266).

Clinical Implications Elevated reducing substances in stool are found in the following conditions:
1. Disaccharidase deficiency (intestinal)
2. Short bowel syndrome
3. Idiopathic lactase deficiency, primary alactasia (enzyme deficiency leading to lactose intolerance)
4. Carbohydrate malabsorption abnormalities due to:
   a. Sprue
   b. Celiac disease
   c. Viral gastroenteritis

Interfering Factors
1. Bacterial fermentation of sugars may give falsely low results if the stool is not tested immediately.
2. Newborns may normally have elevated results.
3. Drug may cause malabsorption (eg, neomycin, kanamycin, methotrexate).

Interventions
Pretest Patient Preparation
1. Explain the purpose of the test and interfering factors.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.
2. If outcome is positive, further testing (lactose intolerance) and/or treatment (dietary therapy) may be necessary.

Leukocytes in Stool

Microscopic examination of the feces for the presence of white blood cells (leukocytes) is performed as a preliminary procedure in determining the cause of diarrhea. Leukocytes are normally not present in stools and are a response to infection or inflammation.

The presence or absence of fecal leukocytes can provide diagnostic information before the isolation of a bacterial pathogen. Neutrophils (>3 neutrophils per high-power field) are seen in the feces in conditions that affect the intestinal wall (e.g., ulcerative colitis, invasive bacterial pathogen infection). Viruses and parasites usually do not cause neutrophils in the stool. The greater the number of leukocytes, the greater the likelihood that an invasive pathogen is present.

Reference Values

Normal Negative for leukocytes

Procedure Collect a random stool specimen (see p. 266). Mucus or a liquid stool specimen can be used. A fresh specimen is preferred, or it may be preserved in PVA.

Clinical Implications

1. Large amounts of leukocytes (primarily neutrophils) accompany the following conditions:
   a. Chronic ulcerative colitis
   b. Bacillary dysentery
   c. Localized abscesses
   d. Fistulas of the sigmoid rectum or anus
   e. Shigellosis
   f. Salmonellosis
   g. Yersinia infection
   h. Invasive Escherichia coli diarrhea
      i. Campylobacter

2. Primarily mononuclear leukocytes appear in typhoid. Few leukocytes are sometimes seen in amebiasis.

3. Absence of leukocytes is associated with the following conditions:
   a. Cholera
   b. Nonspecific diarrhea (e.g., drug or food induced)
   c. Viral diarrhea
   d. Amebic colitis (many red blood cells)
   e. Noninvasive E. coli diarrhea
   f. Toxigenic bacteria (e.g., Staphylococcus, Clostridium)
   g. Parasites (e.g., Giardia, Entamoeba)

Interfering Factors Fecal leukocytes cannot be performed on formalin-preserved specimens.

Interventions

Pretest Patient Preparation

1. Explain the purpose of the test and the collection procedure. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
2. Ensure that the patient avoids barium procedures and laxatives for 1 week before test.
3. Withhold antibiotic therapy until after collection.

Posttest Patient Aftercare

1. Interpret abnormal test results. Monitor for diarrhea. Counsel patient concerning the need for follow-up tests (stool culture) and treatment (drugs, e.g., antibiotics).
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Collection and Transport of 24-, 48-, 72-, and 96-Hour Stool Specimens

This method is used to test for fat, porphyrins, urobilinogen, nitrogen, and electrolytes.

Special Instructions for Submitting Individual Specimens

1. Collect all stool specimens for 1 to 3 days. The entire stool should be collected. Some procedures may require 4 days.
2. Label specimens with day of test (e.g., Day 1, Day 2, Day 3, Day 4), time of day collected, patient's name, and tests ordered. It is important for calculations to disclose the number of days collected.
3. Submit individual specimens to the laboratory as soon as they are collected.

Special Instructions for Submitting Total Specimens

1. Obtain a 1-gallon container from the laboratory (a 1-gallon paint tin or covered plastic pail is preferred).
2. Save all stool and place in the container. Keep refrigerated or in a container with canned ice and replace ice as needed.
3. Transfer the properly labeled container to the laboratory at the end of the collection period.
4. Record dates, duration of collection time period, tests to be performed, patient's name, and other vital information on the collection receptacle.

**Fat in Stool; Fecal Fat Stain**

Fecal fat is the gold standard test for diagnosing steatorrhea (malabsorption). The three major causes of steatorrhea, which is a pathologic increase in fecal fat, are impairment of intestinal absorption, deficiency of pancreatic digestive enzymes, and deficiency of bile.

Specimens from patients suspected of having steatorrhea can be screened microscopically for the presence of excess fecal fat. This procedure can also be used to monitor patients undergoing treatment for malabsorption disorders. In general, there is good correlation between the qualitative and quantitative fecal fat procedures. Lipids included in the microscopic examination of feces are neutral fats (triglycerides), fatty acid salts (soaps), fatty acids, and cholesterol. The presence of these lipids can be observed microscopically by staining with Sudan III, Sudan IV, or oil red O dye. The staining procedure consists of two parts: the neutral fat stain and the split fat stain for fatty acids.

**Reference Values**

**Normal**

1. Qualitative
   a. Neutral fat: <50 fat globules per high-power field
   b. Fatty acids: <100 fat globules per high-power field

2. Quantitative
   a. Adult: 2–7 g/24 h (or 2–7 g/d) and <20% of total solids
   b. Infant: <1.0 g/24 h (or <1.0 g/d) and breast fed 10%–40% of total solids; bottle fed 30%–50% of total solids

**Procedure**

1. Collect a 48- to 96-hour specimen for the quantitative test. A random specimen can be used for the qualitative test. Each individual stool specimen is collected and identified with the name of the patient, time and date of collection, and test to be performed. Also indicate the length (actual time frame) of the collection period. The specimen should be sent immediately to the laboratory.
2. Follow the procedure for the collection of 24-, 48-, or 72-hour specimens.

**Clinical Implications**

1. Increases in fecal fat and fatty acids are associated with malabsorption syndrome caused by the following conditions:
   a. Celiac disease
   b. Crohn's disease
   c. Whipple's disease
   d. Cystic fibrosis
   e. Regional enteritis
   f. Sprue
   g. Atrophy of malnutrition
2. Increases in fecal fat and fatty acids are also found in the following conditions:
   a. Enteritis and pancreatic diseases in which there is a lack of lipase (eg, chronic pancreatitis)
   b. Surgical removal of a section of the intestine
3. Fecal fat test does not provide a diagnostic explanation for the presence of steatorrhea. It is not useful for differentiating among pancreatic diseases.
   a. D-Xylose absorption test may be ordered for the differential diagnosis of malabsorption.

**Interfering Factors**

1. Increased neutral fat may occur under the following nondisease conditions:
   a. Use of rectal suppositories and/or oily creams applied to the perineum
   b. Ingestion of castor oil, mineral oil
   c. Ingestion of dietetic low-calorie mayonnaise, oily salad dressings
   d. Ingestion of high-fiber diet (>100 g/24 h or >100 g/d)
   e. Use of psyllium-based stool softeners (eg, Metamucil)
2. Use of barium and bismuth interfere with test results.
3. Urine contaminates the specimen.
4. A random stool specimen is not an acceptable sample for the quantitative fat test.

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose of the test, interfering factors, and the procedure for the collection of specimens. Follow guidelines in Chapter 1 concerning diverse patient needs and safe, effective, informed pretest care.
2. For a 72- to 96-hour stool collection, ensure the patient has a diet containing 100–150 g of fat, 100 g of protein, and 180 g of carbohydrate for 6 days before and during the test.
3. Do not allow patient to have laxatives for 3 days before the test.
4. Follow the procedure for the collection of 72-hour stool specimens.

**Posttest Patient Aftercare**

1. Resume normal diet.
2. Record appearance, color, and odor of all stools in persons suspected of having steatorrhea. The typical stool in patients with this condition is foamy, greasy, soft, pasty, and foul smelling.
3. Counsel patient concerning test outcome and possible need for further testing (eg, colonoscopy) and treatment (eg, elimination of certain foods from the diet).
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Meat Fibers in Stool; Stool Muscle Fiber**

The presence of undigested meat fibers (ie, muscle fibers) in stool implies impaired intraluminal digestion. There is positive correlation between the presence of meat or muscle fibers and the presence of fat excreted in the stool.

**Reference Values**

*Normal* Negative (no undigested meat fibers present in the normal stool)

**Procedure**

1. Ensure that the patient eats 4 to 6 ounces of red meat for 24 to 72 hours before testing.
2. Collect a random specimen (see p. 266). Specimens obtained with warm saline enema or Fleet Phospho-Soda are acceptable.
3. Record method and type of stool procurement.

**Clinical Implications** Increased amounts of meat fibers are found in the following conditions:

1. Malabsorption syndromes caused by biliary obstruction
2. Pancreatic exocrine dysfunction (cystic fibrosis)
3. Gastrocolic fistula

**Interfering Factors**

1. Specimens should not be obtained with mineral oil, bismuth, or magnesium compounds.
2. Barium procedures and laxatives should be avoided for 1 week before collection.

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose of the test and interfering factors.
2. Ensure that the patient eats a high-meat diet for 72 hours before test.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Resume normal diet.
2. Interpret test outcomes.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Urobilinogen in Stool**

Increased destruction of red blood cells, as in hemolytic anemia, increases the amount of urobilinogen excreted. Liver disease, in general, reduces the flow of bilirubin to the intestine and thereby decreases the fecal excretion of urobilinogen. In addition, complete obstruction of the bile duct reduces urobilinogen to very low levels.

This test investigates hemolytic diseases and hepatic obstructive conditions. Determination of stool urobilinogen is an estimation of the total excretion of bile pigments, which are the breakdown products of hemoglobin.

**Reference Values**

*Normal*

1. 50–300 mg/24 h or 100–400 Ehrlich units/100 g
2. Newborns–6 months: negative

**Procedure**

1. Collect a 48-hour specimen.
2. Protect the specimen from light. Send to the laboratory as soon as possible.

**Clinical Implications**

1. *Increased values* are associated with hemolytic anemias.
2. *Decreased values* are associated with the following conditions:
a. Complete biliary obstruction (clay-colored feces result)
b. Severe liver disease (eg, infectious hepatitis)
c. Oral antibiotic therapy that alters intestinal bacterial flora
d. Aplastic anemia, which results in decreased hemoglobin turnover

Interfering Factors See Appendix J for drugs that affect test outcomes.

Interventions

Pretest Patient Preparation

1. Explain purpose of test.
2. Ensure that the patient does not receive oral antibiotic therapy for 1 week before test.
3. Ensure that the patient avoids laxatives and barium procedure 1 week before test.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes. Counsel patient appropriately regarding further testing. Monitor patient for liver disease, biliary obstruction, and diarrhea.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Trypsin in Stool: Fecal Chymotrypsin

Trypsin is a proteolytic enzyme formed in the small intestine. In older children and adults, trypsin is destroyed by bacteria in the GI tract. Inadequate trypsin secretion can lead to malabsorption and abdominal discomfort. Chymotrypsin, an intestinal proteolytic enzyme secreted by the pancreas, can be used to assess pancreatic function. Fecal chymotrypsin is a more reliable measurement of pancreatic function than trypsin.

Procedural Alert

This test will probably be replaced by immunoassays. It is an unreliable test in older children and adults.

Reference Values

Normal Trypsin, 20–950 U/g or 20–950 µg/g stool Chymotrypsin, 74–1200 µg/g or 74–1200 mg/kg stool

Procedure

1. Collect random specimens and send to the laboratory. Three separate, fresh stools are usually collected.
2. Ensure that the specimen is taken to the laboratory and tested within 2 hours.
3. Give a cathartic before obtaining a specimen from older children (saline or Fleet only).

Clinical Implications Decreased amounts of trypsin occur in the following conditions:

1. Pancreatic deficiency syndromes (0–33 U/g or 0–33 µg/g stool)
2. Cystic fibrosis (sweat chloride test is diagnostic) (<20 U/g or <20 µg/g stool)

Interfering Factors

1. No trypsin activity is detectable in constipated stools owing to prolonged exposure to intestinal bacteria, which inactivates trypsin.
2. Barium and laxatives used less than 1 week before test affect results.
3. In adults, the test is unreliable owing to trypsin inactivation by intestinal flora.
4. Bacterial proteases may produce positive reactions when no trypsin is present.

Interventions

Pretest Patient Preparation

1. Explain purpose of test and interfering factors.
2. Ensure that the patient avoids barium procedures and laxatives for 1 week before stool collection.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret abnormal test results and counsel patient concerning possible need for follow-up testing (eg, sweat testing) and treatment (enzymes).
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

1. Diagnosis of pancreatic insufficiency should not be made until three specimens exhibit no trypsin activity.
2. Bacterial protease may produce positive reactions when no trypsin is present; therefore, both positive and negative reactions should be carefully interpreted.

Stool Electrolytes: Sodium, Chloride, Potassium, and Osmolality
Normal colon function involves absorption of fluid and electrolytes. Stool electrolyte tests are used to assess electrolyte imbalance in patients with diarrhea. Stool electrolytes must be evaluated along with the serum and urine electrolytes as well as clinical findings in the patient. Stool osmolality is used in conjunction with blood serum osmolality to calculate the osmotic gap and to diagnose intestinal disaccharide deficiency.

**Reference Values**

**Normal**

1. Sodium: 5.8–9.8 mEq/L or 5.8–9.8 mmol/d
2. Chloride: 2.5–3.9 mEq/L or 2.5–3.9 mmol/d
3. Potassium: 15.7–20.7 mEq/L or 15.7–20.7 mmol/d
4. Osmolality: 275–295 mOsm/kg
5. Osmotic gap: measured osmolality ($2[Na + K]$)

Reference values vary from laboratory to laboratory. Check with your laboratory for normal values.

**Procedure**

1. Collect a random or 24-hour liquid stool specimen.
2. Keep the specimen covered and refrigerated.

**Clinical Implications**

1. Electrolyte abnormalities occur in the following conditions:
   a. Idiopathic proctocolitis: increased sodium (Na) and chloride (Cl); normal potassium (K)
   b. Ileostomy: increased sodium (Na) and chloride (Cl), low potassium (K)
   c. Cholera: increased sodium (Na) and chloride (Cl)
2. Chloride is greatly increased in stool in the following conditions:
   a. Congenital chloride diarrhea
   b. Acquired chloride diarrhea or secondary chloride diarrhea
   c. Idiopathic proctocolitis
   d. Cholera
3. Stool osmolality 500 mg/dL per day is suspicious for factitious disorders (eg, laxative abuse, ingestion of rat poison). Higher levels indicate high amounts of stool reducing substances. The osmotic gap is increased in osmotic diarrhea caused by the following conditions:
   a. Saline laxatives
   b. Sodium or magnesium citrate
   c. Carbohydrates (lactulose or sorbitol candy)

**Interfering Factors**

1. Formed stools invalidate the results. Stools must be liquid for electrolyte tests.
2. The stool cannot be contaminated with urine.
3. Surreptitious addition of water to the stool specimen considerably lowers the osmolality. Stool osmolality must be less than 240 mOsm/kg (or <240 mmol/kg H$_2$O) to calculate the osmotic gap.
4. See Appendix J for drugs that cause increased values.

**Interventions**

**Pretest Patient Preparation**

1. Explain purpose of test, procedure for stool collection, and interfering factors.
2. Ensure the patient avoids barium procedures and laxatives for 1 week before collection of specimen.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret abnormal test outcomes. Monitor diarrhea episodes and record findings. Assess patient for electrolyte imbalances and counsel regarding further testing and treatment.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**BIBLIOGRAPHY**

Bakeman S: ABCs of Interpretive Laboratory Data, 3rd ed. Greenville, NC, Interpretive Laboratory Data Inc., 1993


OVERVIEW OF CEREBROSPINAL FLUID (CSF)
Description, Formation, and Composition of CSF

Cerebrospinal fluid (CSF) is a clear, colorless fluid formed within the cavities (ie, ventricles) of the brain. The choroid plexus produces about 70% of the CSF by ultrafiltration and secretion. The ependymal lining of the ventricles and cerebral subarachnoid space produce the remainder of the CSF total volume. Approximately 500 mL of CSF fluid is formed per day, although only 90 to 150 mL is present in the system at any one time. Reabsorption of CSF occurs at the arachnoid villi.

CSF circulates slowly from the ventricular system into the space surrounding the brain and spinal cord and serves as a hydraulic shock absorber, diffusing external forces to the skull that might otherwise cause severe injury. The CSF also helps to regulate intracranial pressure (ICP), supply nutrients to the nervous tissues, and remove waste products. The chemical composition of CSF does not resemble an ultrafiltrate of plasma. Certain chemicals in the CSF are regulated by specific transport systems (eg, K⁺, Ca²⁺, Mg²⁺), whereas other substances (eg, glucose, urea, creatinine) diffuse freely. Proteins enter the CSF by passive diffusion at a rate dependent on the plasma-to-CSF concentration gradient. The term blood-brain barrier is used to represent the control and filtration of blood plasma components (eg, restriction of protein diffusion from blood into brain tissue) to the CSF and then to the brain capillaries. The ratio of increased albumin in CSF to blood serum is always caused by blood-brain barrier dysfunction because albumin is found extensively in blood. A decreased CSF flow rate is due to decreased production or restriction or blockage of flow.

Most CSF constituents are present in the same or lower concentrations as in the blood plasma, except for chloride concentrations, which are usually higher. Disease, however, can cause elements ordinarily restrained by the blood-brain barrier to enter the spinal fluid. Erythrocytes and leukocytes can enter the CSF from the rupture of blood vessels or from meningeal reaction to irritation. Bilirubin can be found in the spinal fluid after intracranial hemorrhage. In such cases, the arachnoid granulations and the nerve root sheaths will reabsorb the bloody fluid. Normal CSF pressure will consequently be maintained by the reabsorption of CSF in amounts equal to its production. Blockage causes an increase in the amount of CSF, resulting in hydrocephalus in infants or increased ICP in adults. Of the many factors that regulate the level of CSF pressure, venous pressure is the most important because the reabsorbed fluid ultimately drains into the venous system.

Despite the continuous production (~0.3 mL/min) and reabsorption of CSF and the exchange of substances between the CSF and the blood plasma, considerable pooling occurs in the lumbar sac. The lumbar sac, located at L4 to L5, is the usual site used for puncture to obtain CSF specimens because damage to the nervous system is less likely to occur in this area. In infants, the spinal cord is situated more caudally than in adults (L3–L4 until 9 months of age, when the cord ascends to L1–L2); therefore; a low lumbar puncture should be made in these patients (Table 5.1).

Table 5.1 Normal CSF Values

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volume</td>
<td>Adult: 90–150 mL; child: 60–100 mL</td>
</tr>
<tr>
<td>Appearance</td>
<td>Crystal clear, colorless</td>
</tr>
<tr>
<td>Pressure</td>
<td>Adults: 90–180 mm H₂O; child: 10–100 mm H₂O</td>
</tr>
<tr>
<td>Total cell count</td>
<td>Essentially free cells</td>
</tr>
</tbody>
</table>
**WBCs**

- Adults: 0–5 cells
- Newborn (0–14 d): 0–30 cells

**Differential**

- Lymphocytes: 40%–80%
- Monocytes: 15%–45%
- Polys: 0%–6%

**RBCs (has limited diagnostic value)**

Specific gravity: 1.006–1.008

### Clinical Tests

**Glucose**

- Adults: 40–70 mg/dL (2.2–3.9 mmol/L)
- Newborn (0–14 d): 60–80 mg/dL

**Protein**

- Lumbar: Adults: 15–45 mg/dL (150–450 mg/L)
- Neonates: 15–100 mg/dL (150–1000 mg/L)
- Elderly (>60 y): 15–60 mg/dL
- Cisternal: 15–25 mg/dL (150–250 mg/L)
- Ventricular: 5–15 mg/dL (50–150 mg/L)
- Lactic acid (lactate): 10–24 mg/dL (1.11–2.66 mmol/L)
- Glutamine: 5–20 mg/dL (0.34–1.37 mmol/L)
- Albumin: 10–35 mg/dL (1.52–5.32 µmol/L)
- Urea nitrogen: 6–16 mg/dL (2.14–5.71 mmol/L)
- Creatinine: 0.5–1.2 mg/dL (44–106 µmol/L)
- Uric acid: 0.5–4.5 mg/dL (29.7–268 µmol/L)
- Bilirubin: 0 (none)
- Phosphorous: 1.2–2.0 mg/dL (387–268 µmol/L)
- Ammonia: 10–35 µg/dL (5.87–20.5 µmol/L)

**Electrolytes and pH**

- **pH**
  - Lumbar: 7.28–7.32
  - Cisternal: 7.32–7.34

- **Chloride**
  - 115–130 mEq/L (mmol/L)
- **Sodium**
  - 135–160 mEq/L (mmol/L)
- **Potassium**
  - 2.6–3.0 mEq/L (mmol/L)
- **CO2 content**
  - 20–25 mEq/L (mmol/L)
- **PCO2**
  - 44–50 mm Hg (5.8–6.6 kPa)
- **PO2**
  - 40–44 mm Hg (5.3–5.8 kPa)
- **Calcium**
  - 2.0–2.8 mEq/L
- **Magnesium**
  - 2.4–3.0 mEq/L
- **Osmolality**
  - 280–300 mOsm/kg (280–300 mmol/kg)

**Serology and Microbiology**

- **VDRL**
  - Negative
- **Bacteria**
  - None present
- **Viruses**
  - None present
- **Antibody index**
  - >1.5 indicates chronic inflammatory process
  - <0.4 probably not acute inflammatory process

Be sure to include patient's age because it is needed to evaluate borderline values.

### Explanation of Tests

Cerebrospinal fluid, obtained by lumbar/intrathecal puncture, is the main diagnostic tool for neurologic disorders. A lumbar/intrathecal puncture is done for the following reasons:

1. To examine the spinal fluid for diagnosis of four major disease categories:
   a. Meningitis
   b. Subarachnoid hemorrhage
   c. CNS malignancy (meningeal carcinoma, tumor metastasis)
   d. Autoimmune disease and multiple sclerosis
2. To determine level of CSF pressure, to document impaired CSF flow, or to lower pressure by removing volume of fluid. (Fluid removal should be done with caution.)
3. To identify disease-related immunoglobulin patterns (IgG, IgA, and IgM referenced to albumin) in neurotuberculosis, neuroborreliosis (after a tick bite), or opportunistic infections.
4. To introduce anesthetics, drugs, or contrast media used for radiographic studies and nuclear scans into the spinal cord.
5. Confirm the identity of pathogens involved in acute inflammatory or chronic inflammatory disorders (eg, multiple sclerosis and blood-brain barrier dysfunction).
7. Formulate antibody index (AI) of the IgG class for polyspecific immune response in CNS. Examples: measles, rubella, and zoster (MRZ) antibodies to viruses in multiple sclerosis (MS); HSV antibodies in MS; toxoplasma antibodies in MS; and autoantibodies to dsDNA (double-stranded deoxyribonucleic acid).

8. Identify brain-derived proteins, such as neuron-specific enolase present after brain trauma.

See Figure 5.1 for an example of a CSF analysis report.

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**Clinical Alert**

The MRZ reaction occurs in MS, lupus erythematosus, Sjögren syndrome, or Wegener granulomatosis.

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Certain observations are made each time lumbar puncture is performed:

1. CSF pressure is measured.
2. General appearance, consistency, and tendency of the CSF to clot are noted.
3. CSF cell count is performed to distinguish types of cells present; this must be done within 2 hours of obtaining the CSF sample.
4. CSF protein and glucose concentrations are determined.
5. Other clinical serologic and bacteriologic tests are done when the patient's condition warrants (eg, culture for aerobes and anaerobes or tuberculosis).
6. Tumor markers may be present in CSF; these tests are useful as supplements to CSF cytology analysis (Table 5.2).

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**Table 5.2 Tumor Markers in CSF**

<table>
<thead>
<tr>
<th>Determination</th>
<th>Used in Diagnosis of</th>
<th>Normal Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alpha-fetoprotein (AFP)</td>
<td>CNS dysgerminomas and meningeal carcinomas</td>
<td>&lt;1.5 mg/mL (&lt;1.5 µg/L)</td>
</tr>
<tr>
<td>Beta-Glucuronidase</td>
<td>Possible meningeal adenocarcinoma</td>
<td>&lt;49 mU/L (&lt;0.82 nKat/L) normal; 47–70 mU/L (0.78–1.17 nKat/L), suspicious</td>
</tr>
<tr>
<td></td>
<td>Acute myeloblastic leukemia</td>
<td>&gt;70 mU/L (&gt;1.17 nKat/L) abnormal</td>
</tr>
<tr>
<td>Carcinoembryonic antigen (CEA)</td>
<td>Meningeal carcinomatosis; intradural or extradural, or brain parenchymal metastasis from adenocarcinoma; although the assay appears to be specific for adenocarcinoma and squamous cell carcinoma, increased CEA values in CSF are not seen in all such tumors of the brain</td>
<td>&lt;0.6 mg/mL (&lt;0.6 µg/L)</td>
</tr>
<tr>
<td>Human chorionic gonadotropin (HCG)</td>
<td>Adjunct in determining CNS dysgerminomas and meningeal carcinomatosis</td>
<td>&lt;0.21 U/L (&lt;1.5 IU/L)</td>
</tr>
<tr>
<td>Lysozyme (muramidase)</td>
<td>CNS tumors, especially myoclonal and monocytic leukemia</td>
<td>4–13 µg/mL</td>
</tr>
</tbody>
</table>

Note: The value of tumor markers in CSF for routine clinical diagnosis has not been established.

*Normal values vary greatly; check with your reference laboratory.*
Clinical Alert

1. Blood levels for specific substances should always be measured simultaneously with CSF determinations for meaningful interpretation of results.
2. Before lumbar puncture, check eyegrounds for evidence of papilledema, because its presence may signal potential problems or complications of lumbar puncture.
3. A mass lesion should be precluded by CT scan before lumbar puncture, because this can lead to brain stem herniation.
4. However, if increased pressure is found while performing the lumbar puncture, it should not be necessary to stop the procedure unless neurologic signs are present.

CEREBROSPINAL FLUID TESTS

Lumbar Puncture (Spinal Tap)

Procedure

1. Place the patient in a side-lying position with the head flexed onto the chest and knees drawn up to, but not compressing, the abdomen to “bow” the back. This position helps to increase the space between the lower lumbar vertebrae so that the spinal needle can be inserted more easily between the spinal processes. However, a sitting position with the head flexed to the chest can be used. The patient is helped to relax and instructed to breathe slowly and deeply with his or her mouth open.
2. Select the puncture site, usually between L4 and L5 or lower. There is a small bony landmark at the L5-S1 interspace known as the “surgeon’s delight” that helps to locate the puncture site. The site is thoroughly cleansed with an antiseptic solution, and the surrounding area is draped with sterile towels in such a way that the drapes do not obscure important landmarks (Fig. 5.2).

![FIGURE 5.2 Spinal tap technique. The patient lies on his side with knees flexed and back arched to separate the lumbar vertebrae. The patient is surgically draped and an area overlying the lumbar spine is disinfected (A). The space between lumbar vertebrae L₄ and L₅ is palpated with steriley gloved forefinger (B) and the spinal needle is carefully directed between the spinous processes, through the infraspinous ligaments into the spinal canal (C).]

3. Inject a local anesthetic slowly into the dermis around the intended puncture site.
4. Insert a spinal needle with stylet into the midline between the spines of the lumbar space and slowly advance until it enters the subarachnoid space. The patient may feel the entry as a “pop” of the needle through the dura mater. Once this happens, the patient can be helped to straighten his or her legs slowly to relieve abdominal compression.
5. Remove the stylet with the needle remaining in the subarachnoid space, and attach a pressure manometer to the needle to record the opening CSF pressure.
6. Remove a specimen consisting of up to 20 mL CSF. Take up to four samples of 2 to 3 mL each, place in separate sterile vials, and label sequentially. Tube 1 is used for chemistry and serology; tube 2 is used for microbiology studies; tube 3 is used for hematologic cell counts; and tube 4 is used for special studies such as cryptococcal antigens, syphils testing (Venereal Disease Research Laboratory [VDRL]), protein electrophoresis, and other immunologic studies. A closing pressure reading may be taken before the needle is withdrawn. In cases of increased intracranial pressure (ICP), no more than 2 mL is withdrawn because of the risk that the brain stem may shift.
7. Apply a small sterile dressing to the puncture site.
8. Label tubes correctly with the proper sequential number (1, 2, 3, or 4), the patient’s name, and the date of collection. Specimens of CSF must be immediately delivered to the laboratory, where they should be given to laboratory personnel with specific instructions regarding the testing. CSF samples should never be placed in the refrigerator because refrigeration alters the results of bacteriologic and fungal studies. Analysis should be started immediately. If viral studies are to be done, a portion of the specimen should be frozen.
9. Record procedure start and completion times, patient’s status, CSF appearance, and CSF pressure readings.

Procedural Alert

1. If the opening pressure is >200 mm H₂O in a relaxed patient, no more than 2 mL of CSF should be withdrawn.
2. If the initial pressure is normal, the Queckenstedt’s test may be done. (This test is not done if a central nervous system [CNS] tumor is suspected.) In this test, pressure is placed on both jugular veins to occlude them temporarily and to produce an acute rise in CSF pressure. Normally, pressure rapidly returns to average levels after jugular vein occlusion is removed. Total or partial spinal fluid blockage is diagnosed if the lumbar pressure fails to rise when both jugular veins are compressed or if the pressure requires >20 seconds to fall after compression is released.

Interventions

Pretest Patient Preparation

-
1. Explain the purpose, benefits, and risks of lumbar puncture and explain tests to be performed on the CSF specimen; present a step-by-step description of the actual procedure. Emphasize the need for patient cooperation. Assess for contraindications or impediments such as arthritis. Sedation or analgesia may be used.

2. Help the patient to relax by having him or her breathe slowly and deeply. The patient must refrain from breath holding, straining, moving, and talking during the procedure.

3. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have the patient lie prone (flat or horizontal, or on the abdomen) for approximately 4 to 8 hours. Turning from side to side is permitted as long as the body is kept in a horizontal position.

2. Women may have difficulty voiding in this position. The use of a fracture bedpan may help.

3. Fluids are encouraged to help prevent or relieve headache, which is a possible result of lumbar puncture.

4. Interpret test outcomes. Assess and monitor for abnormal outcomes and complications such as paralysis (or progression of paralysis, as with spinal tumor), hematoma, meningocele, asphyxiation of infants due to tracheal obstruction from pushing the head forward, and infection. Institute infection control precautions if test outcomes reveal an infectious process.

5. Observe for neurologic changes such as altered level of consciousness, change of pupils, change in temperature, increased blood pressure, irritability, and numbness and tingling sensations, especially in the lower extremities.

6. If headache occurs, administer analgesics as ordered and encourage a longer period of prone bed rest. If headache persists, a “blood patch” may need to be done, in which a small amount of the patient's own blood is introduced into the spinal canal at the same level that the canal was previously entered. For reasons not totally understood, this blood patch very effectively stops spinal headaches within a very short period.

7. Check the puncture site for leakage.

8. Document the procedure completion and any problems encountered or complaints voiced by the patient.

9. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Clinical Alert**

1. Extreme caution should be used when performing lumbar puncture:
   a. If ICP is elevated, especially in the presence of papilledema or split cranial sutures. However, with some cases of increased ICP, such as with a coma, intracranial bleeding, or suspected meningitis, the need to establish a diagnosis is absolutely essential and outweighs the risks of the procedure.
   b. A relative contraindication would be ICP from a suspected mass lesion. To reduce the risk for brain herniation, a less invasive procedure such as a CT scan or MRI should be done.

2. Contraindications to lumbar puncture include the following conditions:
   a. Suspected epidural infection
   b. Infection or severe dermatologic disease in the lumbar area, which may be introduced into the spinal canal
   c. Severe psychiatric or neurotic problems
   d. Chronic back pain
   e. Anatomic malformations, scarring in puncture site areas, or previous spinal surgery at the site

3. If there is CSF leakage at the puncture site, notify the physician immediately and document findings.

4. Follow standard precautions (see Appendix A) when handling CSF specimens.

**CSF Pressure**

The CSF pressure is directly related to pressure in the jugular and vertebral veins that connect with the intracranial dural sinuses and the spinal dura. In conditions such as congestive heart failure or obstruction of the superior vena cava, CSF pressure is increased, whereas in circulatory collapse, CSF pressure is decreased.

Pressure measurement is done to detect impairment of CSF flow or to lower the CSF pressure by removing a small volume of CSF fluid. Provided initial pressure is not elevated and there is no marked fall in the pressure as fluid is removed, 10 to 20 mL of CSF may be removed without danger to the patient. Elevation of the opening CSF pressure may be the only abnormality found in patients with cryptococcal meningitis and pseudotumor cerebri. Repeated lumbar punctures are performed for ICP elevation in cryptococcal meningitis to decrease the CSF pressure.

**Reference Values**

*Normal*  
Adult: 90–180 mm H₂O in the lateral recumbent position. (This value is position dependent and will change with a horizontal or sitting position.) Child (<8 years of age): 10–100 mm H₂O

**Procedure**

1. Measure the CSF pressure before any fluid is withdrawn.
2. Take up to four samples of 2 to 3 mL each, place in separate sterile vials, and label sequentially. Tube 1 is used for chemistry and serology; tube 2 is used for microbiology studies; tube 3 is used for hematology cell counts; and tube 4 is used for special studies.

**Clinical Implications**

1. Increases in CSF pressure can be a significant finding in the following conditions:
   a. Intracranial tumors; abscess; lesions
   b. Meningitis (bacterial, fungal, viral, or syphilitic)
   c. Hypoosmolality as a result of hemodialysis
   d. Congestive heart failure
e. Superior vena cava syndrome
g. Cerebral edema
h. Thrombosis of venous sinuses
i. Conditions inhibiting CSF absorption
2. Decreases in pressure can be a significant finding in the following conditions:
a. Circulatory collapse
b. Severe dehydration
c. Hyperosmolality
d. Leakage of spinal fluid
e. Spinal-subarachnoid block
3. Significant variations between opening and closing CSF pressure can be found in the following conditions:
a. Tumors or spinal blockage above the puncture site when there is a large pressure drop (no further fluid should be withdrawn)
b. Hydrocephalus when there is a small pressure drop that is indicative of a large CSF pool

Interfering Factors
1. Slight elevations of CSF pressure may occur in an anxious patient who holds his or her breath or tenses his or her muscles.
2. If the patient's knees are flexed too firmly against the abdomen, venous compression will cause an elevation in CSF pressure. This can occur in patients of normal weight and in those who are obese.

Interventions
Pretest Patient Preparation
1. See page 296 for care before lumbar puncture.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.
Posttest Patient Aftercare
1. Interpret abnormal pressure levels and monitor and intervene appropriately to prevent complications.
2. See page 296 for care after lumbar puncture.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

CSF Color and Appearance
Normal CSF is crystal clear, with the appearance and viscosity of water. Abnormal CSF may appear hazy, cloudy, smoky, or bloody. Clotting of CSF is abnormal and indicates increased protein or fibrinogen levels.

The initial appearance of CSF can provide various types of diagnostic information. Inflammatory diseases, hemorrhage, tumors, and trauma produce elevated cell counts and corresponding changes in appearance.

Reference Values
Normal Clear and colorless

Procedure Compare the CSF with a test tube of distilled water held against a white background. If there is no turbidity, newsprint can be read through normal CSF in the tube.

Clinical Implications
1. Abnormal appearance (Table 5.3)—causes and indications:

Table 5.3 Color Changes in CSF Suggestive of Disease States

<table>
<thead>
<tr>
<th>Appearance</th>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Opalescent, slightly yellow, with delicate clot</td>
<td>Tuberculous meningitis</td>
</tr>
<tr>
<td>Opalescent to purulent, slightly yellow, with coarse clot</td>
<td>Acute pyogenic meningitis</td>
</tr>
<tr>
<td>Slightly yellow; may be clear or opalescent, with delicate clot</td>
<td>Acute anterior poliomyelitis</td>
</tr>
<tr>
<td>Bloody, purulent, may be turbid</td>
<td>Primary amebic meningoencephalitis</td>
</tr>
<tr>
<td>Generally clear, but may be xanthochromic</td>
<td>Tumor of brain or cord</td>
</tr>
<tr>
<td>Xanthochromic</td>
<td>Toxoplasmosis</td>
</tr>
<tr>
<td>Viscous</td>
<td>Metastatic colon cancer, severe meningeal infection, cryptoccocus, injury</td>
</tr>
</tbody>
</table>

a. Blood. The blood is evenly mixed in all three tubes in subarachnoid and cerebral hemorrhage. Table 5.4 describes differentiation of bloody spinal tap from cerebral hemorrhage. Clear CSF fluid does not rule out intracranial hemorrhage.

Table 5.4 Differentiation of Bloody CSF Caused by Subarachnoid Hemorrhage Versus Traumatic Lumbar Puncture

<table>
<thead>
<tr>
<th>CSF Findings</th>
<th>Subarachnoid Hemorrhage</th>
<th>Traumatic Lumbar Puncture</th>
</tr>
</thead>
</table>
CSF Pressure
- Often increased
- Normal: Tubes 1 and 2 more bloody than tube 3 or 4

Blood in tubes for collecting CSF
- Mixture with blood is uniform in all tubes
- Often clots

CSF clotting
- Does not clot
- Present if >8–12 h since cerebral hemorrhage
- Absent unless patient is jaundiced

Xanthochromia
- Present if >8–12 h since cerebral hemorrhage
- Absent unless patient is jaundiced

Immediate repeat of lumbar puncture at higher level
- CSF same as initial puncture
- CSF clear (if atraumatic)

*CSF with RBCs >6000 per mm³ appears grossly bloody

b. Turbidity is graded from 1+ (slightly cloudy) to 4+ (very cloudy) and may be caused by the following conditions:
1. Leukocytes (pleocytosis)
2. Erythrocytes
3. Microorganisms such as fungi and amebae
4. Protein
5. Epidural fat aspirated (pale pink to dark yellow)
6. Contrast media

c. Xanthochromia (pale pink to dark yellow) can be caused by the following conditions:
1. Oxyhemoglobin from lysed red blood cells (RBCs) present in CSF before lumbar puncture
2. Methemoglobin
3. Bilirubin (>6 mg/dL or >103 µmol/L)
4. Increased protein (>150 mg/dL or >1500 mg/L)
5. Melanin (meningeal melanocarcinoma)
6. Carotene (systemic carotenemia)
7. Prior bleeding within 2–36 hours (eg, traumatic puncture >72 hours before)

d. Yellow color (bilirubin >10 mg/dL or >171 µmol/L) due to a prior hemorrhage (10 hour to 4 weeks before)

### Clinical Alert
1. Spinal fluid should be cultured for bacteria, fungi, and tuberculosis. In children, *Haemophilus influenzae* type B is the most common cause of bacterial meningitis; in adults, the most common bacterial pathogens for meningitis are meningococci and pneumococci.
2. Spinal fluid with any degree of cloudiness should be treated with extreme care because this could be an indication of contagious disease.

### Interfering Factors
1. CSF can look xanthochromic from contamination with methylate used to disinfect the skin.
2. If the blood in the specimen is due to a traumatic spinal tap, the CSF in the third tube should be clearer than that in tube 1 or 2; a traumatic tap makes interpretation of results very difficult to impossible.

### Interventions

#### Pretest Patient Preparation
1. Observations of color and appearance of CSF are always noted.
2. See page 296 for care before lumbar puncture.

#### Posttest Patient Aftercare
1. Recognize abnormal color and presence of turbidity and monitor patient appropriately.
2. See page 296 for care after lumbar puncture.

### CSF Microscopic Examination of Cells; Total Cell Count; Differential Cell Count

Normal CSF contains a small number of lymphocytes and monocytes in a ratio of approximately 70:30 in adults. A higher proportion of monocytes is present in young children. An increase in the number of white blood cells (WBCs) in CSF is termed pleocytosis. Disease processes may lead to abrupt increases or decreases in numbers of cells.

CSF is examined for the presence of RBCs and WBCs. The cells are counted and identified by cell type; the percentage of cell type is compared with the total number of WBCs or RBCs present. In general, inflammatory disease, hemorrhage, neoplasms, and trauma cause an elevated WBC count.

### Reference Values

**Normal** Normal CSF is essentially free of cells (Table 5.5 and Table 5.6).

<table>
<thead>
<tr>
<th>Differential</th>
<th>Adults</th>
<th>Newborn (0–14 d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lymphocytes</td>
<td>40%–80%</td>
<td>5%–35%</td>
</tr>
<tr>
<td>Monocytes</td>
<td>15%–45%</td>
<td>50%–90%</td>
</tr>
<tr>
<td>Polys (Neutrophils)</td>
<td>0%–6%</td>
<td>0%–8%</td>
</tr>
</tbody>
</table>
Table 5.6 Major Cells Seen in Microscopic Examination of CSF

<table>
<thead>
<tr>
<th>Cell Types</th>
<th>Occurrence</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blast forms</td>
<td>Acute leukemia</td>
<td>Lymphoblasts or myeloblasts</td>
</tr>
<tr>
<td>Ependymal and choroidal cells</td>
<td>Trauma (diagnostic procedures)</td>
<td>Clusters with distinct nuclei and distinct cell walls</td>
</tr>
<tr>
<td>Lymphocytes</td>
<td>Normal</td>
<td>All stages of development possible</td>
</tr>
<tr>
<td></td>
<td>Viral, tubercular, and fungal meningitis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Multiple sclerosis</td>
<td></td>
</tr>
<tr>
<td>Macrophages</td>
<td>Viral, tubercular, and fungal meningitis</td>
<td>May contain phagocytized RBCs (appearing as empty vacuoles or ghost cells) and hemosiderin granules</td>
</tr>
<tr>
<td></td>
<td>RBCs in spinal fluid</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Contrast media</td>
<td></td>
</tr>
<tr>
<td>Malignant cells</td>
<td>Metastatic carcinomas</td>
<td>Clusters with fusing of cell borders and nuclei</td>
</tr>
<tr>
<td>Monocytes</td>
<td>Chronic bacterial meningitis</td>
<td>Mixed with lymphocytes and neutrophils</td>
</tr>
<tr>
<td></td>
<td>Viral and tubercular meningitis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Multiple sclerosis</td>
<td></td>
</tr>
<tr>
<td>Neutrophils</td>
<td>Bacterial meningitis</td>
<td>Granules may be less prominent than in blood</td>
</tr>
<tr>
<td></td>
<td>Early cases of viral, tubercular, and fungal meningitis</td>
<td></td>
</tr>
<tr>
<td>Pia arachnoid mesothelial (PAM) cells</td>
<td>Normal, mixed reactions, including lymphocytes, neutrophils, monocytes, and plasma cells</td>
<td>Resemble young monocytes with a round, not indented, nucleus</td>
</tr>
<tr>
<td>Plasma cells</td>
<td>Multiple sclerosis</td>
<td>Transitional and classic forms seen</td>
</tr>
<tr>
<td></td>
<td>Tuberculosis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Meningitis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sarcoidosis</td>
<td></td>
</tr>
</tbody>
</table>

Adults: 0–5 WBCs/µL or 0–5 × 10^6 WBCs/L Newborn: 0–30 WBCs/µL or 0–30 × 10^6 WBCs/L Child: 0–15 WBCs/µL or 0–15 × 10^6 WBCs/L

Procedure Use tube 3 for counting the cells present in the CSF sample. The cells are counted by a manual counting chamber or by electronic means. A CSF smear is made, and various types of cells present are counted to determine differentiation of cells.

Clinical Implications

1. The total CSF cell count (includes neutrophils, lymphocytes, mixed cells, and cells after hemorrhage) is the most sensitive index of acute inflammation of the CNS.
2. WBC counts >500 WBCs/µL or >500 × 10^6 WBCs/L usually arise from a purulent infection and are preponderantly granulocytes (ie, neutrophils). Neutrophilic reaction classically suggests meningitis caused by a pyogenic organism, in which case the WBC count can exceed 1000 WBCs/µL or 1000 × 10^6 WBCs/L and even reach 20,000 WBCs/µL or 20,000 × 10^6 WBCs/L.
   a. Increases in neutrophils are associated with the following conditions:
      1. Bacterial meningitis (see Table 5.7)

Table 5.7 Abnormal CSF Findings in Types of Meningitis

<table>
<thead>
<tr>
<th></th>
<th>Bacterial</th>
<th>Viral</th>
<th>Tubercular</th>
<th>Fungal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total WBCs Differential</td>
<td>Increased</td>
<td>Increased</td>
<td>Increased</td>
<td>Increased</td>
</tr>
<tr>
<td></td>
<td>Neutrophils present</td>
<td>Lymphocytes present</td>
<td>Lymphocytes and monocytes present</td>
<td>Lymphocytes and monocytes present</td>
</tr>
<tr>
<td>Protein</td>
<td>Marked increase</td>
<td>Moderate increase</td>
<td>Moderate to marked increase; clots occur with protein &gt;150 mg/dL (&gt;1500 mg/L)</td>
<td>Moderate to marked increase</td>
</tr>
<tr>
<td>Glucose</td>
<td>Markedly decreased</td>
<td>Normal</td>
<td>Decreased</td>
<td>Normal to decreased</td>
</tr>
<tr>
<td>Lactate</td>
<td>Increased</td>
<td>Normal</td>
<td>Increased</td>
<td>Increased</td>
</tr>
<tr>
<td>LDH fractions</td>
<td>LD isoenzymes 4 and 5 increased</td>
<td>LD isoenzymes 1, 2, and 3 increased</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Limulus amebocyte Lysate: indicator of endotoxin produced by gram-negative bacteria (Not affected by antibiotic therapy)</td>
<td>Positive</td>
<td>—</td>
<td>Pellicle formation when protein &gt;150 mg/dL (&gt;1500 mg/L)</td>
<td>Positive India ink with neoformans</td>
</tr>
</tbody>
</table>
2. Early viral meningitis
3. Early tubercular meningitis
4. Fungal mycositic meningitis
5. Amoebic encephalomyelitis
6. Early stages of cerebral abscess

b. Noninfectious causes of neutrophilia include the following:
1. Reaction to CNS hemorrhage
2. Injection of foreign materials into the subarachnoid space (eg, x-ray contrast medium, anticancer drugs)
3. CSF infarct
4. Metastatic tumor in contact with CSF
5. Reaction to repeated lumbar puncture

3. WBC counts of 300–500/µL or 300–500 × 10⁶ with preponderantly lymphocytes are indicative of the following conditions:
   a. Viral meningitis
   b. Syphilis of CNS (ie, meningoencephalitis)
   c. Tuberculous meningitis
   d. Parasitic infestation of the CNS
   e. Bacterial meningitis due to unusual organisms (eg, *Listeria* species)
      f. Multiple sclerosis (MS) (reactive lymph present)
   g. Encephalopathy caused by drug abuse
   h. Guillain-Barré syndrome (15%)
      i. Acute disseminated encephalomyelitis
   j. Sarcoidosis of meninges
   k. Human T-lymphotropic virus type III (HTLV III)
   l. Aseptic meningitis due to peptic focus adjacent to meninges
   m. Fungal meningitis
   n. Polyneuritis

4. WBC counts with <40% monocytes occur in the following conditions:
   a. Chronic bacterial meningitis
   b. Toxoplasmosis and amoebic meningitis
   c. MS
   d. Rupture of brain abscess

5. Malignant cells (lymphocytes or histiocytes) may be present with primary and metastatic brain tumors, especially when there is meningeal extension.

6. Increased numbers of plasma cells occur in the following conditions:
   a. Acute viral infections
   b. MS
   c. Sarcoidosis
   d. Syphilitic meningoencephalitis
   e. Subacute sclerosing panencephalitis
   f. Tuberculous meningitis
   g. Parasitic infestations of CSF
   h. Guillain-Barré syndrome
      i. Lymphocytic reactions

7. Plasma cells are responsible for an increase in immunoglobulin G (IgG) and altered patterns in immunoelectrophoresis.

8. Macrophages are present in tuberculous or viral meningitis and in reactions to erythrocytes, foreign substances, or lipids in the CSF.

9. Ependymal and plexus cells may be present after surgical procedures or trauma to the CNS (not clinically significant).

10. Blast cells appear in CSF when acute leukemia is present (lymphoblasts or myeloblasts).

11. Eosinophils are present in the following conditions:
   a. Parasitic infections
   b. Fungal infections
   c. Rickettsial infections (Rocky Mountain spotted fever)
   d. Idiopathic hypereosinophilic syndrome
   e. Reaction to foreign materials in CSF (eg, drugs, shunts)
   f. Sarcoidosis

---

**Clinical Alert**

Neutrophilic reaction classically suggests meningitis caused by a pyogenic organism.

**Interventions**

**Pretest Patient Preparation**

1. See page 296 for [care before lumbar puncture](#).
2. Follow guidelines in Chapter 1 regarding [safe, effective, informed pretest care](#).

**Posttest Patient Aftercare**

1. Interpret abnormal cell counts. Monitor, intervene, and counsel as appropriate for infection and malignancy.
2. See page 296 for [care after lumbar puncture](#).
3. Follow guidelines in Chapter 1 regarding [safe, effective, informed posttest care](#).
CSF Glucose

The CSF glucose level varies with the blood glucose levels. It is usually about 60% of the blood glucose level. A blood glucose specimen should be obtained at least 60 minutes before lumbar puncture for comparisons. Any changes in blood sugar are reflected in the CSF approximately 1 hour later because of the lag in CSF glucose equilibrium time.

This measurement is helpful in determining impaired transport of glucose from plasma to CSF, increased use of glucose in the CNS, and glucose utilization by leukocytes and microorganisms. The finding of a markedly decreased CSF glucose level accompanied by an increased WBC count with a large percentage of neutrophils is indicative of bacterial meningitis.

Reference Values

| Normal | Adult: 40–70 mg/dL or 2.2–3.9 mmol/L | Child: 60–80 mg/dL or 3.3–4.4 mmol/L | CSF-to-plasma glucose ratio: <0.5 |

Procedure
Place 1 mL of CSF in a sterile tube. The glucose test should be done on tube 1 when three tubes of CSF are taken. Accurate evaluation of CSF glucose requires a plasma glucose measurement. A blood glucose level ideally should be drawn 1 hour before the lumbar puncture.

Clinical Implications

1. Decreased CSF glucose levels are associated with the following conditions:
   a. Acute bacterial meningitis
   b. Tuberculosis, fungal, and amebic meningitis
   c. Systemic hypoglycemia
   d. Subarachnoid hemorrhage
2. CSF glucose levels are uncommonly decreased in the following conditions:
   a. Malignant tumor with meningeal involvement
   b. Acute syphilitic meningitis
   c. Nonbacterial meningoencephalitis
3. Increased CSF glucose levels are associated with diabetes and diabetic hyperglycemia. Elevated CSF levels are always a result of high plasma values.

Interfering Factors

1. Falsely decreased levels may be due to cellular and bacterial metabolism if the test is not performed immediately after specimen collection.
2. A traumatic tap may produce misleading results owing to glucose present in blood.
3. See Appendix J for drugs that affect test outcomes.

Clinical Alert

1. All types of organisms consume glucose; therefore, decreased glucose levels reflect abnormal activity.
2. The panic value for CSF glucose level is <20 mg/dL (<1.1 mmol/L); below this level, damage to the CNS will occur.
3. The findings of a markedly decreased CSF glucose and an increased WBC count with a high percentage of neutrophils are indicative of bacterial meningitis.

Interventions

Pretest Patient Preparation

1. See page 296 for care before lumbar puncture.
2. Explain the need for a blood specimen test for glucose to compare with CSF glucose.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret abnormal CSF glucose levels and correlate with the presence of meningitis, cancer, hemorrhage, and diabetes. Monitor and intervene appropriately to prevent complications.
2. See page 296 for care after lumbar puncture.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

CSF Glutamine

Glutamine is synthesized in brain tissue from ammonia and alpha-ketoglutarate. Production of glutamine provides a mechanism for removing the ammonia, a toxic metabolic waste product, from the CNS.

The determination of CSF glutamine level provides an indirect test for the presence of excess ammonia in the CSF. As the concentration of ammonia in the CSF increases, the supply of alpha-ketoglutarate becomes depleted; consequently, glutamine can no longer be produced to remove the toxic ammonia, and coma ensues. A CSF glutamine test is therefore frequently requested for patients with coma of unknown origin. A glutamine value of >35 mg/dL (>2.4 mmol/L) usually results in loss of consciousness.

Reference Values

| Normal | 8–18 mg/dL or 0.4–1.2 mmol/L |
Procedure

1. Use 1 mL of CSF for the glutamine test. Tube 1 is used for this chemistry test.
2. Centrifuge the samples if cells are present.

Clinical Implications Increased CSF glutamine levels are associated with the following conditions:

1. Hepatic encephalopathy (glutamine values >35 mg/dL or >2.4 mmol/L are diagnostic)
2. Reye’s syndrome
3. Encephalopathy secondary to hypercapnia or sepsis

Interventions

Pretest Patient Preparation

1. See page 296 for care before lumbar puncture.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret abnormal glutamine levels and correlate with clinical symptoms. Monitor and intervene appropriately to prevent complications.
2. See page 296 for care after lumbar puncture.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

CSF Lactic Acid

The source of CSF lactic acid is CNS anaerobic metabolism. Lactic acid in CSF varies independently with the level of lactic acid in the blood. Destruction of tissue within the CNS because of oxygen deprivation causes the production of increased CSF lactic acid levels. Thus, elevated CSF lactic acid levels can result from any condition that decreases the flow of oxygen to brain tissues.

The CSF lactic acid test is used to differentiate between bacterial and nonbacterial meningitis. Elevated CSF lactate levels are not limited to meningitis and can result from any condition that decreases the flow of oxygen to the brain. CSF lactate levels are frequently used to monitor severe head injuries.

Reference Values

Normal Adult: 10–22 mg/dL or 1.1–2.4 mmol/L Newborn: 10–60 mg/dL or 1.1–6.7 mmol/L

Procedure

1. Collect 0.5 mL of CSF in a sterile test tube; tube 1 should be used.
2. Refrigerate the sample.

Clinical Implications Increased CSF lactic acid levels are associated with the following conditions:

1. Bacterial meningitis (>38 mg/dL or >4.2 mmol/L)
2. Brain abscess or tumor
3. Cerebral ischemia
4. Cerebral trauma
5. Seizures
6. Stroke (cerebral infarct)

Interfering Factors Traumatic tap causes elevated levels: RBCs contain large amounts of lactate. Hemolyzed or xanthochromic specimens will give falsely elevated results.

Interventions

Pretest Patient Preparation

1. See page 296 for care before lumbar puncture.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes; monitor and intervene appropriately to detect CNS disease and prevent complications. Results must be interpreted in light of clinical symptoms.
2. See page 296 for care after lumbar puncture.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Clinical Alert

Increases in CSF lactic acid levels must be interpreted in light of the clinical findings and in conjunction with glucose levels, protein levels, and cell counts in the CSF. Equivocal results in some instances of aseptic meningitis may lead to erroneous diagnosis of a bacterial etiology. Increased lactate in CSF following head injury suggests poor prognosis.
CSF Lactate Dehydrogenase (LD/LDH); CSF Lactate Dehydrogenase (LDH) Isoenzymes

Although many different enzymes have been measured in CSF, only lactate dehydrogenase (LDH) appears useful clinically. Sources of LDH in normal CSF include diffusion across the blood-CSF barrier, diffusion across the brain-CSF barrier, and LDH activity in cellular elements of the CSF, such as leukocytes, bacteria, and tumor cells. Because brain tissue is rich in LDH, damaged CNS tissue can cause increased levels of LDH in the CSF.

High levels of LDH occur in about 90% of cases of bacterial meningitis and in only 10% of cases of viral meningitis. When high levels of LDH do occur in viral meningitis, the condition is usually associated with encephalitis and a poor prognosis. Tests of LDH isoenzymes have been used to improve the specificity of LDH measurements and are useful for making the differential diagnosis of viral versus bacterial meningitis (see Chap. 6 for a complete description of isoenzymes). Elevated LDH levels following resuscitation predict a poor outcome in patients with hypoxic brain injury.

Reference Values
Normal Adults: <20 U/L or approximately 10% of serum levels

Procedure
1. Obtain 1 mL of CSF for the LDH test; use tube 1 for LDH examination.
2. Take the sample to the laboratory as quickly as possible.

Clinical Implications
1. Increased CSF/LDH levels are associated with the following conditions:
   a. Bacterial meningitis (90% of cases)
   b. Viral meningitis (10% of cases)
   c. Massive cerebrovascular accident
   d. Leukemia or lymphoma with meningeal infiltration
   e. Metastatic carcinoma of the CNS
2. The presence of CSF/LDH isoenzymes 1, 2, and 3 reflects a CNS lymphocytic reaction, suggesting viral meningitis.
3. The CSF/LDH isoenzyme pattern reflects a granulocytic (neutrophilic) reaction with LDH isoenzymes 4 and 5, suggesting bacterial meningitis.
4. High levels of CSF/LDH isoenzymes 1 and 2 suggest extensive CNS damage and a poor prognosis (ie, they are indicative of destruction of brain tissue).
5. CSF/LDH isoenzymes 3 and 4 suggest lymphatic leukemia or lymphoma.

Interfering Factors For the LDH test to be valid, CSF must not be contaminated with blood. A traumatic lumbar tap will make results difficult to interpret.

Interventions
Pretest Patient Preparation
1. See page 296 for care before lumbar puncture.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret abnormal LDH test patterns and monitor and intervene appropriately to detect and prevent complications.
2. See page 296 for care after lumbar puncture.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

CSF Total Protein

The CSF normally contains very little protein because the protein in the blood plasma does not cross the blood-brain barrier easily. Protein concentration normally increases caudally from the ventricles to the cisterns and finally to the lumbar sac.

The CSF protein is a nonspecific but reliable indication of CNS pathology such as meningitis, brain abscess, MS, and other degenerative processes causing neoplastic disease. Elevated CSF protein levels may be caused by increased permeability of the blood-brain barrier, decreased resorption of the arachnoid villi, mechanical obstruction of the CSF flow, or increased intrathecal immunologic synthesis.

Reference Values
Normal Results vary by method used; check with the laboratory for reference values. Total protein: Adults: 15–45 mg/dL or 150–450 mg/L (lumbar) Adults: 15–25 mg/dL or 150–250 mg/L (cisternal) Adults: 5–15 mg/dL or 50–150 mg/L (ventricular) Neonates: 15–100 mg/dL or 150–1000 mg/L (lumbar) Elderly patients (>60 years of age): 15–60 mg/dL or 150–600 mg/L (lumbar)

Procedure
1. Obtain 1 mL of CSF for protein analysis. Tube 1 should be used.
2. Measure serum protein levels concurrently to interpret CSF protein values.

Clinical Implications
1. Increased CSF protein occurs in the following situations:
a. Traumatic tap with normal CSF pressure: CSF initially streaked with blood, clearing in subsequent tubes
b. Increased permeability of blood-CSF barrier: CSF protein 100–500 mg/dL (1000–5000 mg/L)
   1. Infectious conditions
      i. Bacterial meningitis: Gram stain usually positive; culture may be negative if antibiotics have been administered
      ii. Tuberculosis: CSF protein 50–300 mg/dL (500–3000 mg/L); mixed cellular reaction typical
      iii. Fungal meningitis: CSF protein 50–300 mg/dL (500–3000 mg/L); special stains helpful
      iv. Viral meningitis: CSF protein usually <200 mg/dL (<2000 mg/L)
   2. Noninfectious conditions
      i. Subarachnoid hemorrhage: xanthochromia 2–4 hours after onset
      ii. Intracerebral hemorrhage: CSF protein 20–200 mg/dL (200–2000 mg/L); marked fall in pressure after removing small amounts of CSF; xanthochromia
      iii. Cerebral thrombosis: slightly increased CSF protein in 40% of cases (usually, <100 mg/dL or <1000 mg/L)
      iv. Endocrine disorders, diabetic neuropathy, myxedema, hyperadrenalism, hypoparathyroidism: CSF protein 50–150 mg/dL (500–1500 mg/L) in ~50% of cases
      v. Metabolic disorders, uremia, hypercalcemia, hypercapnia, dehydration: CSF protein slightly elevated (usually, <100 mg/dL or <1000 mg/L)
      vi. Drug toxicity, ethanol, phenytoin, phenothiazines: CSF protein slightly elevated in about 40% of cases (usually, <200 mg/dL or <2000 mg/L)
c. Obstruction to circulation of CSF occurs in the following circumstances:
   1. Mechanical obstruction (eg, tumor, abscess), herniated disk: rapid fall in pressure (yellow CSF, contains excess protein)
   2. Loculated effusion of CSF: repeated taps may show a progressive increase in CSF protein; diagnosis by myelography
d. Increased CSF/IgG synthesis occurs in the following conditions:
   1. MS: CSF protein level slightly increased
   2. Subacute sclerosing panencephalitis: increased CSF protein
   3. Neurosyphilis: CSF protein normal or slightly increased (usually, <100 mg/dL or <1000 mg/L)
e. Increased CSF/IgG synthesis and increased permeability of blood-CSF barrier occur in the following conditions:
   1. Guillain-Barré syndrome (infectious polyneuritis): CSF protein usually 100–400 mg/dL (1000–4000 mg/L)
   2. Collagen diseases (eg, periarteritis, lupus): CSF protein usually < 400 mg/dL (or <4000 mg/L)
   3. Chronic inflammatory demyelinating polyradiculopathy
f. Decreased CSF protein occurs in the following conditions:
   1. Leakage of CSF due to trauma
   2. Removal of a large volume of CSF
   3. Intracranial hypertension
   4. Hyperthyroidism
   5. Young children between 6 months and 2 years of age

**Clinical Alert**

More than 1000 mg/dL (>10,000 mg/L) of protein in CSF suggests subarachnoid block. In a complete spinal block, the lower the tumor location, the higher the CSF protein value.

**Interfering Factors**

1. Hemolyzed or xanthochromic drugs may falsely depress results.
2. Traumatic tap will invalidate the protein results.
3. See Appendix J for drugs that affect test outcomes.

**Interventions**

**Pretest Patient Preparation**

1. See page 296 for *care before lumbar puncture*.
2. Follow guidelines in Chapter 1 regarding *safe, effective, informed pretest care*.

**Posttest Patient Aftercare**

1. Interpret abnormal CSF protein levels; monitor for both infectious and noninfectious conditions and intervene appropriately to prevent and detect complications.
2. See page 296 for *care after lumbar puncture*.
3. Follow guidelines in Chapter 1 regarding *safe, effective, informed posttest care*.

**CSF Albumin and Immunoglobulin G (IgG)**

Albumin composes the majority of the proteins in CSF. The albumin and IgG that are present in normal CSF are derived from the serum. Increased levels of either or both are indicative of damage to the blood-CNS barrier.

The combined measurement of albumin and IgG is used to evaluate the integrity and permeability of the blood-CSF barrier and to measure the synthesis of IgG within the CNS. The IgG index is the most sensitive method to determine local CNS synthesis of IgG and to detect increased permeability of the blood-CNS barrier.
The IgG index method is superior to the IgG-to-albumin ratio or measurement of IgG only. Normal persons usually have an index < 0.60. With MS, the index is > 0.77.

Reference Values

<table>
<thead>
<tr>
<th>Normal</th>
<th>Albumin: 10–35 mg/dL or 1.5–5.3 µmol/L</th>
<th>IgG: &lt;4.0 mg/dL or &lt;40 mg/L</th>
<th>CSF-to-serum albumin index: &lt;9.0</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IgG index: 0.3–0.60</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>IgG index: 0.3–0.60</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Procedure

1. Obtain 0.5 mL of CSF in a sterile tube.
2. Freeze the sample if the determination is not done immediately.

Clinical Implications

1. Increased CSF albumin occurs in most of the same conditions as increased total protein, especially:
   a. Lesions of the choroid plexus
   b. Blockage of CSF flow
   c. Bacterial meningitis
d. Guillain-Barré syndrome
e. Many infectious diseases, such as typhoid fever, tularemia, diphtheria, and septicemia
f. Malignant neoplasms of the CNS
2. Increased CSF/IgG-to-albumin index (increased IgG, normal albumin) occurs in the following conditions:
   a. MS
   b. Subacute sclerosing leukoencephalitis
c. Neurosyphilis
d. Chronic phases of CNS infections (subacute sclerosing panencephalitis [SSPE])

Interfering Factors

A traumatic tap will invalidate the results.

Interventions

Pretest Patient Preparation

1. See page 296 for care before lumbar puncture.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes; monitor and intervene appropriately to prevent and detect complications.
2. See page 296 for care after lumbar puncture.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

CSF Protein Electrophoresis; Oligoclonal Bands; Multiple Sclerosis Panel

Agarose gel electrophoresis of concentrated CSF is used to detect oligoclonal bands, defined as two or more discrete bands in the gamma region that are absent or of less intensity than in the concurrently tested patient's serum.

Fractionation (ie, electrophoresis) of CSF is used to evaluate bacterial and viral infections and tumors of the CNS. However, the most important application of CSF protein electrophoresis is the detection and diagnosis of MS. Abnormalities of CSF in MS include an increase in total protein, primarily from IgG, which is the main component of the gamma-globulin fraction. Abnormal immunoglobulins migrate as discrete, sharp bands, called oligoclonal bands. This is the pattern observed in MS: a pattern of discrete bands within the gamma-globulin portion of the electrophoretic pattern. However, oligoclonal bands are found in the CSF of patients with other types of nervous system disorders of the immune system, including human immunodeficiency virus (HIV).

Electrophoresis is also the method of choice to determine whether a fluid is actually CSF. Identification can be made based on the appearance of an extra band of transferrin (referred to as “TAV”), which occurs in CSF and not in serum.

Reference Values

| Normal Globalins: Oligoclonal banding: none present; alpha 1: 2%–7% IgG synthesis rate: 0.0–8.0 mg/day; alpha 2: 4%–12% IgG-to-albumin ratio: 0.09–0.25; beta: 8%–18%; Prealbumin: 2%–7%; gamma: 3%–12% Albumin: 56%–76% |
|---------|-------------------------------------------------|---------------------------------|------------------------|

Procedure

1. Obtain 3 mL of CSF for this test. Use tube 1. The sample must be frozen if the test is not performed immediately.
2. Apply a sample of the concentrate to a thin-layer agarose gel. Subject the agarose gel to electrophoresis. CSF is concentrated approximately 80-fold by selective permeability. Serum electrophoresis must be done concurrently for interpretation of the bands.

Clinical Implications
1. Increases in CSF IgG or in the IgG-to-albumin index occur in the following conditions:
   a. MS
   b. Subacute sclerosis panencephalitis
   c. Tumors of the brain and meninges
   d. Chronic CNS infections
   e. Some patients with meningitis, Guillain-Barré syndrome, lupus erythematosus involving the CNS, and other neurologic conditions

2. Increases in the CSF albumin index occur in the following conditions:
   a. Obstruction of CSF circulation
   b. Damage to the CNS blood-brain barrier
   c. Diabetes mellitus
   d. Systemic lupus erythematosus of the CNS
   e. Guillain-Barré syndrome
   f. Polyneuropathy
   g. Cervical spondylosis

3. Increased CSF gamma-globulin and the presence of oligoclonal bands occur in the following conditions:
   a. MS
   b. Neurosyphilis
   c. Subacute sclerosing panencephalitis
   d. Cerebral infarction
   e. Viral and bacterial meningitis
   f. Progressive rubella panencephalitis
   g. Cryptococcal meningitis
   h. Idiopathic polyneuropathies
   i. Burkitt’s lymphoma
   j. HIV-1 (acquired immunodeficiency syndrome [AIDS])
   k. Guillain-Barré syndrome

4. Increased CSF synthesis of IgG occurs in the following conditions:
   a. MS (90% of definite cases)
   b. Inflammatory neurologic diseases
   c. Postpolio syndrome

**Clinical Alert**

1. A serum electrophoresis must be done at the same time as the CSF electrophoresis. An abnormal result is the finding of two or more bands in the CSF that are not present in the serum specimen. (See Figure 5.3.)


2. Oligoclonal bands are not specific for multiple sclerosis; however, the sensitivity is 83% to 94%.
3. Diagnostic differentiation between MS and CSF autoimmune disease relies on further testing (eg, antinuclear antibodies [ANAs] in blood [see Chap. 8]).

**Interfering Factors**

1. A traumatic tap invalidates the results.
2. Recent myelography affects the results.

**Interventions**

**Pretest Patient Preparation**

1. See page 296 for care before lumbar puncture.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome; monitor for MS and other CNS disorders and intervene appropriately to prevent and detect complications.
2. See page 296 for care after lumbar puncture.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

CSF Syphilis Serology

Reference Values

**Normal** Negative (ie, nonreactive) for syphilis. Neurosyphilis is characterized by an increase in protein, an increase in the number of lymphocytes, and a positive test for syphilis (see Chap. 8). Use CSF VDRL test, only if serum VDRL test is positive, to rule in, not rule out, neurosyphilis. Do not use VDRL to evaluate the results of syphilis therapy.

BIBLIOGRAPHY


OVERVIEW OF CHEMISTRY STUDIES

General Biochemical Profiles

Use of the Autoanalyzer

NOTE

DIABETES TESTING (TYPE 1 AND TYPE 2), BLOOD GLUCOSE, BLOOD SUGAR, AND RELATED TESTS AND CRITERIA FOR DIAGNOSING DIABETES

C-Peptide
Glucagon
Insulin

Fasting Blood Glucose (FBG); Fasting Blood Sugar (FBS); Fasting Plasma Glucose (FPG); Casual Plasma Glucose (PG)

Clinical Alert

Gestational Diabetes Mellitus (GDM); O'Sullivan Test (1-h Gestational Diabetes Mellitus Screen)

Glucose Tolerance Test (GTT); Oral Glucose Tolerance Test (OGTT)

Glycosylated Hemoglobin (Hb A1c); Glycohemoglobin (G-Hb); Glycated Hemoglobin (GhB); Diabetic Control Index; Glycated Serum Protein (GSP), Fructosamine

Lactose Tolerance; Breath Hydrogen Test

END PRODUCTS OF METABOLISM AND OTHER TESTS

Ammonia (NH3)
Bilirubin
Neonatal Bilirubin, Total and Fractionated (“Baby Bili”)

Blood Urea Nitrogen (BUN, Urea Nitrogen)

Albumin

Porphobilinogen (PAB)

CHOLESTEROL AND OTHER LIPIDS

Cholesterol

Cystatin C

Uric Acid

Lead (Pb)

Osteocalcin (Bone GLA Protein)

HORMONE TESTS

Androstenedione

Aldosterone

Antidiuretic Hormone (ADH); Arginine Vasopressin Hormone

Atrial Natriuretic Factor (ANF), ANP and BNP

Chart 6.1 Heart Failure

Chart 6.2 Grading Heart Diseases

Cortisol (Hydrocortisone)

Cortisol Suppression (Dexamethasone Suppression; DST)

Cortisol Stimulation (Cosyntropin, Cortrosyn Stimulation); Adrenocorticotropin Hormone (ACTH) Stimulation

Gastrin

Growth Hormone (HGH), Somatotropin

Parathyroid Hormone Assay, Parathyrin; Parathormone (PTH-C-Terminal)

Somatomedin C (SM-C), Insulin-like Growth Hormone

FERTILITY TESTS

Chorionic Gonadotropin; Human Chorionic Gonadotropin (hCG) ß Subunit; Pregnancy Test

Prostate-Specific Antigen (PSA)

Ankylase and Lipase

Aspartate Transaminase (Aminotransferase, AST); Serum Glutamic-Oxaloacetic Transaminase (SGOT)

Creatine Phosphokinase (CPK); Creatine Kinase (CK)

GASTROINTESTINAL ENZYME TESTS

Amylase and Lipase

Aspartate Transaminase (Aminotransferase, AST); Serum Glutamic-Oxaloacetic Transaminase (SGOT)

Cardiac Troponin T (cTnT); Troponin I (cTnI)

Creatine Phosphokinase (CPK); Creatine Kinase (CK); CPK and CK Isoenzymes

Galactose-1-Phosphate Uridylytransferase (GPT); Galactokinase; Galactose-1-Phosphosphate

Hexosaminidase, Total and Isoenzyme A

Lactate Dehydrogenase (LD, LDH)

Lactate Dehydrogenase (LDH, LD) Isoenzymes (Electrophoresis)

Renin (Angiotensin); Plasma Renin Angiotensin (PRA)

Renin Stimulation/Challenges Test

Interpretation of Renin Stimulation Test

?-Glutamyltransferase (?-Glutamyl Transpeptidase, GGT, ?GT)

Homocysteine (tHcy)

Chart 6.5 Homocysteine Testing

A1-Antitrypsin (AA1)
OVERVIEW OF CHEMISTRY STUDIES

Blood chemistry testing identifies many chemical blood constituents. It is often necessary to measure several blood chemicals to establish a pattern of abnormalities. A wide range of tests can be grouped under the headings of enzymes, electrolytes, blood sugars, lipids, hormones, proteins, vitamins, minerals, and drug investigation. Other tests have no common denominator. Selected tests serve as screening devices to identify target organ damage. When collecting specimens for chemistry studies, refer to Standard/Universal Precautions in Appendix A, Latex and Rubber Allergy Precautions in Appendix B, and Guidelines for Specimen Transport and Storage in Appendix E, and always refer to Appendix J for Effects of the Mostly Commonly Used Drugs on Frequently Ordered Laboratory Tests.

General Biochemical Profiles

Profiles are a group of select tests that screen for certain conditions. Some of the more common profiles or panels are listed in Table 6.1.

Table 6.1 Common Screening Profiles

<table>
<thead>
<tr>
<th>Group Headings</th>
<th>Tests Suggested</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiac markers (MI)</td>
<td>Chem panels, cardiac troponin, CK, MB, homocysteine</td>
</tr>
<tr>
<td>Electrolyte panel</td>
<td>Na, K, Cl, CO₂, pH</td>
</tr>
<tr>
<td>Kidney functions/disease</td>
<td>BUN, phosphorus, LDH, creatinine, creatinine clearance, total protein, A/G ratio, albumin, calcium, glucose, CO₂</td>
</tr>
<tr>
<td>Lipids (coronary risk)</td>
<td>Cholesterol, triglycerides, HDL, lipoprotein electrophoresis (LDL, VLDL, HDL)</td>
</tr>
<tr>
<td>Liver function/disease</td>
<td>Total bilirubin, alkaline phosphatase, GGT, total protein, A/G ratio, albumin, AST, LDH, viral hepatitis panel, PT</td>
</tr>
<tr>
<td>Thyroid function</td>
<td>T₃ uptake, free T₄, Total T₄, T₇, FTI, TSH</td>
</tr>
<tr>
<td>Basic metabolic screen</td>
<td>Chloride, sodium, potassium, carbon dioxide, glucose, BUN, creatinine</td>
</tr>
<tr>
<td>Syndrome X (metabolic syndrome)</td>
<td>Blood lipid glucose</td>
</tr>
</tbody>
</table>

A/G ratio, albumin/globulin ratio; AST, aspartate aminotransferase; BUN, blood urea nitrogen; CK, creatine kinase; FTI, free thyroxine index; HDL, high-density lipoproteins; LDH, lactate dehydrogenase; LDL, low-density lipoproteins; PT, prothrombin time; TSH, thyroid-stimulating hormone; VLDL, very-low-density lipoprotein.

Use of the Autoanalyzer

Sophisticated automated instrumentation makes it possible to conduct a wide variety of chemical tests on a single sample of blood and to report results in a timely manner. Numerical results may be reported with low, high, panic, toxic, or D (ie, fails Delta check) comments along with normal reference range. Computerized interfaces allow direct transmission of results between laboratory and clinical settings. "Hard copy" printouts can then become a permanent part of the health care record. Not only does this method of record keeping provide a baseline for future comparisons, but it also allows unsuspected diseases to be uncovered and can lead to early diagnosis when symptoms are vague or absent. Chemistry tests may be termed chem panels, SMAC, chem 2 zyme profiles, and SMAS. These terms refer to the company that produces the auto analyzer. A list of standard panels follows in Table 6.2.
### Table 6.2 Standard Panels

<table>
<thead>
<tr>
<th>Panel Tests</th>
<th>Specimen Collection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ARTHITIS PANEL (ARTH PN)</strong></td>
<td></td>
</tr>
<tr>
<td>Uric acid, ESR, ANA (antinuclear antibody screen), rheumatoid factor</td>
<td>Two 7-mL red topped tubes and 1 lavender-topped tube</td>
</tr>
<tr>
<td><strong>BASIC METABOLIC PANEL (BC MET)</strong></td>
<td></td>
</tr>
<tr>
<td>Creatinine, CO₂, chloride, glucose, potassium, sodium, BUN, calcium</td>
<td>1 mL unhemolyzed serum (one SST tube)</td>
</tr>
<tr>
<td><strong>COMPREHENSIVE METABOLIC PANEL (CM MET)</strong></td>
<td></td>
</tr>
<tr>
<td>Albumin, chloride, creatinine, glucose, potassium, sodium, total protein, BUN</td>
<td>1 mL unhemolyzed serum (one SST tube)</td>
</tr>
<tr>
<td><strong>ELECTROLYTES (LYTES)</strong></td>
<td></td>
</tr>
<tr>
<td>CO₂, chloride, potassium, sodium</td>
<td>1 mL unhemolyzed serum (one SST tube)</td>
</tr>
<tr>
<td><strong>HEPATIC FUNCTION PANEL (HEPFUN)</strong></td>
<td></td>
</tr>
<tr>
<td>ALT, albumin, alkaline phosphatase, AST, direct bilirubin and total bilirubin, total protein</td>
<td>1 mL unhemolyzed serum (one SST tube)</td>
</tr>
<tr>
<td><strong>ACUTE HEPATITIS PANEL (ACUTE HEP)</strong></td>
<td></td>
</tr>
<tr>
<td>Hepatitis A, AB, IgM, hepatitis B core antibody, IgM, hepatitis B surface antigen, IgM, hepatitis C, AB</td>
<td>One 7-mL red-topped tube</td>
</tr>
<tr>
<td><strong>LIPID PANEL (LIPID PN)</strong></td>
<td></td>
</tr>
<tr>
<td>Cholesterol, HDL, triglycerides (LDL and CHO/HDL ratio included, as calculated values)</td>
<td>2 mL serum (one SST tube)</td>
</tr>
<tr>
<td><strong>OBSTETRIC PANEL (OB PN)</strong></td>
<td></td>
</tr>
<tr>
<td>Type and Rh, antibody screen, RPR, rubella Ab-IgG, hepatitis B surface antigen</td>
<td>One 7-mL red-topped tube, one lavender-topped tube, and one SST tube</td>
</tr>
<tr>
<td><strong>CBC WITH DIFF</strong></td>
<td></td>
</tr>
<tr>
<td>Type and Rh, antibody screening and studies if indicated, RPR for syphilis, rubella Ab-IgG, hepatitis B surface antigen</td>
<td>One lavender-topped tube, one 7-mL red-topped tube, and one SST tube</td>
</tr>
<tr>
<td><strong>PRENATAL SCREEN (PRESCP)</strong></td>
<td></td>
</tr>
<tr>
<td>Type and RH, antibody screening and studies if indicated, RPR for syphilis, rubella Ab-IgG, hepatitis B surface antigen</td>
<td>One lavender-topped tube, one 7-mL red-topped tube, and one SST tube</td>
</tr>
</tbody>
</table>

### NOTE

Normal or reference values for any chemistry determination vary with the method or assay employed. For example, differences in substrates or temperature at which the assay is run will alter the “normal” range. Thus, “normal ranges” vary from laboratory to laboratory.

The following is a list of routine automated tests performed commonly in the chemistry department:

1. Alanine aminotransferase (ALT)
2. Albumin
3. Alkaline phosphatase
4. Amylase
5. Aspartate aminotransferase (AST)
6. Bilirubin, direct
7. Bilirubin, total
8. Calcium
9. Carbon dioxide (CO₂)
10. Chloride
11. Cholesterol
12. Cholesterol-HDL
13. Creatine kinase
14. Creatinine
15. ?-Glutamyl transferase (GGT)
16. Glucose
17. Iron
18. Lactate dehydrogenase (LDH)
19. LDL cholesterol (calculated)
20. Magnesium
21. Phosphorous, inorganic
22. Potassium
23. Protein, total
24. Sodium
25. Total iron binding (calculated)
26. Triglycerides
27. Unbound iron binding (UIBC)
28. Urea nitrogen
29. Uric acid

**DIABETES TESTING (TYPE 1 AND TYPE 2), BLOOD GLUCOSE, BLOOD SUGAR, AND RELATED**
TESTS AND CRITERIA FOR DIAGNOSING DIABETES

C-Peptide

C-peptide is formed during the conversion of pro-insulin to insulin. Pro-insulin is cleaved (holds α and β insulin chains together in the pro-insulin molecule) into insulin and biologically inactive C-peptide. C-peptide assay provides distinction between exogenous and endogenous circulating insulin.

The main use of C-peptide is to evaluate hypoglycemia. C-peptide levels provide reliable indicators for pancreatic, β, and secretory functions and insulin secretions. In a patient with type 1 diabetes mellitus, C-peptide measurements can be an index of insulin production and mark endogenous β-cell activity. C-peptide levels can also be used to confirm suspected surreptitious insulin injections (i.e., factitious hypoglycemia). Findings in these patients reveal that insulin levels are usually high, insulin antibodies may be high, but C-peptide levels are low or undetectable. This test also monitors the patient's recovery after excision of an insulinoma. Rising C-peptide levels suggest insulinoma tumor recurrence or metastases.

Reference Values

Normal Fasting: 0.51–2.72 ng/mL or 0.17–0.90 mmol/L Varies with laboratory.

Procedure

1. Draw a 1-mL venous blood sample from a fasting patient using a red-topped chilled tube. Serum is needed for test. Date and time must be correct. Centrifuge blood for 30 minutes.
2. Separate the blood at 4°C and freeze if it will not be tested until later.
3. Remember that a sample for glucose testing is usually drawn at the same time.

Clinical Implications

1. Increased C-peptide values occur in the following conditions:
   a. Endogenous hyperinsulinism (insulinemia)
   b. Oral hypoglycemic drug ingestion
   c. Pancreas or β-cell transplantation
   d. Insulin-secreting neoplasms (islet cell tumor)
   e. Type 2 diabetes mellitus (non-insulin-dependent)
2. Decreased C-peptide values occur in the following conditions:
   a. Factitious hypoglycemia (surreptitious insulin administration)
   b. Radical pancreatectomy
   c. Type 1 diabetes mellitus
3. C-peptide stimulation test can determine the following:
   a. Distinguishes between type 1 and type 2 diabetes mellitus.
   b. Patients with diabetes whose C-peptide stimulation values are >1.8 ng/mL (>0.59 nmol/L) can be managed without insulin treatment.

Interfering Factors

Increased C-peptide:

1. Renal failure
2. Ingestion of sulfonylurea

<table>
<thead>
<tr>
<th>Clinical Alert</th>
</tr>
</thead>
<tbody>
<tr>
<td>To differentiate insulinoma from factitious hypoglycemia, an insulin/C-peptide ratio can be performed.</td>
</tr>
<tr>
<td>&lt;1.0 Ratio: increased endogenous insulin secretion</td>
</tr>
<tr>
<td>&gt;1.0 Ratio: exogenous insulin</td>
</tr>
</tbody>
</table>

Interventions

Pretest Patient Care

1. Explain the test purpose and blood-drawing procedure. Obtain history of signs and symptoms of hypoglycemia.
2. Ensure that the patient fasts, except for water, for 8 to 12 hours before blood is drawn.
3. Remember that radiisotope test, if necessary, should take place after blood is drawn for C-peptide levels.
4. If the C-peptide stimulation test is done, give IV glucagon after a baseline value blood sample is drawn.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume normal activities.
2. Interpret test results and monitor as appropriate. Explain possible need for further testing. See Chapter 8 for insulin antibody testing.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Glucagon

Glucagon is a peptide hormone that originates in the α cells of the pancreatic islets of Langerhans. This hormone
promotes glucose production in the liver. Normally, glucagon is a counterbalance to insulin. Glucagon provides a sensitive, coordinated control mechanism for glucose production and storage. For example, low blood glucose levels cause glucagon to stimulate glucose release into the bloodstream, whereas elevated blood glucose levels reduce the amount of circulating glucagon to about 50% of that found in the fasting state. The kidneys also affect glucagon metabolism. Elevated fasting glucagon levels in the presence of renal failure return to normal levels following successful renal transplantation. Abnormally high glucagon levels drop toward normal once insulin therapy effectively controls diabetes. However, when compared with a healthy person, glucagon secretion in the person with diabetes does not decrease after eating carbohydrates. Moreover, in healthy persons, arginine infusion causes increased glucagon secretion.

This test measures glucagon production and metabolism. A glucagon deficiency reflects pancreatic tissue loss. Failure of glucagon levels to rise during arginine infusion confirms glucagon deficiency. Hyperglucagonemia (ie, elevated glucagon levels) occurs in diabetes, acute pancreatitis, and situations in which catecholamine secretion is stimulated (eg, pheochromocytoma, infection).

Reference Values

**Normal**
- Adults: 20–100 pg/mL or 20–100 ng/L
- Children: 0–148 pg/mL or 0–148 ng/L
- Newborns: 0–1750 pg/mL or 0–1750 ng/L

Normal ranges vary with different laboratories.

**Clinical Alert**

During a glucose tolerance test (GTT) in healthy persons, glucagon levels will decline significantly compared with baseline fasting levels as normal hyperglycemia takes place during the first hour of testing.

**Procedure**

1. Draw a 5-mL blood sample from a fasting person into a chilled EDTA Vacutainer tube containing aprotinin (Trasylol) proteinase inhibitor. Special handling is required because glucagon is very prone to enzymatic degradation. Tubes used to draw blood must be chilled before the sample is collected and placed on ice afterward, and plasma must be frozen as soon as possible after centrifuging.
2. Observe standard precautions. Place specimen in a biohazard bag.

**Clinical Implications**

1. **Increased glucagon levels** are associated with the following conditions:
   a. Acute pancreatitis (eg, pancreatic a-cell tumor)
   b. Diabetes mellitus: persons with severe diabetic ketoacidosis are reported to have fasting glucagon levels five times normal despite marked hyperglycemia.
   c. Glucagonoma (familial) may be manifested by three different syndromes:
      1. The first syndrome exhibits a characteristic skin rash, necrolytic migratory erythema, diabetes mellitus or impaired glucose tolerance, weight loss, anemia, and venous thrombosis. This form usually shows elevated glucagon levels (>1000 pg/mL or >1000 ng/L) (diagnostic).
      2. The second syndrome occurs with severe diabetes.
      3. The third form is associated with multiple endocrine neoplasia syndrome and can show relatively lower glucagon levels as compared with the others.
   d. Chronic renal failure
   e. Hyperlipidemia
   f. Stress (trauma, burns, surgery)
   g. Uremia
   h. Hepatic cirrhosis
   i. Hyperosmolality
   j. Acute pancreatitis
   k. Hypoglycemia
2. **Reduced levels of glucagon** are associated with the following conditions:
   a. Loss of pancreatic tissue
      1. Pancreatic neoplasms
      2. Pancreatectomy
   b. Chronic pancreatitis
   c. Cystic fibrosis

**NOTE**

After glucose load, there is no suppression of glucagon in patients with glucagonoma.

**Interventions**

**Pretest Patient Care**

1. Explain purpose of test and blood-drawing procedure. A minimum 8-hour fast (no calorie intake for at least 8 hours) is necessary before the test.
2. Promote relaxation in a low-stress environment; stress alters normal glucagon levels.
3. Do not administer radiopharmaceuticals within 1 week before the test.
4. Follow guidelines in Chapter 1 for *safe, effective, informed pretest care*.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. Interpret test outcome and monitor for the three different syndromes of glucagonoma.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Insulin**

Insulin, a hormone produced by the pancreatic β cells of the islets of Langerhans, regulates carbohydrate metabolism together with contributions from the liver, adipose tissue, and other target cells. Insulin is responsible for maintaining blood glucose levels at a constant level within a defined range. The rate of insulin secretion is primarily regulated by the level of blood glucose perfusing the pancreas; however, it can also be affected by hormones, the autonomic nervous system, and nutritional status.

Insulin levels are valuable for establishing the process of an insulinoma (ie, tumor of the islets of Langerhans). This test is also used for investigating the causes of fasting hypoglycemic states and neoplasm differentiation. The insulin study can be done in conjunction with a GTT or fasting blood glucose (FBG) test or a fasting plasma glucose (FPG) test.

**Reference Values**

- **Normal Immunoreactive**
  - Adults: 0–35 µIU/mL or 0–243 pmol/L
  - Children: 0–10 µIU/mL or 0–69 pmol/L
- **Free**
  - Adults: 0–17 µIU/mL or 0–118 pmol/L
  - Children (prepubertal): 0–13 µIU/mL or 0–90 pmol/L

**Procedure**

1. Obtain a 5-mL blood sample from a fasting person; serum is preferred. Observe standard precautions. Heparinized blood may be used.
2. If done in conjunction with a GTT, draw the specimens before administering oral glucose and again 30, 60, and 120 minutes after glucose ingestion (the same times as the GTT).

**Clinical Implications**

1. **Increased insulin values** are associated with the following conditions:
   a. Insulinoma (pancreatic islet tumor). Diagnosis is based on the following findings:
      1. Hyperinsulinemia with hypoglycemia (glucose <30 mg/dL or <1.66 mmol/L)
      2. Persistent hypoglycemia together with hyperinsulinemia (>20 µIU/mL or >139 pmol/L) after tolbutamide injection (rapid rise and rapid fall)
      3. Failed C-peptide suppression with a plasma glucose level <30 mg/dL or <1.66 mmol/L and insulin/glucose ratio >0.3.
   b. Type 2 diabetes mellitus, untreated
   c. Acromegaly
   d. Cushing's syndrome
   e. Endogenous administration of insulin (factitious hypoglycemia)
   f. Obesity (most common cause)
   g. Pancreatic islet cell hyperplasia
2. **Decreased insulin values** are found in the following conditions:
   a. Type 1 diabetes mellitus, severe
   b. Hypopituitarism

**Clinical Alert**

Panic range: >35 µIU/mL or >243 pmol/L (fasting)

**Interfering Factors**

1. Surreptitious insulin or oral hypoglycemic agent ingestion or injection causes elevated insulin levels (with low C-peptide values).
2. Oral contraceptives and other drugs cause falsely elevated values.
3. Recently administered radioisotopes affect test results.
4. In the second to third trimester of pregnancy, there is a relative insulin resistance with a progressive decrease of plasma glucose and immunoreactive insulin.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. Ensure that the patient fasts from all food and fluid, except water, unless otherwise directed.
3. Be aware that because insulin release from an insulinoma may be erratic and unpredictable, it may be necessary for the patient to fast for as long as 72 hours before the test.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activity and diet.
2. Interpret test results and counsel appropriately. Obese patients may have insulin resistance and unusually high fasting and postprandial (after eating) insulin levels. Explain possible need for further testing and treatment.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.
A potentially fatal situation may exist if the insulinoma secretes unpredictably high levels of insulin. In this case, the blood glucose may drop to such dangerously low levels as to render the person comatose and unable to self-administer oral glucose forms. Patients and their families must learn how to deal with such an emergency and to be vigilant until the problem is treated.

Fasting Blood Glucose (FBG); Fasting Blood Sugar (FBS); Fasting Plasma Glucose (FPG); Casual Plasma Glucose (PG)

Glucose is formed from carbohydrate digestion and conversion of glycogen to glucose by the liver. The two hormones that directly regulate blood glucose are glucagon and insulin. Glucagon accelerates glycogen breakdown in the liver and causes the blood glucose level to rise. Insulin increases cell membrane permeability to glucose, transports glucose into cells (for metabolism), provided there are enough insulin receptors. Insulin binds to these receptors on the surface of target cells such as are found in fat and muscle. This opens the channels so that glucose can pass into cells, where it can be converted into energy. As cellular glucose metabolism occurs, blood glucose levels fall. Adrenocorticotropic hormone (ACTH), adrenocorticosteroids, epinephrine, and thyroxine also play key roles in glucose metabolism. See Chapter 11 for genetic causes of type 1 and type 2 diabetes mellitus.

The American Diabetes Association (ADA) has begun using the term pre-diabetes, also known as impaired glucose tolerance or impaired fasting glucose. Individuals with pre-diabetes demonstrate higher levels of blood plasma glucose (PG) (110–125 mg/dL or 6.1–6.9 mmol/L) than normals (<110 mg/dL or <6.1 mmol/L) and, if left untreated, go on to develop type 2 diabetes within 10 years.


In most cases, significantly elevated fasting plasma glucose levels (ie, >140 mg/dL or >7.77 mmol/L; hyperglycemia) are, in themselves, usually diagnostic of diabetes. However, mild, borderline cases may present with normal fasting glucose values. If diabetes is suspected, a GTT can confirm the diagnosis. Occasionally, other diseases may produce elevated plasma glucose levels; therefore, a comprehensive history, physical examination, and workup should be done before a definitive diagnosis of diabetes is established.

**Clinical Alert**

A. New NIH guidelines endorse diabetic testing of all adults = 45 years every 3 years. The American Diabetes Association recommends the following guidelines for testing:
1. Testing should be considered if patient is >45 years of age.
2. Testing is strongly recommended if patient is >45 years of age and overweight.
3. Testing should be considered if patient is <45 years of age and overweight with another risk factor.

B. Diabetes mellitus, a group of metabolic disorders, is characterized by hyperglycemia and abnormal protein, fat, and carbohydrate metabolism due to defects in insulin secretion, ie, inadequate and deficient insulin action on target organs, or both.

C. 1. Symptoms of diabetes plus random/casual plasma glucose concentration ≥200 mg/dL (11.1 mmol/L). Random/casual is defined as any time of day without regard to time since last meal. The classic symptoms of diabetes include polyuria, polydipsia, and unexplained weight loss.
2. or FPG =126 mg/dL (7.0 mmol/L). Fasting is defined as no caloric intake for at least 8 h.
3. or 2-h PG =200 mg/dL (11.1 mmol/L) during an OGTT. The test should be performed as described by WHO, using a glucose load containing the equivalent of 75 g anhydrous glucose dissolved in water.

In the absence of unequivocal hyperglycemia with acute, metabolic decompensation, these criteria should be confirmed by repeat testing on a different day. OGTT is not recommended for routine clinical use. Source: Diabetes Care 25: 742–749, 2000.

**Reference Values**
Normal Fasting adults: =110 mg/dL or =6.1 mmol/L Fasting children (2–18 years): 60–100 mg/dL or 3.3–5.6 mmol/L Fasting young children (0–2 years): 60–110 mg/dL or 3.3–6.1 mmol/L Fasting premature infants: 40–65 mg/dL or 2.2–3.6 mmol/L

Procedure

1. Draw a 5-mL venous blood sample from a fasting person. In known cases of diabetes, blood drawing should precede insulin or oral hypoglycemic administration. Observe standard precautions. Serum is acceptable if separated from red cells within an hour. A gray-topped tube, which contains sodium chloride, is acceptable for 24 hours without separation.

2. Be aware that self-monitoring of blood glucose by the person with diabetes can be done by finger-stick blood drop sampling several times per day if necessary. Several devices are commercially available for this procedure; they are relatively easy to use and have been established as a major component in satisfactory diabetes control. Calibration of monitoring devices should be done on a regular basis.

3. Be aware that noninvasive methods using skin pads to check blood glucose level are being developed for self-monitoring that eliminate the dreaded finger-prick test, for example, a Gluco-watch (Cygnes, Redwood City, CA), worn on the wrist and powered by an AAA battery.

NOTE

When whole blood glucose values are not equivalent to plasma values, plasma values are about 10–15% higher than whole blood values. However, some of the newer meters now convert the whole blood values to plasma, thus giving a better comparison between the lab values and bedside or home testing.

Patient Checklist for Self-Monitoring of Blood Glucose (SMBG) Testing This list is a general outline. Each brand of meter has its own instructions. Read the instructions on each new meter carefully to get accurate results. Know whether your monitor and strips give whole blood or plasma results.

1. General instructions
   a. Make sure your hands are clean, dry, and warm.
   b. Prick your finger with the lancet.
   c. Squeeze out a drop of capillary blood.
   d. Drop the blood onto the test strip or sensor.
   e. Wait for the test strip or sensor to develop.
   f. Compare the test strip to the chart or insert it in the meter.
   g. Safely dispose of your lancet in an approved sharps container.
   h. Record blood glucose results with date and time.

2. If you have type 1 diabetes mellitus, you should also monitor your urine for ketones to alert you to possibly dangerous complications such as diabetic ketoacidosis (eg, during stress or acute illness).

3. Test more often on days when you are ill, when your blood glucose is too high, when your meal or exercise plan changes, when you travel, or if you feel that your blood glucose is low.

4. If you do not feel you are getting accurate results, talk to your diabetes educator and/or contact the manufacturer of your meter. Make sure you are using the meter properly.

5. Several blood glucose meters currently available are approved by the U.S. Food and Drug Administration (FDA), the agency that approves medical devices, for what's called "alternate site testing."

6. Alternate sites (other than fingertips) include forearm, bicep area, palm of hand, between fingers, and sometimes the calf.

7. Tips for using alternate sites:
   a. Rub the site you will use to check your blood glucose vigorously before you prick your skin. This increases blood flow to the site.
   b. Use one type of meter. Do not alternate between different meters. This will help you get consistent results.
   c. Consistently use the same alternate site. For example, always use your forearms. This will help you get consistent results.

Clinical Implications

1. Elevated blood glucose (hyperglycemia) occurs in the following conditions:
   a. Diabetes mellitus: a fasting glucose of >126 mg/dL (>7.0 mmol/L) or a 2-hour postprandial load plasma glucose...
>200 mg/dL (>11.1 mmol/L) during an oral GTT.

b. Other conditions that produce elevated blood plasma glucose levels include the following:
   1. Cushing's disease (increased glucocorticoids cause elevated blood sugar levels)
   2. Acute emotional or physical stress situations (eg, myocardial infarction [MI], cerebrovascular accident, convulsions)
   3. Pheochromocytoma, acromegaly, gigantism
   4. Pituitary adenoma (increased secretion or growth hormone causes elevated blood glucose levels)
   5. Hemochromatosis
   6. Pancreatitis (acute and chronic), neoplasms of pancreas
   7. Glucagonoma
   8. Advanced liver disease
   9. Chronic renal disease
   10. Vitamin B deficiency: Wernicke's encephalopathy
   11. Pregnancy (may signal potential for onset of diabetes later in life)

2. **Decreased blood plasma glucose (hypoglycemia)** occurs in the following conditions:
   a. Pancreatic islet cell carcinoma (insulinomas)
   b. Extrapancreatic stomach tumors (carcinoma)
   c. Addison's disease (adrenal insufficiency), carcinoma of adrenal gland
   d. Hypopituitarism, hypothyroid, ACTH deficiency
   e. Starvation, malabsorption (starvation does not cause hypoglycemia in normal persons)
   f. Liver damage (alcoholism, chloroform poisoning, arsenic poisoning)
   g. Premature infant; infant delivered to a mother with diabetes
   h. Enzyme-deficiency diseases (eg, galactosemia, inherited maple syrup disease, von Gierke's syndrome)
      i. Insulin overdose (accidental or deliberate)
   j. Reactive hypoglycemia, including alimentary hyperinsulinism, pre-diabetes, endocrine deficiency
   k. Postprandial hypoglycemia may occur after GI surgery and is described with hereditary fructose intolerance, galactosemia, and leucine sensitivity.

3. According to the ADA criteria, there are three definitive tests for diabetes:
   a. Symptoms of diabetes plus a random/casual plasma glucose >200 mg/dL (>11.1 mmol/L), or
   b. A fasting plasma glucose >126 mg/dL (>6.99 mmol/L), or
   c. An oral glucose tolerance test with a 2-hour postload (75-g glucose load) level >200 mg/dL (>11.1 mmol/L)

4. Using any of the three methods, the criterion must be reconfirmed on a subsequent day.

5. The classification of diabetes diagnosis reflects a shift to the etiology or pathology of the disease from a classification based on pharmacological treatment.

6. Impaired fasting glucose (IFG) or impaired glucose tolerance (IGT) is referred to as pre-diabetes. See Figure 6.1.

### Interfering Factors

1. **Elevated glucose:**
   a. Steroids, diuretics, other drugs (see Appendix J)
   b. Pregnancy (a slight blood glucose elevation normally occurs)
   c. Surgical procedures and anesthesia
   d. Obesity or sedentary lifestyle
   e. Parenteral glucose administration (eg, from total parenteral nutrition)
   f. IV glucose (recent or current)
   g. Heavy smoking

2. **Decreased glucose:**
   a. Hematocrit >55%
   b. Intense exercise
   c. Toxic doses of aspirin, salicylates, and acetaminophen
   d. Other drugs, including ethanol, quinine, and haloperidol

### Interventions

#### Pretest Patient Care

1. Explain test purpose and blood-drawing procedure.
2. Tell patient that the test requires at least an overnight fast; water is permitted. Instruct the patient to defer insulin or oral hypoglycemics until after blood is drawn, unless specifically instructed to do otherwise.
3. Note the last time the patient ate in the record and on the laboratory requisition.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

#### Posttest Patient Aftercare

1. Tell the patient that he or she may eat and drink after blood is drawn.
2. Interpret test results and monitor appropriately for hyperglycemia and hypoglycemia. Counsel regarding necessary lifestyle changes (eg, diet, exercise, glucose monitoring, medication).
3. Give the patient the following checklist:
   a. Take special care of your feet.
   b. Use a lubricant or unscented hand cream on dry, scaly skin.
   c. Look for calluses on your soles. Rub them gently with a pumice stone.
   d. Make sure new shoes fit properly; wear freshly washed socks or stockings.
   e. Never go barefoot.
   f. Avoid using hot water bottles, tubs of hot water, or heating pads on your feet.
   g. Trim your toenails straight across.
   h. Make sure your doctor inspects your feet as part of every visit.
   i. Use a team approach to help you make decisions about your care. The team may include your doctor, a nurse
diabetes educator, a dietitian, your pharmacist, and your family.

j. Use other health professionals to help with your care. These may include an eye doctor (ophthalmologist or optometrist), an exercise physiologist, a podiatrist (a foot specialist), and a psychologist.

k. Follow the most healthful lifestyle you can.

4. Persons with glucose levels >200 mg/dL (>11.1 mmol/L) should be placed on a strict intake and output program.

5. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

1. If a person with known or suspected diabetes experiences headaches, irritability, dizziness, weakness, fainting, or impaired cognition, a blood glucose test or finger-stick test must be done before giving insulin. Similar symptoms may be present for both hypoglycemia and hyperglycemia. If a blood glucose level cannot be obtained and one is uncertain regarding the situation, glucose may be given in the form of orange juice, sugar-containing soda, or candy (eg, Life-Savers or jelly beans). Make certain the person is sufficiently conscious to manage eating or swallowing. In the acute care setting, IV glucose may be given if a blood glucose level cannot be obtained and one is unsure of the situation. Instruct persons prone to hypoglycemia to carry sugar-type items on their person and to wear a necklace or bracelet that identifies the person as diabetic.

2. Frequent blood glucose monitoring, including self-monitoring, allows better control and management of diabetes than urine glucose monitoring.

3. When blood glucose values are > 300 mg/dL (>16.6 mmol/L), urine output increases, as does the risk for dehydration.

4. Panic values/critical values for fasting blood glucose: <40 mg/dL (<2.22 mmol/L) may cause brain damage (women and children), <50 mg/dL (<2.77 mmol/L) (men); >400 mg/dL (>22.2 mmol/L) may cause coma

5. Diabetes is a “disease of the moment”: persons living with diabetes are continually affected by fluctuations in blood glucose levels and must learn to manage and adapt their lifestyle within this framework. For some, adaptation is relatively straightforward; for others, especially those identified as being “brittle,” lifestyle changes and management are more complicated, and these patients require constant vigilance, attention, encouragement, and support.

6. Each person with diabetes may experience certain symptoms in his or her own unique way and in a unique pattern.

Clinical Alert

1. Infants with tremor, convulsion, or respiratory distress should have STAT glucose done, particularly in the presence of maternal diabetes, or with hemolytic disease of the newborn.

2. Newborns that are too small or too large for gestational age should have glucose level measured in the first day of life.

3. Diseases related to neonatal hypoglycemia:
   a. Glycogen storage diseases
   b. Galactosemia
   c. Hereditary fructose intolerance
   d. Ketogenic hypoglycemia of infancy
   e. Carnitine deficiency (Reye's syndrome)

Gestational Diabetes Mellitus (GDM); O’Sullivan Test (1-h Gestational Diabetes Mellitus Screen)

Glucose intolerance during pregnancy (gestational diabetes mellitus [GDM]) is associated with an increase in perinatal morbidity and mortality, especially in women who are aged >25 years, overweight, or hypertensive. Additionally, more than one half of all pregnant patients with an abnormal GTT do not have any of the same risk factors. It is therefore recommended that all pregnant women be screened for gestational diabetes.

The O’Sullivan test, based on an OGTT, is done to detect gestational diabetes and screen nonsymptomatic pregnant women. During pregnancy, abnormal carbohydrate metabolism is evaluated by screening all pregnant women at first prenatal visit, then again at 24 to 28 weeks.

One-step approach: An oral glucose load of 50 g is administered, and blood is examined for glucose levels 1 hour after administration. Women with a family history of diabetes or previous gestational diabetes should undergo the O’Sullivan test at 15 to 19 weeks of gestation and again at 24 to 28 weeks of gestation.

Two-step approach: Measure plasma or serum glucose 1 hour after GTT (glucose challenge).

Reference Values
Normal 130–140 mg/dL or 7.2–7.9 mmol/L (1 hour after 50 g of glucose)

Procedure

1. Draw a 5-mL venous blood sample (sodium fluoride) after 8–14 hour fast, at least 3 days of unrestricted diet and activity after glucose load.

2. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications

1. Abnormal GDM test result is after 100 g glucose load and after 75 g glucose load reveals glucose intolerance.

2. A 3-hour gestational GTT must then be done.
3. A positive result in a pregnant woman means she is at much greater risk (7 times) for having gestational diabetes mellitus (GDM).
4. GDM is any degree of glucose intolerance with onset during pregnancy or first recognized pregnancy.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose (to evaluate abnormal carbohydrate metabolism and predict diabetes in later life) and procedure. No fasting is usually required. Obtain pertinent history of diabetes and record any signs or symptoms of diabetes.
2. Instruct the woman about obtaining a urine sample for glucose testing to check before drinking the glucose load. Positive urine glucose should be checked with the physician before glucose load. Those with glycosuria >250 mg/dL (>13.8 mmol/L) must have a blood glucose test before O'Sullivan or GDM testing.
3. Give the patient 75–100 g of glucose beverage (150 mL dissolved in water, or Trutol or Orange DEX).
4. Explain to the patient that no eating, drinking, smoking, or gum chewing is allowed during the test. The patient should not leave the office. She may void if necessary.
5. After 1 hour, draw one NaFl or EDTA tube (5-mL venous blood) using standard venipuncture technique. If a 75 g glucose is given, also collect a 2-hour specimen. If a 100 g glucose load is given, obtain 2- and 3-hour specimens.

**Posttest Patient Aftercare**

1. Normal activities, eating, and drinking may be resumed.
2. Interpret test results and explain to patient that a normal outcome is <140 mg/dL (<7.8 mmol/L).
3. A follow-up 3-hour gestational GTT or OGTT is indicated for all abnormal screenings.
4. Six weeks after delivery, the patient should be retested and reclassified. In most cases, glucose will return to normal.

**Glucose Tolerance Test (GTT); Oral Glucose Tolerance Test (OGTT)**

In a healthy individual, the insulin response to a large oral glucose dose is almost immediate. It peaks in 30 to 60 minutes and returns to normal levels within 3 hours when sufficient insulin is present to metabolize the glucose ingested at the beginning of the test. The test should be performed according to WHO guidelines using glucose load containing the equivalent of 75–100 g of anhydrous glucose dissolved in water or other solution.

If fasting and postload glucose test results are borderline, the GTT can support or rule out a diagnosis of diabetes mellitus; it can also be a part of a workup for unexplained hypertriglyceridemia, neuropathy, impotence, renal diseases, or retinopathy. This test may be ordered when there is sugar in the urine or when the fasting blood sugar level is significantly elevated. The GTT/OGTT should not be used as a screening test for nonpregnant adults or children.

**Indications for Test** The GTT/OGTT should be done on certain patients, particularly those with the following indications (few indications still meet wide acceptance):

1. Family history of diabetes
2. Obesity
3. Unexplained episodes of hypoglycemia
4. History of recurrent infections (boils and abscesses)
5. In women, history of delivery of large infants, stillbirths, neonatal death, premature labor, and spontaneous abortions
6. Transitory glycosuria or hyperglycemia during pregnancy, surgery, trauma, stress, MI, and ACTH administration

**Reference Values**

**Normal Fasting plasma glucose (PG):** Adults: 110 mg/dL or 6.1 mmol/L 30-minute: Adults: 110–170 mg/dL or 6.1–9.4 mmol/L 60-minute (1-hour) plasma glucose (PG) after glucose load: Adults: <184 mg/dL or <10.2 mmol/L. 120-minute (2-hour GTT test) 2-hour plasma glucose (PG) after glucose load: Adults: <138 mg/dL or <7.7 mmol/L. Children: <140 mg/dL or <7.8 mmol/L

3-hour plasma glucose (PG) after glucose load: Adults: 70–120 mg/L or 3.9–6.7 mmol/L. All four blood values must be within normal limits to be considered normal.

**Procedure** This is a timed test for glucose tolerance. A 2-hour plasma glucose test is done after glucose load to detect diabetes in individuals other than pregnant women; the 3-hour test is done for pregnant women; and the 4-hour test evaluates possible hypoglycemia.

1. Have patient eat a diet of >150 g of carbohydrates for 3 days before the test.
2. Ensure that the following drugs are discontinued 3 days before the test because they may influence test results:
   a. Hormones, oral contraceptives, steroids
   b. Salicylates, anti-inflammatory drugs
   c. Diuretic agents
   d. Hypoglycemic agents
   e. Antihypertensive drugs
   f. Anticonvulsants (see Appendix J)
3. Insulin and oral hypoglycemics should be withheld until the test is completed.
4. Record the patient's weight.
   a. Pediatric doses of glucose are based on body weight, calculated as 1.75 g/kg not to exceed a total of 75 g.
   b. Pregnant women: 100 g glucose
   c. Nonpregnant adults: 75 g glucose
   d. Possible gestational diabetes: 100 g glucose
5. A 5-mL sample of venous blood is drawn. Serum or gray-topped tubes are used. The patient should fast 12 to 16
hours before testing. After the blood is drawn, the patient drinks all of a specially formulated glucose solution within a 5-minute time frame.

6. Blood samples are obtained 30 minutes, 1 hour, 2 hours, and 3 hours after glucose ingestion.

7. Specimens taken 4 hours after ingestion are significant for detecting hypoglycemia and may be ordered (5-hour specimens have been discredited).

8. Tolerance tests can also be performed for pentose, lactose, galactose, and d-xylose.

9. The GTT is not indicated in these situations:
   a. Persistent fasting hyperglycemia >140 mg/dL or >7.8 mmol/L
   b. Persistent fasting normal plasma glucose
   c. Patients with overt diabetes mellitus
   d. Persistent 2-hour plasma glucose >200 mg/dL or >11.1 mmol/L

10. Test has limited value in diagnosis of diabetes mellitus in children and is rarely indicated for that purpose.

Clinical Implications

1. The presence of abnormal GTT values (decreased tolerance to glucose) is based on the International Classification for Diabetes Mellitus and the following glucose intolerance categories:
   a. At least two GTT values must be abnormal for a diagnosis of diabetes mellitus to be validated.
   b. In cases of overt diabetes, no insulin is secreted; abnormally high glucose levels persist throughout the test.
   c. Glucose values that fall above normal values but below the diagnostic criteria for diabetes or impaired glucose tolerance (IGT) should be considered nondiagnostic.

2. See Table 6.3 for an interpretation of glucose tolerance levels.

<table>
<thead>
<tr>
<th>Table 6.3 Glucose Tolerance Test (GTT) Levels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conventional Units (mg/dL)</td>
</tr>
<tr>
<td>-----------------------------------------------</td>
</tr>
<tr>
<td>Fasting adult</td>
</tr>
<tr>
<td>Adult diabetes mellitus 1-h glucose</td>
</tr>
<tr>
<td>and 2-h glucose</td>
</tr>
<tr>
<td>Fasting adult</td>
</tr>
<tr>
<td>Adult impaired glucose tolerance 1-h glucose</td>
</tr>
<tr>
<td>and 2-h glucose</td>
</tr>
<tr>
<td>Juvenile diabetes mellitus (fasting glucose)</td>
</tr>
<tr>
<td>and 1-h glucose</td>
</tr>
<tr>
<td>and 2-h glucose</td>
</tr>
<tr>
<td>Impaired glucose tolerance in children (fasting glucose) and 2-h glucose</td>
</tr>
</tbody>
</table>

3. A diagnosis of gestational diabetes mellitus (GDM) is based on the following blood glucose results (more than two tests must be met and exceeded): fasting, >95 mg/dL (>5.3 mmol/L); 1-hour, >180 mg/dL (>10.8 mmol/L); 2-hour, >155 mg/dL (>8.6 mmol/L); and 3-hour, >140 mg/dL (>7.8 mmol/L).
   a. All pregnant women should be tested for gestational diabetes with a 50-g dose of glucose at 24 to 28 weeks of gestation. Pregnant women with abnormal GTT are at risk for preeclampsia/eclampsia and delivery of a large infant.
   b. If abnormal results occur during pregnancy, repeat GTT at the first postpartum visit.
   c. During labor, maintain maternal glucose levels at 80 to 100 mg/dL (4.4–5.5 mmol/L); beware of markedly increased insulin sensitivity in the immediate postpartum period.

4. Decreased glucose tolerance occurs with high glucose values in the following conditions:
   a. Diabetes mellitus
   b. Postgastrectomy
   c. Hyperthyroidism
   d. Excess glucose ingestion
   e. Hyperlipidemia types III, IV, and V
   f. Hemochromatosis
   g. Cushing's disease (steroid effect)
   h. CNS lesions
   i. Pheochromocytoma

5. Decreased glucose tolerance with hypoglycemia can be found in persons with von Gierke's disease, severe liver damage, or increased epinephrine levels.

6. Increased glucose tolerance with flat curve (ie, glucose does not increase, but may decrease to hypoglycemic levels) occurs in the following conditions:
   a. Pancreatic islet cell hyperplasia or tumor
   b. Poor intestinal absorption caused by diseases such as sprue, celiac disease, or Whipple's disease
   c. Hypoparathyroidism
   d. Addison's disease
   e. Liver disease
   f. Hypopituitarism, hypothyroidism

Interfering Factors

1. Smoking increases glucose levels.
2. Altered diets (eg, weight reduction) before testing can diminish carbohydrate tolerance and suggest "false
diabetes.”

3. Glucose levels normally tend to increase with aging.

4. Prolonged oral contraceptive use causes significantly higher glucose levels in the second hour or in later blood specimens.

5. Infectious diseases, illnesses, and operative procedures affect glucose tolerance. Two weeks of recovery should be allowed before performing the test.

6. Certain drugs impair glucose tolerance levels (this list is not all inclusive; see Appendix J for other drugs):
   a. Insulin
   b. Oral hypoglycemics
   c. Large doses of salicylates, anti-inflammatories
   d. Thiazide diuretics
   e. Oral contraceptives
   f. Corticosteroids
   g. Estrogens
   h. Heparin
   i. Nicotinic acid
   j. Phenothiazines
   k. Lithium
   l. Metyrapone (Metopirone) If possible, these drugs should be discontinued for at least 3 days before testing. Check with clinician for specific orders.

7. Prolonged bed rest influences glucose tolerance results. If possible, the patient should be ambulatory. A GTT in a hospitalized patient has limited value.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. A written reminder may be helpful.
   a. A diet high in carbohydrates (150 g) should be eaten for 3 days preceding the test. Instruct the patient to abstain from alcohol.
   b. The patient should fast for at least 12 hours but not more than 16 hours before the test. Only water may be ingested during fasting time and test time. Use of tobacco products is not permitted during testing.
   c. Patients should rest or walk quietly during the test period. They may feel weak, faint, or nauseated during the test. Vigorous exercise alters glucose values and should be avoided during testing.

2. Collect blood specimens at the prescribed times and record exact times collected. Urine glucose testing is no longer recommended.

3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have the patient resume normal diet and activities at the end of the test. Encourage eating complex carbohydrates and protein if permitted.

2. Administer prescribed insulin or oral hypoglycemics when the test is done. Arrange for the patient to eat within a short time (30 minutes) after these medications are taken.

3. Interpret test results and counsel appropriately. Patients newly diagnosed with diabetes will need diet, medication, and lifestyle modification instructions.

4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

1. GTT is contraindicated in patients with a recent history of surgery, MI, or labor and delivery; these conditions can produce invalid values.

2. If fasting glucose is >140 mg/dL (>7.8 mmol/L) on two separate occasions, or if the 2-hour postprandial blood glucose is >200 mg/dL (>11.1 mmol/L) on two separate occasions, GTT is not necessary for a diagnosis of diabetes mellitus to be established.

3. The GTT is of limited diagnostic value for children.

4. The GTT should be postponed if the patient becomes ill, even with common illnesses such as the flu or a severe cold.

5. Record and report any reactions during the test. Weakness, faintness, and sweating may occur between the second and third hours of the test. If this occurs, a blood sample for a glucose level should be drawn immediately and the GTT aborted.

6. Should the patient vomit the glucose solution, the test is declared invalid; it can be repeated in 3 days (72 hours).

Glycosylated Hemoglobin (Hb A1c); Glycohemoglobin (G-Hb); Glycated Hemoglobin (GhB); Diabetic Control Index; Glycated Serum Protein (GSP), Fructosamine

Glycohemoglobin is a normal, minor type of hemoglobin. Glycosylated hemoglobin is formed at a rate proportional to the average glucose concentration by a slow, nonenzymatic process within the red blood cells (RBCs) during their 120-day circulating life span. Glycohemoglobin is blood glucose bound to hemoglobin. In the presence of hyperglycemia, an increase in glycohemoglobin causes an increase in Hb A1c. If the glucose concentration increases because of insulin deficiency, then glycosylation is irreversible.

Glycosylated hemoglobin values reflect average blood sugar levels for the 2- to 3-month period before the test. This test provides information for evaluating diabetic treatment modalities (every 3 months), is useful in determining treatment for
juvenile-onset diabetes with acute ketoacidosis, and tracks control of blood glucose in milder cases of diabetes. It can be a valuable adjunct in determining which therapeutic choices and directions (eg, oral antihypoglycemic agents, insulin, β-cell transplantations) will be most effective. A blood sample can be drawn at any time. The measurement is of particular value for specific groups of patients: diabetic children, diabetic patients in whom the renal threshold for glucose is abnormal, unstable type 1 diabetic patients (taking insulin) in whom blood sugar levels vary markedly from day to day, type 2 diabetic patients who become pregnant, and persons who, before their scheduled appointments, change their usual habits, dietary or otherwise, so that their metabolic control appears better than it actually is.

Reference Values

**Normal**
Results are expressed as percentage of total hemoglobin. Values vary slightly by method and laboratory. G-Hb: 4.0%–7.0% Hb A<sub>1c</sub>: 4.0%–6.7% of total hemoglobin H

**Procedure**
1. Obtain a 5-mL venous blood sample with EDTA purple-topped anticoagulant additive. Serum may not be used.
2. Observe standard precautions. Place specimen in a biohazard bag.

**Clinical Implications**
1. Values are frequently increased in persons with poorly controlled or newly diagnosed diabetes.
2. With optimal control, the Hb A<sub>1c</sub> moves toward normal levels.
3. A diabetic patient who recently comes under good control may still show higher concentrations of glycosylated hemoglobin. This level declines gradually over several months as nearly normal glycosylated hemoglobin replaces older RBCs with higher concentrations.
4. Increases in glycosylated hemoglobin occur in the following conditions:
   a. Iron-deficiency anemia
   b. Splenectomy
   c. Alcohol toxicity
   d. Lead toxicity
5. Decreases in glycosylated hemoglobin occur in the following conditions:
   a. Hemolytic anemia
   b. Chronic blood loss
   c. Pregnancy
   d. Chronic renal failure

**Interfering Factors**
1. Presence of Hb F and H causes falsely elevated values.

**Interventions**

**Pretest Patient Care**
1. Explain test purpose and blood-drawing procedure. Observe standard precautions. Fasting is not required.
2. Note that this test is not meant for short-term diabetes mellitus management; instead, it assesses the efficacy of long-term management modalities over several weeks or months. It is not useful more often than 4–6 weeks.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
1. Interpret test outcome and counsel patient appropriately for management of diabetes. If test results are not consistent with clinical findings, check the patient for Hb F, which elevates Hb A<sub>1c</sub> results.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Clinical Alert**

A number of different tests can determine glycosylated hemoglobin levels. The most specific of these measures is Hb A<sub>1c</sub>. There are different expected values for each test. Keep in mind that Hb A<sub>1</sub> is always 2% to 4% higher than Hb A<sub>1c</sub>. When interpreting results, be certain of the specific test used.

**Critical Value**
1. GHB: >10.1%
2. A<sub>1c</sub>: >8.1% (corresponds with glucose >200 mg/dL or >11.1 mmol/L)

**Lactose Tolerance; Breath Hydrogen Test**

Lactose intolerance often begins in infancy, with symptoms of diarrhea, vomiting, failure to thrive, and malabsorption. The patient becomes asymptomatic when lactose is removed from the diet. This syndrome is caused by a deficiency of sugar-splitting enzymes (lactase) in the intestinal tract.

This test is actually a GTT done to diagnose intestinal disaccharidase (lactase) deficiency. Glucose is measured, and it is the increase or lack of increase over the fasting specimen that is used for the interpretation. Breath samples reveal increased hydrogen levels, which are caused by lactose buildup in the intestinal tract. Colonic bacteria metabolize the
lactose and produce hydrogen gas.

Reference Values
Normal Change in glucose from normal value of >30 mg/dL or >1.7 mmol/L Inconclusive: 20–30 mg/dL or 1.1–1.7 mmol/L Abnormal: <20 mg/dL or <1.1 mmol/L Hydrogen (breath): <10 ppm increase from baseline is abnormal

Procedure
1. Follow instruction given for the GTT.
2. Draw a blood specimen from a fasting patient. The patient then drinks 50 g of lactose mixed with 200 mL of water (2 g lactose/kg body weight).
3. Draw blood lactose samples at 0, 30-, 60-, and 90-minute intervals.
4. Take hydrogen breath samples at the same time intervals as the blood specimens. Contact your laboratory for collection procedures.

Clinical Implications
1. Lactose intolerance occurs as follows:
   a. A “flat” lactose tolerance finding (ie, no rise in glucose) points to a deficiency of sugar-splitting enzymes, as in irritable bowel syndrome. This type of deficiency is more prevalent in American Indians, African Americans, Asians, and Jews.
   b. A monosaccharide tolerance test such as the glucose/galactose tolerance test should be done as a follow-up.
      1. The patient ingests 25 g of both glucose and galactose.
      2. A normal increase in glucose indicates a lactose deficiency.
   c. Secondary lactose deficiency found in:
      1. Infectious enteritis
      2. Bacterial overgrowth in intestines
      3. Inflammatory bowel disease, Crohn’s disease
      4. Giardia lamblia infection
      5. Cystic fibrosis of pancreas
   d. The hydrogen breath test is abnormal in the lactose deficiency test because:
      a. Malabsorption causes hydrogen (H₂) production through the process of fermentation of lactose in the colon.
      b. The H₂ formed is directly proportional to the amount of test dose lactose not digested by lactase.
   e. In diabetes:
      a. Blood glucose values may show increases of >20 mg/dL (>1.11 mmol/L) despite impaired lactose absorption.
      b. In diabetes, there may be an abnormal lactose tolerance curve due to faulty metabolism, not necessarily from lactose intolerance.

Interventions
Pretest Patient Preparation
1. Explain test purpose and procedure. The patient must fast for 8–12 hours before the test.
2. Do not allow the patient to eat dark bread, peas, beans, sugars, or high-fiber foods within 24 hours of the test.
3. Do not permit smoking during the test and for 8 hours before testing; no gum chewing.
4. Do not allow antibiotics to be taken for 2 weeks before the test unless specifically ordered.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Have the patient resume normal diet and activity.
2. Interpret test results and counsel appropriately. Patients with irritable bowel syndrome with gas, bloating, abdominal pain, constipation, and diarrhea have lactose deficiency. Restricting milk intake relieves symptoms.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

END PRODUCTS OF METABOLISM AND OTHER TESTS

Ammonia (NH₃)
Ammonia, an end product of protein metabolism, is formed by bacteria acting on intestinal proteins together with glutamine hydrolysis in the kidneys. The liver normally removes most of this ammonia via the portal vein circulation and converts the ammonia to urea. Because any appreciable level of ammonia in the blood affects the body's acid-base balance and brain function, its removal from the body is essential. The liver accomplishes this by synthesizing urea so that it can be excreted by the kidneys.

Blood ammonia levels are used to diagnose Reye's syndrome, to evaluate metabolism, and to determine the progress of severe liver disease and its response to treatment. Blood ammonia measurements are useful in monitoring patients on hyperalimentation therapy.

Reference Values
Normal Adults: 15–56 µg/dL or 9–33 µmol/L Children: 36–85 µg/dL or 21–50 µmol/L 10 days–2 years: (<2 weeks): 95–157 µg/dL or 56–92 µmol/L Birth–10 days: 109–182 µg/dL or 64–107 µmol/L Values test somewhat higher in capillary blood samples. Values can vary greatly with testing method used.

Procedure
1. Obtain a 5-mL venous plasma sample from a fasting patient. A green-topped (heparin) or purple-topped (EDTA)
tube may be used. Observe standard precautions.
2. Place the sample in an iced container. The specimen must be centrifuged at 4°C. Promptly remove plasma from cells. Perform the test within 20 minutes or freeze plasma immediately.
3. Note all antibiotics the patient is receiving; these drugs lower ammonia levels.

**Clinical Implications** Increased ammonia levels occur in the following conditions:

1. Reye's syndrome
2. Liver disease, cirrhosis
3. Hepatic coma (does not reflect degree of coma)
4. Gl hemorrhage
5. Renal disease
6. HHH syndrome: hyperornithinemia, hyperammonemia, homocitrullinuria
7. Transient hyperammonemia of newborn
8. Certain inborn errors of metabolism or urea except for argininosuccinicaciduria
9. Gl tract infection with distention and stasis
10. Total parenteral nutrition
11. Ureterosigmoidostomy

**Interfering Factors**

1. Ammonia levels vary with protein intake and many drugs.
2. Exercise may cause an increase in ammonia levels.
3. Ammonia levels may be increased by use of a tight tourniquet or by tightly clenching the fist while samples are drawn.
4. Ammonia levels can rise rapidly in the blood tubes.
5. Hemolysed blood gives falsely elevated levels.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. Instruct the patient to fast (if possible) for 8 hours before the blood test. Water is permitted.
2. Do not allow the patient to smoke for several hours before the test (raises levels).
3. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes, monitor appropriately, and begin treatment.
2. Remember that in patients with impaired liver function demonstrated by elevated ammonia levels, the blood ammonia level can be lowered by reduced protein intake and by use of antibiotics to reduce intestinal bacteria counts.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Clinical Alert**

Ammonia should be measured in all cases of unexplained lethargy and vomiting, in encephalitis, or in any neonate with unexplained neurologic deterioration.

**Bilirubin**

Bilirubin results from the breakdown of hemoglobin in the red blood cells and is a byproduct of hemolysis (ie, red blood cell destruction). It is produced by the reticuloendothelial system. Removed from the body by the liver, which excretes it into the bile; it gives the bile its major pigmentation. Usually, a small amount of bilirubin is found in the serum. A rise in serum bilirubin levels occurs when there is an excessive destruction of red blood cells or when the liver is unable to excrete the normal amounts of bilirubin produced.

There are two forms of bilirubin in the body: indirect or unconjugated bilirubin, which is protein bound, and direct or conjugated bilirubin, which circulates freely in the blood until it reaches the liver, where it is conjugated with glucuronide transferase and then excreted into the bile. An increase in protein-bound bilirubin (unconjugated bilirubin) is more frequently associated with increased destruction of red blood cells (hemolysis); an increase in free-flowing bilirubin is more likely seen in dysfunction or blockage of the liver. A routine examination measures only the total bilirubin. A normal level of total bilirubin rules out any significant impairment of the excretory function of the liver or excessive hemolysis of red cells. Only when total bilirubin levels are elevated will there be a call for differentiation of the bilirubin levels by conjugated and unconjugated types.

The measurement of bilirubin allows evaluation of liver function and hemolytic anemias. This test is not suitable for infants younger than 15 days (see Neonatal Bilirubin on page 342).

**Reference Values**

**Normal** Adults Total: 0.3–1.0 mg/dL or 5–17 µmol/L Conjugated (direct): 0.0–0.2 mg/dL or 0.0–3.4 µmol/L

**Procedure**
1. Obtain a 5-mL nonhemolyzed sample from a fasting patient. Observe standard precautions. Serum is used.
2. Protect the sample from ultraviolet light (sunlight).
3. Avoid air bubbles and unnecessary shaking of the sample during blood collection.
4. If the specimen cannot be examined immediately, store it away from light and in a refrigerator.

Clinical Implications

1. Total bilirubin elevations accompanied by jaundice may be due to hepatic, obstructive, or hemolytic causes.
   a. Hepatocellular jaundice results from injury or disease of the parenchymal cells of the liver and can be caused by the following conditions:
      1. Viral hepatitis
      2. Cirrhosis
      3. Infectious mononucleosis
      4. Reactions of certain drugs such as chlorpromazine
   b. Obstructive jaundice is usually the result of obstruction of the common bile or hepatic ducts due to stones or neoplasms. The obstruction produces high conjugated bilirubin levels due to bile regurgitation.
   c. Hemolytic jaundice is due to overproduction of bilirubin resulting from hemolytic processes that produce high levels of unconjugated bilirubin. Hemolytic jaundice can be found in the following conditions:
      1. After blood transfusions, especially those involving many units
      2. Pernicious anemia
      3. Sickle cell anemia
      4. Transfusion reactions (ABO or Rh incompatibility)
      5. Crigler-Najjar syndrome (a severe disease that results from a genetic deficiency of a hepatic enzyme needed for the conjugation of bilirubin)
      6. Erythroblastosis fetalis (see Neonatal Bilirubin, page 342)
   d. Miscellaneous diseases
      1. Dubin-Johnson syndrome
      2. Gilbert's disease (familial hyperbilirubinemia)
      3. Nelson's disease (with acute liver failure)
      4. Pulmonary embolism/infarct
      5. Congestive heart failure

2. Elevated indirect unconjugated bilirubin levels occur in the following conditions:
   a. Hemolytic anemias due to a large hematoma
   b. Trauma in the presence of a large hematoma
   c. Hemorrhagic pulmonary infarcts
   d. Crigler-Najjar syndrome (rare)
   e. Gilbert's disease (conjugated hyperbilirubinemia; rare)

3. Elevated direct conjugated bilirubin levels occur in the following conditions:
   a. Cancer of the head of the pancreas
   b. Choledocholithiasis
   c. Dubin-Johnson syndrome

Interfering Factors

1. A 1-hour exposure of the specimen to sunlight or high-intensity artificial light at room temperature will decrease the bilirubin content.
2. No contrast media should be administered 24 hours before measurement; a high-fat meal may also cause decreased bilirubin levels by interfering with the chemical reactions.
3. Air bubbles and shaking of the specimen may cause decreased bilirubin levels.
4. Certain foods (eg, carrots, yams) and drugs (see Appendix J) increase the yellow hue in the serum and can falsely increase bilirubin levels when tests are done using certain methods (eg, spectrophotometer).
5. Prolonged fasting raises the bilirubin level, as does anorexia.

Clinical Alert

Panic Value for Bilirubin in Adults

>12 mg/dL or >200 µmol/L

Interventions

Pretest Patient Care

1. Explain test purpose and procedure and relation of results to jaundice.
2. Ensure that the patient is fasting, if possible.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

NOTE

Excessive amounts of bilirubin eventually seep into the tissues, which assume a yellow hue as a result. This yellow color is a clinical sign of jaundice. In newborns, signs of jaundice may indicate hemolytic anemia or congenital icterus. Total bilirubin must be >2.5 mg/dL (>41.6 µmol/L) to detect jaundice in adults.

Posttest Patient Aftercare
1. Interpret test outcome and monitor appropriately.
2. Have patient resume normal activities.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Neonatal Bilirubin, Total and Fractionated (“Baby Bili”)**

In newborns, signs of jaundice may indicate hemolytic anemia or congenital icterus. If bilirubin levels reach a critical point in the infant, damage to the CNS may occur in a condition known as kernicterus. Therefore, in these infants, the level of bilirubin is the deciding factor in whether or not to perform an exchange transfusion. Total bilirubin must be >5.0 mg/dL to detect jaundice in newborns.

Jaundice may also be seen in babies who are breast feeding as a result of low milk intake and subsequent lack of vitamin K–dependent clotting factors. This condition usually resolves within 1 week.

Neonatal bilirubin is used to monitor erythroblastosis fetalis (hemolytic disease of the newborn), which usually causes jaundice in the first 2 days of life. All other causes of neonatal jaundice, including physiologic jaundice, hematomahemorrhage, liver disease, and biliary disease, should also be monitored. Normal, full-term neonates experience a normal, neonatal, physiologic, transient hyperbilirubinemia by the third day of life, which rapidly falls by the fifth to tenth day of life. This test cannot be used after the tenth day of life owing to the formation of endogenous carotenoids.

**Reference Values**

**Normal Newborns** (0–7 days) Total: 1.0–10.0 mg/dL or 17–170 µmol/L Conjugated (direct): 0.0–0.8 mg/dL or 0–136 µmol/L Unconjugated (indirect): 0.0–10.0 mg/dL or 0–170 µmol/L Cord blood total: Full term: <2.5 mg/dL or <43 µmol/L Premature: <2.9 mg/dL or <50 µmol/L See Table 6.4 for a comparison of premature and full-term infants.

**Table 6.4 Neonatal Total Comparison**

<table>
<thead>
<tr>
<th>Test</th>
<th>Serum Bilirubin (mg/dL)</th>
<th>SI Units (µmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;24 h</td>
<td>&lt;8.0</td>
<td>&lt;137</td>
</tr>
<tr>
<td>&lt;48 h</td>
<td>&lt;12.0</td>
<td>&lt;205</td>
</tr>
<tr>
<td>3–5 day</td>
<td>&lt;15.0</td>
<td>&lt;256</td>
</tr>
<tr>
<td>7 day</td>
<td>&lt;15.0</td>
<td>&lt;256</td>
</tr>
</tbody>
</table>

Note: Most labs are not doing conjugated and unconjugated anymore.

**Procedure**

1. Draw blood from heel of newborn using a capillary pipette and amber Microtainer tube; 0.5 mL of serum is needed. Cord blood may also be used.
2. Protect sample from light.

**Clinical Implications**

1. **Elevated total bilirubin** (neonatal) is associated with the following conditions:
   a. Erythroblastosis fetalis occurs as a result of blood incompatibility between mother and fetus.
      1. Rh (D) antibodies and other Rh factors
      2. ABO antibodies
      3. Other blood groups, including KIDD, KELL, and DUFFY (see Chapter 8)
   b. Galactosemia
   c. Sepsis
   d. Infectious diseases (eg, syphilis, toxoplasmosis, cytomegalovirus)
   e. Red blood cell enzyme abnormalities
      1. Glucose-6-phosphate dehydrogenase (G6PD) deficiency
      2. Pyruvate kinase (PK) deficiency
      3. Spherocytosis
   f. Subdural hematoma, hemangiomas
2. **Elevated unconjugated (indirect) neonatal bilirubin** is associated with the following conditions:
   a. Erythroblastosis fetalis
   b. Hypothyroidism
   c. Crigler-Najjar syndrome
   d. Obstructive jaundice
   e. Infants of diabetic mothers
3. **Elevated conjugated (direct) neonatal bilirubin** is associated with the following conditions:
   a. Biliary obstruction
   b. Neonatal hepatitis
   c. Sepsis
Clinical Alert

Panic Value for Neonatal Bilirubin

>15 mg/dL or >256 µmol/L (mental retardation can occur)

Interventions

Pretest Patient Care

1. Explain test purpose and procedure and its relation to jaundice to the mother.
2. See Chapter 1 guidelines for safe, informed, effective pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately.
2. Be aware that for slight elevations (ie, <10.0 mg/dL or <170 µmol/L), phototherapy may be initiated.
3. Monitor neonatal bilirubin levels to determine indication for exchange transfusion. Tests should be done every 12 hours in jaundiced newborns. See Table 6.5 for exchange transfusion indications.

Table 6.5 Indications for Exchange Transfusion

<table>
<thead>
<tr>
<th>Birth Weight (g)</th>
<th>Serum Bilirubin (mg/dL)</th>
<th>Serum Bilirubin (µmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;1000</td>
<td>10.0</td>
<td>170</td>
</tr>
<tr>
<td>1001–1250</td>
<td>13.0</td>
<td>222</td>
</tr>
<tr>
<td>1251–1500</td>
<td>15.0</td>
<td>256</td>
</tr>
<tr>
<td>1501–2000</td>
<td>17.0</td>
<td>291</td>
</tr>
<tr>
<td>2001–2500</td>
<td>18.0</td>
<td>309</td>
</tr>
<tr>
<td>&gt;2500</td>
<td>20.0</td>
<td>&gt;342</td>
</tr>
</tbody>
</table>

4. Transfuse at one step earlier in the presence of the following conditions:
   a. Coombs’ test positive
   b. Serum protein <5 g/dL
   c. Metabolic acidosis (pH < 7.25)
   d. Respiratory distress (with O<sub>2</sub> <50 mm Hg or 6.6 kPA)
   e. Certain clinical findings (eg, hypothermia, CNS, or other clinical deterioration; sepsis; hemolysis)

Other criteria for exchange transfusion are suddenness and rate of bilirubin increase and when such an increase occurs; for example, an increase of 3 mg/dL (51 µmol/L) in 12 hours, especially after bilirubin has already leveled off, must be followed by frequent serial determinations, especially if it occurs on the first or seventh day of life rather than on the third day. Beware of a rate of bilirubin increase of >1 mg/dL (>17 µmol/L) during the first day of life. Serum bilirubin of 10 mg/dL (170 µmol/L) after 24 hours or 15 mg/dL (256 µmol/L) after 48 hours despite phototherapy usually indicates that serum bilirubin will reach 20 mg/dL (342 µmol/L).

Blood Urea Nitrogen (BUN, Urea Nitrogen)

Urea forms in the liver and, along with CO<sub>2</sub>, constitutes the final product of protein metabolism. The amount of excreted urea varies directly with dietary protein intake, increased excretion in fever, diabetes, and increased adrenal gland activity.

The test for BUN, which measures the nitrogen portion of urea, is used as an index of glomerular function in the production and excretion of urea. Rapid protein catabolism and impairment of kidney function will result in an elevated BUN level. The rate at which the BUN level rises is influenced by the degree of tissue necrosis, protein catabolism, and the rate at which the kidneys excrete the urea nitrogen. A markedly increased BUN is conclusive evidence of severe impaired glomerular function. In chronic renal disease, the BUN level correlates better with symptoms of uremia than does the serum creatinine.

Reference Values

Normal
- Adults: 6–20 mg/dL or 2.1–7.1 mmol/L
- Elderly patients (>60 years): 8–23 mg/dL or 2.9–8.2 mmol/L
- Children: 5–18 mg/dL or 1.8–6.4 mmol/L

Procedure

1. Obtain a 5-mL venous blood sample. Serum is preferred.
2. Observe standard precautions.

Clinical Implications

1. Increased BUN levels (azotemia) occur in the following conditions:
   a. Impaired renal function caused by the following conditions:
      1. Congestive heart failure
      2. Salt and water depletion
      3. Shock
      4. Stress
      5. Acute MI
b. Chronic renal disease such as glomerulonephritis and pyelonephritis  
c. Urinary tract obstruction  
d. Hemorrhage into GI tract  
e. Diabetes mellitus with ketoacidosis  
f. Excessive protein intake or protein catabolism as occurs in burns or cancer  
g. Anabolic steroid use  

2. **Decreased BUN levels** are associated with the following conditions:  
   a. Liver failure (severe liver disease), such as that resulting from hepatitis, drugs, or poisoning  
   b. Acromegaly  
   c. Malnutrition, low-protein diets  
   d. Impaired absorption (celiac disease)  
   e. Nephrotic syndrome (occasional)  
   f. Syndrome of inappropriate antidiuretic hormone (SIADH)  

**Interfering Factors**  
1. A combination of a low-protein and high-carbohydrate diet can cause a decreased BUN level.  
2. The BUN is normally lower in children and women because they have less muscle mass than adult men.  
3. Decreased BUN values normally occur in late pregnancy because of increased plasma volume (physiologic hydremia).  
4. Older persons may have an increased BUN when their kidneys are not able to concentrate urine adequately.  
5. IV feedings only may result in overhydration and decreased BUN levels.  
6. Many drugs may cause increased or decreased BUN levels.  

**Clinical Alert**  
1. If a patient is confused, disoriented, or has convulsions, the BUN level should be checked. If the level is high, it may help to explain these signs and symptoms.  
2. Panic value for BUN is >100 mg/dL (>35 mmol/L).  

**Interventions**  

**Pretest Patient Care**  
1. Explain test purpose and blood-drawing procedure. Assess dietary history.  
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.  

**Posttest Patient Aftercare**  
1. Interpret test outcome and monitor as appropriate for impaired kidney function.  
2. Be aware that in patients with an elevated BUN level, fluid and electrolyte regulation may be impaired.  
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.  

**Albumin**  
Albumin (along with total protein) is a part of a diverse microenvironment. Its primary function is the maintenance of colloidal osmotic pressure (COP) in the vascular and extravascular spaces (eg, urine, cerebrospinal fluid, and omniotic fluid). Albumin is a source of nutrition and also a part of a complex buffer system. It is a “negative” acute-phase reactant. It decreases in response to acute inflammatory infectious processes.  

Albumin is used to evaluate nutritional status, albumin loss in acute illness, liver disease and renal disease with proteinuria, hemorrhage, burns, exudates or leaks in the GI tract, and other chronic diseases. Hypoalbuminuria is an independent risk factor for older adults for mortality—admission serum albumin in geriatric patients is a predictor of outcome.  

**Reference Values**  
**Normal**  
*Children:* 2.9–5.5 g/dL or 29–55 g/L  
*Adults:* 3.5–4.8 g/dL or 35–48 g/L  
After age 40 years and in persons living in subtropics and tropics (secondary to parasitic infections), level slowly declines.  

**Procedure**  
1. Obtain 5 mL of serum in a light green tube. Fasting is not necessary.  
2. Centrifuge within 30 minutes of blood draw. Place specimen in a biohazard bag.  
3. Observe standard procedures.  
4. Urine specimens may also be collected (see Chapter 3).  

**Clinical Implications**  
1. Increased albumin is not associated with any naturally occurring condition. When albumin is increased, the only cause is decreased plasma water that increases the albumin proportionally: dehydration.  
2. Decreased albumin is associated with the following conditions:  
   a. Acute and chronic inflammation and infections  
   b. Cirrhosis, liver disease, alcoholism  
   c. Nephrotic syndrome, renal disease (increased loss in urine)  
   d. Crohn's disease, colitis  
   e. Congenital analbuminurea  
   f. Burns, severe skin disease
g. Heart failure  

h. Starvation, malnutrition, malabsorption, anorexia (decreased synthesis)  
  i. Thyroid diseases: Cushing's disease, thyrotoxicosis

Interfering Factors  Albumin is decreased in:

1. Pregnancy (last trimester, owing to increased plasma volume)  
2. Oral birth control (estrogens) and other drugs (see Appendix J)  
3. Prolonged bed rest  
4. IV fluids, rapid hydration, overhydration

Clinical Alert

Panic range: <1.5 g/dL or 15 g/L  
Levels at 2.0–2.5 g/dL or 20–25 g/L may be the cause of edema.  
Low levels occur with prolonged hospital stay.  
Lipemic specimens with a high fat content interfere.

Interventions

Pretest Patient Care

1. Explain test purpose and specimen collection procedure. No fasting is required.  
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately. Explain possible need for treatment (replacement therapy).  
2. Low levels are associated with edema. Assess patient for these signs and symptoms.  
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.  
4. Be aware that further tests may have to be done:  
   a. Total protein  
   b. Protein electrophoresis  
   c. 24-hour urine protein

Prealbumin (PAB)

In 1995, the Joint Commission on Accreditation of Healthcare Organizations (JCAHO) first issued standards that hospitals assess a patient’s nutritional status and that all patients at risk for malnutrition be identified. Visceral proteins most often used in nutrition assessment include albumin, prealbumin, C-reactive protein, and retinol-binding protein. When used in combination, they can very accurately reflect a subclinical deficit and assess response to restorative therapy.

For years albumin was the widely accepted marker for malnutrition. However, mounting evidence points to prealbumin (PAB) as the better choice. Because albumin has a half-life of 21 days, it is slow to respond to a patient’s recent increase in nutrients and, therefore, is not a good indicator of recent changes in protein levels. In contrast, prealbumin responds more rapidly and gives a timelier picture of a change in dietary status. Because of its short half-life (2 days), PAB responds quickly to a decrease in nutritional intake and nutritional restoration. It reflects the current nutritional status within a patient’s body, not the status from 3 weeks ago.

Reference Values

Normal 19–38 mg/dL (190–380 mg/L) by nephelometry

Procedure

1. Collect 7-mL blood serum sample in a red-topped tube. Observe standard precautions.  
2. Place specimen in a biohazard bag for transport to the laboratory.

Clinical Implications

1. Hospital laboratories, in conjunction with dieticians, administration, pharmacists, nurses, and physicians, may develop a clinical pathway that includes running a PAB upon admission of each surgical, ICU, and medicinal patient.  
2. Values of 0–5, 5–10, and 10–15 mg/dL (0–50, 50–100, and 100–150 mg/L) indicate severe, moderate, and mild protein depletion, respectively.

Interventions

Pretest Patient Care

1. Explain test purpose. PAB is useful in assessing nutritional status, especially in monitoring the response to nutritional support in the acutely ill patient.  
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.
Posttest Patient Aftercare

1. Interpret test outcomes and determine the need for possible follow-up testing. Hospital protocol may require patients to be retested twice a week until discharge if their PAB level is less than 18 mg/dL (<180 mg/L). Possible treatment includes replacement/restorative therapy.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Cholinesterase, Serum (Pseudocholinesterase); Cholinesterase, Red Blood Cell (Acetylcholinesterase)

The cholinesterase of serum is referred to as pseudocholinesterase to distinguish it from the true cholinesterase of the red blood cell (RBC). Both of these enzymes act on acetylcholine and other cholinesters. Alkylphosphates are potent inhibitors of both serum and RBC cholinesterase.

Patients who are homozygous for the atypical gene that controls serum cholinesterase activity have low levels of cholinesterase that are not inhibited by dibucaine. Persons with normal serum cholinesterase activity show 70% to 90% inhibition by dibucaine.

The red cell (true cholinesterase) enzyme is specific for the substrate acetylcholine.

These are two separate tests. The primary use of serum cholinesterase measurement (pseudocholinesterase) is to monitor the effect of muscle relaxants (eg, succinylcholine), which are used in surgery. Patients for whom suxamethonium anesthesia is planned should be tested using the dibucaine inhibition test for the presence of atypical cholinesterase variants that are incapable of hydrolyzing this widely used muscle relaxant.

The RBC cholinesterase test is used when poisoning by pesticides such as Parathion or Malathion is suspected. Severe insecticide poisoning causes headaches, visual distortions, nausea, vomiting, pulmonary edema, confusion, convulsions, reparatory paralysis, and coma.

Reference Values

**Normal** Serum cholinesterase: 4.9–11.9 U/mL or 4.9–11.9 (× 1.00) kU/L Dibucaine inhibition: 79%–84% RBC cholinesterase 30–40 U/g hemoglobin Values vary with substrate and method. These are two different tests. Values are low at birth and for the first 6 months of life

Procedures

1. For serum cholinesterase, obtain a 5-mL blood sample; 3 mL of serum is needed. This is stable for 1 week at 4°–25°C. Observe standard precautions.
2. For RBC cholinesterase, draw a blood sample using sodium heparin as an anticoagulant; do not use serum. Observe standard precautions. This is stable for 1 week at 4°–25°C.

Clinical Implications

1. **Decreased or no serum cholinesterase** occurs in the following conditions:
   a. Congenital inherited recessive disease. These patients are not able to hydrolyze drugs such as muscle relaxants used in surgery. These patients may have a prolonged period of apnea and may die if they are given succinylcholine.
   b. Poisoning from organic phosphate insecticides.
   c. Liver diseases, hepatitis, cirrhosis with jaundice
   d. Conditions that may have decreased blood albumin, such as malnutrition, anemia, infections, skin diseases, and acute MI
   e. Congestive heart failure
2. **Decreased RBC cholinesterase levels** occur in the following conditions:
   a. Congenital inherited recessive disease
   b. Organic phosphate poisoning
   c. Paroxysmal nocturnal hemoglobinemia
   d. Megaloblastic anemia (returns to normal with therapy)
3. **Increased serum cholinesterase** is associated with
   a. Type IV hyperlipidemia
   b. Nephrosis
   c. Obesity
   d. Diabetes
4. **Increased RBC cholinesterase** is associated with:
   a. Reticulocytosis
   b. Sickle cell anemia
   c. Hemolytic anemias
5. **Increased RBC cholinesterase** in amniotic fluid, along with elevated AFP, is presumptive evidence of open neural tube defect (not normally present in amniotic fluid)

Interventions

**Pretest Patient Care**

1. Explain test purpose and procedure
2. Draw blood for serum cholinesterase 2 days before surgery.
3. Be aware that blood should not be drawn in the recovery room; prior administration of surgical drugs and anesthesia invalidates the test results.
Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome and counsel appropriately.
2. Consider patients exhibiting <70% inhibition as an atypical cholinesterase variant, and be aware that the administration of succinylcholine or similar type drugs may pose a risk.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Clinical Alert**

1. In industrial exposure, workers should not return to work until cholinesterase values rise to at least 75% of normal. Red blood cell cholinesterase regenerates at the rate of 1% per day. Plasma cholinesterase regenerates at the rate of 25% in 7 to 10 days and returns to baseline in 4 to 6 weeks.
2. Cholinesterase activity is completely and irreversibly inhibited by organophosphate pesticides.

**Creatinine**

Creatinine is a byproduct in the breakdown of muscle creatine phosphate resulting from energy metabolism. It is produced at a constant rate depending on the muscle mass of the person and is removed from the body by the kidneys. Production of creatinine is constant as long as muscle mass remains constant. A disorder of kidney function reduces excretion of creatinine, resulting in increased blood creatinine levels. Thus, creatinine levels give an approximation of the glomerular filtration rate.

This test diagnoses impaired renal function. It is a more specific and sensitive indicator of kidney disease than BUN, although in chronic renal disease, both BUN and creatinine are ordered to evaluate renal problems because the BUN/creatinine ratio provides more information.

**Reference Values**

**Normal**  
Adult men: 0.9–1.3 mg/dL or 80–115 µmol/L  
Adult women: 0.6–1.1 mg/dL or 53–97 µmol/L  
Children (3–18 years): 0.5–1.0 mg/dL or 44–88 µmol/L  
Young children (0–3 years): 0.3–0.7 mg/dL or 27–62 µmol/L  
BUN/creatinine ratio: 10:1 to 20:1

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum is preferred, but heparinized blood can be used. Place specimen in a biohazard bag.
2. Observe standard precautions.

**Clinical Implications**

1. **Increased blood creatinine levels** occur in the following conditions:  
   a. Impaired renal function  
   b. Chronic nephritis  
   c. Obstruction of urinary tract  
   d. Muscle disease  
      1. Gigantism  
      2. Acromegaly  
      3. Myasthenia gravis  
      4. Muscular dystrophy  
      5. Poliomyelitis  
   e. Congestive heart failure  
   f. Shock  
   g. Dehydration  
   h. Rhabdomyolysis  
   i. Hyperthyroidism
2. **Decreased creatinine levels** occur in the following conditions:  
   a. Small stature  
   b. Decreased muscle mass  
   c. Advanced and severe liver disease  
   d. Inadequate dietary protein  
   e. Pregnancy (0.4–0.6 mg/dL or 36–53 µmol/L is normal; >0.8 mg/dL or >71 µmol/L is abnormal and should be noted)
3. **Increased ratio (>20:1)** with normal creatinine occurs in the following conditions:  
   a. Increased BUN (prerenal azotemia), heart failure, salt depletion, dehydration  
   b. Catabolic states with tissue breakdown  
   c. GI hemorrhage  
   d. Impaired renal function plus excess protein intake, production, or tissue breakdown
4. **Increased ratio (>20:1)** with elevated creatinine occurs in the following conditions:  
   a. Obstruction of urinary tract  
   b. Prerenal azotemia with renal disease
5. **Decreased ratio (<10:1)** with decreased BUN occurs in the following conditions:  
   a. Acute tubular necrosis  
   b. Decreased urea synthesis as in severe liver disease or starvation  
   c. Repeated dialysis
d. SIADH
e. Pregnancy
6. Decreased ratio (<10:1) with increased creatinine occurs in the following conditions:
   a. Phenacetin therapy (accelerates conversion of creatine to creatinine)
   b. Rhabdomyolysis (releases muscle creatinine)
   c. Muscular patients who develop renal failure

Interfering Factors
1. High levels of ascorbic acid and cephalosporin antibiotics can cause a falsely increased creatinine level; these agents also interfere with BUN/creatinine ratio.
2. Drugs that influence kidney function plus other medications can cause a change in the blood creatinine level (see Appendix J).
3. A diet high in meat can cause increased creatinine levels.
4. Creatinine is falsely decreased by bilirubin, glucose, histidine, and quinidine compounds.
5. Ketoacidosis may increase serum creatinine substantially.

Clinical Alert
1. Panic value is 10 mg/dL or 890 µmol/L in nondialysis patients.
2. Creatinine level should always be checked before administering nephrotoxic chemotherapeutics such as methotrexate, cisplatin, cyclophosphamide, mithramycin, and semustine.

Interventions

Pretest Patient Care
1. Explain test purpose and procedure.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Care
1. Interpret test results and monitor as appropriate for impaired renal function.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Cystatin C

Cystatin C is a low-molecular-weight protein inhibitor found in blood serum and is an indicator of glomerular filtration in kidney function.

This test is done to assess glomerular filtration rate (GFR) in the elderly. Cystatin C may be a more reliable indicator of renal function in the elderly than is the creatinine level. GFR and kidney size decline with age and thus creatinine levels may be unreliable as an indicator of GFR.

Reference Values
Normal
Young adults: <0.70 mg/mL (<2.9 µmol/mL) Elderly adults: <0.85 mg/mL (<3.5 µmol/mL)

Procedure
1. No fasting is required.
2. Obtain a venous blood sample.

Clinical Implications Cystatin C levels abnormally increase in association with impaired renal function and loss of kidney homeostasis, as in acute renal failure, chronic renal failure, diabetic nephropathy, and infections.

Interventions

Pretest Patient Care
1. Explain purpose and sampling procedure for cystatin C.
2. Assess for signs of abnormal kidney function (hypertension, pain, edema, uremia, disorders of urination, and urine composition). Some conditions have no symptoms of nephrotic syndrome.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Care
1. Interpret outcomes and provide the patient with support and counseling.
2. Explain follow-up testing and possible treatment for kidney disease.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Uric Acid

Uric acid is formed from the breakdown of nucleic acids and is an end product of purine metabolism. Uric acid is transported by the plasma from the liver to the kidney, where it is filtered and where about 70% is excreted. The remainder of uric acid is excreted into the GI tract and degraded. A lack of the enzyme uricase allows this poorly soluble substance to accumulate in body fluids.
The basis for this test is that an overproduction of uric acids occurs when there is excessive cell breakdown and catabolism of nucleonic acids (as in gout), excessive production and destruction of cells (as in leukemia), or an inability to excrete the substance produced (as in renal failure). Measurement of uric acid is used most commonly in the evaluation of renal failure, gout, and leukemia. In hospitalized patients, renal failure is the most common cause of elevated uric acid levels, and gout is the least common cause.

Reference Values

**Normal**
- Men: 3.4–7.0 mg/dL or 202–416 µmol/L
- Women: 2.4–6.0 mg/dL or 143–357 µmol/L
- Children: 2.0–5.5 mg/dL or 119–327 µmol/L

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum is preferred; heparinized blood is acceptable. Place specimen in a biohazard bag.
2. Observe standard precautions.

**Clinical Implications**

1. *Elevated uric acid levels (hyperuricemia)* occur in the following conditions:
   - a. Gout (the amount of increase is not directly related to the severity of the disease)
   - b. Renal diseases and renal failure, prerenal azotemia
   - c. Alcoholism (ethanol consumption)
   - d. Down syndrome
   - e. Lead poisoning
   - f. Leukemia, multiple myeloma, lymphoma
   - g. Lesch-Nyhan syndrome (hereditary gout)
   - h. Starvation, weight-loss diets
   - i. Metabolic acidosis, diabetic ketoacidosis
   - j. Toxemia of pregnancy (serial determination to follow therapy)
   - k. Liver disease
   - l. Hyperlipidemia, obesity
   - m. Hypoparathyroidism, hypothyroidism
   - n. Hemolytic anemia, sickle cell anemia
   - o. Following excessive cell destruction, as in chemotherapy and radiation treatment (acute elevation sometimes follows treatment)
   - p. Psoriasis
   - q. Xanthinuria (deficiency of xanthine oxidase)

2. *Decreased levels of uric acid* occur in the following conditions:
   - a. Fanconi’s syndrome
   - b. Wilson's disease
   - c. SIADH
   - d. Some malignancies (eg, Hodgkin's disease, multiple myeloma)
   - e. Xanthinuria (deficiency of xanthine oxidase)

**Interfering Factors**

1. Stress and strenuous exercise will falsely elevate uric acid.
2. Many drugs cause increase or decrease of uric acid (see Appendix J).
3. Purine-rich diet (eg, liver, kidney, sweetbreads) increases uric acid levels.
4. High levels of aspirin decrease uric acid levels.
5. Low purine intake, coffee, and tea decrease uric acid levels.

**Interventions**

**Pretest Patient Care**

1. Advise patient of test purpose and blood-drawing procedure; fasting is preferred.
2. Promote relaxation; avoid strenuous exercise.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. Interpret test results and monitor appropriately for renal failure, gout, or leukemia. Uric acid level should fall in patients who are treated with uricosuric drugs such as allopurinol, probenecid, and sulfinpyrazone.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Clinical Alert**

1. Monitor uric acid levels during treatment of leukemia.
2. Acute, dangerous levels may occur following administration of cytotoxic drugs.

**Lead (Pb)**

Lead is absorbed into the body through both the respiratory and GI tracts. It also moves transplacentally to the fetus.
Absorption through these different routes varies and is affected by age, nutritional status, particle size, and chemical form of the lead. Absorption is inversely proportional to particle size; this factor makes lead-bearing dust important. Adults absorb 6% to 10% of dietary lead and retain very little of it; however, children from birth to 2 years of age have been shown to absorb 40% to 50% and to retain 20% to 25% of dietary lead. Spontaneous excretion of lead in urine by infants and young toddlers is normally about 1 µg/kg/24 hours, which may increase somewhat in cases of acute poisoning. Dietary intake of lead is <1 µg/kg of lead, which provides a margin of safety in the sense that a child goes into positive lead balance when intake exceeds 5 µg/kg of body weight. Early symptoms of lead poisoning include anorexia, apathy or irritability, fatigue, and anemia. Toxic effects include GI distress, joint pain, colic, headache, stupor, convulsions, and coma. Another test that may be used to evaluate lead intoxication is free erythrocyte protoporphyrin. However, a blood lead assay is the definitive test.

The blood lead assay is used to screen adults and children for lead poisoning (plumbism). In adults, high levels are caused mainly by industrial exposure from lead-based paints, gasoline, and ceramics. High-risk children usually are aged 3 to 12 years and live in or visit old or dilapidated housing with lead-based paint. A single paint chip can contain as much as 10,000 µg of lead.

Reference Values

**Normal** 0–10 µg/dL or 0–0.48 µmol/L

**Procedure**

1. Obtain a sample by finger stick using lead-free heparinized capillary tubes or venous blood drawn in a 3-mL trace element–free tube. Place specimen in a lead-free biohazard bag or container.
2. Do not separate plasma from cells. Refrigerate the sample.
3. Observe standard precautions.

**Clinical Implications** Blood lead levels in adults:

1. <10 µg/dL or <0.48 µmol/L: normal without occupational exposure
2. <20 µg/dL or <0.97 µmol/L: acceptable with occupational exposure
3. >20 µg/dL or >1.9 µmol/L: report to state occupational agency
4. >60 µg/dL or >2.9 µmol/L: remove from occupational exposure and begin chelation therapy

Table 6.6 lists the U.S. Centers for Disease Control and Prevention (CDC) classifications for levels of blood lead. See Table 6.7 for the effects of blood lead in children.

<table>
<thead>
<tr>
<th>Class Blood Lead *</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>I  &lt;10 µg/dL or 0.48 µmol/L</td>
<td>Not lead poisoned</td>
</tr>
<tr>
<td>IIA 10–14 µg/dL or 0.48–0.68 µmol/L</td>
<td>Rescreen frequently and consider prevention activities</td>
</tr>
<tr>
<td>IIB 15–19 µg/dL or 0.72–0.92 µmol/L</td>
<td>Institute nutritional and educational interventions</td>
</tr>
<tr>
<td>III 20–44 µg/dL or 0.97–2.1 µmol/L</td>
<td>Evaluate environment and consider chelation therapy</td>
</tr>
<tr>
<td>IV 45–69 µg/dL or 2.17–3.33 µmol/L</td>
<td>Institute environmental intervention and chelation therapy</td>
</tr>
<tr>
<td>V &gt;69 µg/dL or 3.33 µmol/L</td>
<td>Medical emergency</td>
</tr>
</tbody>
</table>

*Owing to possible contamination during collection, elevated levels should be confirmed with a second specimen before therapy is instituted.

Table 6.7 Effects of Increased Blood Lead Levels on Children

<table>
<thead>
<tr>
<th>Blood Lead Level</th>
<th>Effects in Children</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;10 µg/dL or &gt;0.48 µmol/L</td>
<td>Reduced IQ, hearing, and growth</td>
</tr>
<tr>
<td>&gt;20 µg/dL or &gt;0.97 µmol/L</td>
<td>Impaired nerve function</td>
</tr>
<tr>
<td>&gt;30 µg/dL or &gt;1.45 µmol/L</td>
<td>Reduced vitamin D metabolism</td>
</tr>
<tr>
<td>&gt;40 µg/dL or &gt;1.93 µmol/L</td>
<td>Damage to blood-forming system</td>
</tr>
<tr>
<td>&gt;50 µg/dL or &gt;2.41 µmol/L</td>
<td>Severe stomach cramps</td>
</tr>
<tr>
<td>&gt;60 µg/dL or &gt;2.90 µmol/L</td>
<td>Severe anemia</td>
</tr>
<tr>
<td>&gt;80 µg/dL or &gt;3.86 µmol/L</td>
<td>Severe brain damage</td>
</tr>
<tr>
<td>&gt;125 µg/dL or &gt;6.04 µmol/L</td>
<td>Death</td>
</tr>
</tbody>
</table>


**Interfering Factors**

1. Failing to use lead-free Vacutainer tubes invalidates results.
2. An elevated level should be confirmed with a new second specimen to ensure that the specimen was not contaminated.
Clinical Alert

1. Critical values:
   a. <15 years of age, >20 µg/dL or >0.97 µmol/L; ≥15 years of age, >30 µg/dL or >1.45 µmol/L
   b. Patients with blood lead concentrations >80 µg/dL or >3.86 µmol/L (panic value) should be hospitalized immediately and treated as medical emergencies.
   c. A single lead determination cannot distinguish between chronic and acute exposure.

2. Following chelation therapy, lead levels are assessed at varying intervals, and it is not unusual to see a slight increase due to lead leeching from bones.

3. Pregnant women with blood lead levels (BLL) > 10 µg/dL or >0.48 µmol/L are at risk for delivering a child with a BLL also > 10 µg/dL or >0.48 µmol/L.

Pretest Patient Care

1. Explain test purpose and procedure.
2. Explain the importance of follow-up if lead levels are elevated.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results, counsel, and monitor appropriately for elevated lead levels. Explain chelation therapy and possible need for further testing, eg, iron deficiency and blood protoporphyrins.
   a. Parental compliance is necessary. Parent education about lead poisoning can be given face-to-face, by pamphlet distribution, or in both ways.
   b. The most important component of medical management is to facilitate reduction in the child's exposure to the environmental lead. In providing intervention for the child with an elevated blood lead level, the initial step is to obtain a detailed environmental history. The causes of childhood lead poisoning are multiple and must take into account potential environmental hazards as well as characteristics of the individual child. Once a child is found to have lead intoxication, all potential sources must be identified and removed from the child's environment.
   c. The recommended diet for a child with lead toxicity is simply a good diet with adequate protein and mineral intake and limitation of excess fat. It is no longer necessary to exclude canned foods and beverages when the cans are manufactured in the United States because the manufacture of cans with lead-soldered seams ended in the United States in 1991.
   d. Iron deficiency can enhance absorption and toxicity of lead and often coexists with overexposure to lead. All children with a blood lead concentration >20 µg/dL or >0.97 µmol/L whole blood should have appropriate testing for iron deficiency.
   e. In class IV lead intoxication, chelation is necessary. Chelation therapy must be done in conjunction with eliminating the source of the lead poisoning. Chelation therapy, when promptly administered, can be life-saving and can reduce the period of morbidity associated with lead toxicity.
   f. Additional follow-up tests may be ordered, including free erythrocyte protoporphyrin or erythrocyte HNC protoporphyrin.

3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Osteocalcin (Bone G1a Protein)

Osteocalcin, also referred to as bone G1a protein, is a protein produced by the osteoblasts and dentin and has a function in bone mineralization and calcium ion homeostasis. A small amount of osteocalcin, an integral part in bone formation, is released into the blood and therefore can serve as a marker for recent bone formation. Osteocalcin levels are influenced by age (rapid growth), gender (males somewhat higher), and are increased during menopause. This test is used to screen for osteoporosis in postmenopausal women, assess risk for fractures, and determine eligibility for treatment for osteoporosis. Osteocalcin is a specific marker for bone formation and is regulated by 1, 25-dehydroxy vitamin D.

Reference Values

Normal Osteocalcin: 8.1 ± 4.6 µg/L or 1.4 ± 0.8 nmol/L Carboxylated osteocalcin: 9.9 ± 0.5 µg/L or 1.7 ± 0.1 nmol/L Undercarboxylated osteocalcin: 3.7 ± 1.0 µg/L or 0.6 ± 0.2 nmol/L

Normal Using RIA Adult male: 3.0–13.0 ng/mL or 3.0–13.0 µg/L Premenopausal female: 0.4–8.2 ng/mL or 0.4–8.2 µg/L Postmenopausal female: 1.5–11.0 ng/mL or 1.5–11.0 µg/L There is a diurnal variation, a peak during the night and a decrease in the morning.

Procedure Collect a venous blood sample of serum on ice, separate within 1 hour, and immediately freeze. Avoid a freeze–thaw cycle.

Interfering Factors

1. Increased during bed rest and no increase in bone formation.
2. Increased with impaired renal function and no increase in bone formation.

Clinical Implications

1. Abnormally increased levels indicate increased bone formation in persons with hyperparathyroidism, fractures, and acromegaly.
2. Decreased levels are associated with hypoparathyroidism, a deficiency of growth hormone, and medications such as glucocorticoids, bisphosphonates, and calcitonin.

Interventions
Pretest Patient Care

1. Explain purpose and procedure of test. Record age and menopausal state. Tell patient that the risk for osteoporosis steadily increases with age. Also obtain pertinent personal and family history of osteoporotic fractures, history of falls, etc.
2. Follow Chapter 1 guideline for **safe, effective, informed pretest care**.

Posttest Patient Care

1. Interpret test outcomes and counsel regarding further tests (eg, dual-energy x-ray absorptiome [DXA] [bone density of the femoral neck] or quantitative ultrasound) and possible treatment (eg, medical: alendronate, raloxifene). Sixteen percent of postmenopausal women will be found to have lumbar spine osteoporosis. Other blood test markers of bone resorption include pyridinolines, telopeptides, acid phosphatase, and urine tests of hydroxyproline and galactosyl hydroxylysine. These markers are known as collagen cross-links.
2. See Chapter 1 for **safe, effective, informed posttest care**.

HORMONE TESTS

Androstenedione

Androstenedione is one of the major androgens produced by the ovaries in females, and to a lesser extent in the adrenal in both genders. This hormone is converted to estrogens by hepatic enzymes. Levels rise sharply after puberty and peak at age 20 years.

This hormone measurement is helpful in the evaluation of conditions characterized by hirsutism and virilization. In females, there is poor correlation of plasma levels with clinical severity.

Reference Values

| Normal | Newborns: 20–290 ng/dL or 0.7–10.1 mmol/L Prepuberty: 8–50 ng/dL or 0.3–1.7 mmol/L Women: 75–205 ng/dL or 2.6–7.2 mmol/L Men: 85–275 ng/dL or 3.0–9.6 mmol/L Postmenopausal women: <10 ng/dL or 0.35 mmol/L (abrupt decline at menopause) Different laboratories may have variation in reference values. |

Procedure

1. Obtain a 5-mL venous blood sample in the morning and place on ice. Serum or EDTA can be used. Observe standard precautions. Place specimen in a biohazard bag.
2. In women, collect this specimen 1 week before or after the menstrual period. Record date of last menstrual period on the laboratory form.

Clinical Implications

1. **Increased androstenedione values** are associated with the following conditions:
   a. Stein-Leventhal syndrome
   b. Cushing's syndrome
   c. Certain ovarian tumors (polycystic ovarian syndrome)
   d. Ectopic ACTH-producing tumor
   e. Late-onset congenital adrenal hyperplasia
   f. Ovarian stromal hyperplasia
   g. Osteoporosis in females
2. **Decreased androstenedione values** are found in the following conditions:
   a. Sickle cell anemia
   b. Adrenal and ovarian failure

<table>
<thead>
<tr>
<th>Clinical Alert</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;1000 mg/dL or &gt;34.9 mmol/L (suggests virilizing tumor)</td>
</tr>
</tbody>
</table>

Interventions

Pretest Patient Care

1. Explain purpose of test and blood-drawing procedure. Obtain pertinent history of signs and symptoms (eg, excessive hair growth and infertility).
2. Ensure that patient is fasting and that blood is drawn at peak production (7:00 a.m.). Lowest levels are at 4:00 p.m.
3. Collect specimen 1 week before menstrual period in women.
4. Follow guidelines in Chapter 1 for **safe, effective, informed pretest care**.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results and counsel appropriately for ovarian and adrenal dysfunction.
3. Follow guidelines in Chapter 1 for **safe, effective, informed posttest care**.

Aldosterone

Aldosterone is a mineralocorticoid hormone produced in the adrenal zona glomerulosa under complex control by the
The renin-angiotensin system. Its action is on the renal distal tubule, where it increases resorption of sodium and water at the expense of increased potassium excretion.

This test is useful in detecting primary or secondary aldosteronism. Patients with primary aldosteronism characteristically have hypertension, muscular pains and cramps, weakness, tetany, paralysis, and polyuria. It is also used to evaluate causes of hypertension (found in 1% of hypertension cases).

**NOTE**

A random aldosterone test is of no diagnostic value unless a plasma renin activity is done at the same time.

**Reference Values**

**Normal (In upright position)**

<table>
<thead>
<tr>
<th>Age</th>
<th>Reference Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adults</td>
<td>7–30 ng/dL or 0.19–0.83 nmol/L</td>
</tr>
<tr>
<td>Adolescents</td>
<td>4–48 ng/dL or 0.11–1.33 nmol/L</td>
</tr>
<tr>
<td>Children</td>
<td>5–80 mg/dL or 0.14–2.22 nmol/L</td>
</tr>
<tr>
<td>Low-sodium diet</td>
<td>values 3–5 times higher</td>
</tr>
</tbody>
</table>

**Procedure**

1. Take plasma with the patient in an upright position for 2 hours and with unrestricted salt intake.
2. Obtain a 5-mL venous blood specimen in a heparinized or EDTA Vacutainer tube. Serum, EDTA, or heparinized blood may be used. The cells must be separated from plasma immediately. Blood should be drawn with patient sitting. Observe standard precautions.
3. Specify and record the time of the venipuncture. Circadian rhythm exists in normal subjects, with levels of aldosterone peaking in the morning. Specify if the blood has been drawn from the adrenal vein (values are much higher: 200–800 ng/dL or 5.5–22.6 mmol/L).
4. Be aware that a 24-hour urine specimen with boric acid preservative may also be ordered. Refrigerate immediately following collection.
5. Have patient follow a normal sodium diet 2–4 weeks before test.
6. Ensure that low potassium is treated before test.

**Clinical Implications**

1. **Elevated levels of aldosterone (primary aldosteronism)** occur in the following conditions:
   a. Aldosterone-producing adenoma (Conn's disease)
   b. Adrenocortical hyperplasia (pseudoprimary aldosteronism)
   c. Indeterminate hyperaldosteronism
   d. Glucocorticoid remediable hyperaldosteronism
2. **Secondary aldosteronism**, in which aldosterone output is elevated because of external stimuli or greater activity in the renin-angiotensin system, occurs in the following conditions:
   a. Salt depletion
   b. Potassium loading
   c. Laxative abuse
   d. Cardiac failure
   e. Cirrhosis of liver with ascites
   f. Nephrotic syndrome
   g. Bartter's syndrome
   h. Diuretic abuse
   i. Hypovolemia and hemorrhage
   j. After 10 days of starvation
   k. Toxemia of pregnancy
3. **Decreased aldosterone levels** are found in the following conditions:
   a. Aldosterone deficiency
   b. Addison's disease
   c. Syndrome of renin deficiency (very rare)
   d. Low aldosterone levels associated with hypertension are found in Turner's syndrome, diabetes mellitus, and alcohol intoxication

**Interfering Factors**

1. Values are increased by upright posture.
2. Recently administered radioactive medications affect test outcomes.
3. Heparin therapy causes levels to fall. See Appendix I for drugs that increase and decrease levels.
4. Thermal stress, late pregnancy, and starvation cause levels to rise.
5. Aldosterone levels decrease with age.
6. Many drugs—diuretics, antihypertensives, progestogens, estrogens, and licorice—should be terminated 2–4 weeks before test.

**Clinical Alert**

1. The simultaneous measurement of aldosterone and renin is helpful in differentiating primary from secondary hyperaldosteronism. Renin levels are high in secondary aldosteronism and low in primary aldosteronism.
2. Potassium deficiencies should be corrected before testing for aldosterone.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedures. Assess for history of diuretic or laxative abuse. If 24-hour urine specimen is required, follow protocols in Chapter 3.
2. Discontinue diuretic agents, progestational agents, estrogens, and black licorice for 2 weeks before the test.
3. Ensure that the patient's diet for 2 weeks before the test is normal (other than the previously listed restrictions) and should include 3 g/day (135 mEq/L/day) of sodium. Check with your laboratory for special protocols.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities and diet.
2. Interpret test results and monitor appropriately for aldosteronism and aldosterone deficiency.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Antidiuretic Hormone (ADH); Arginine Vasopressin Hormone**

ADH is excreted by the posterior pituitary gland. When ADH activity is present, small volumes of concentrated urine are excreted. When ADH is absent, large amounts of diluted urine are produced. Higher secretion occurs at night, with erect posture, and with pain, stress, or exercise. Measurement of the level of ADH is useful in the differential diagnosis of polyuric and hyponatremic states. ADH testing aids in diagnosis of urine concentration disorders, especially diabetes insipidus, SIADH, psychogenic water intoxication, and syndromes of ectopic ADH production.

**Reference Values**

*Normal* <2.5 pg/mL or <2.3 pmol/L

**Procedure**

1. Draw venous blood samples, 5 mL, into prechilled tubes and put on ice. Plasma with EDTA anticoagulant is needed. Observe standard precautions. Place specimen in a biohazard bag.
2. Ensure that patient is in a sitting position and calm during blood collection.

**Clinical Implications**

1. Increased secretion of ADH is associated with the following conditions:
   a. SIADH (with respect to plasma osmolality)
   b. Ectopic ADH production (systemic neoplasm)
   c. Nephrogenic diabetes insipidus
   d. Acute intermittent porphyria
   e. Guillain-Barré syndrome
   f. Brain tumor, diseases, injury, neurosurgery
   g. Pulmonary diseases (tuberculosis)
2. Decreased secretion of ADH occurs in the following conditions:
   a. Central diabetes insipidus (hypothalamic or neurogenic)
   b. Psychogenic polydipsia (water intoxication)
   c. Nephrotic syndrome

**Interfering Factors**

1. Recently administered radioisotopes cause spurious results.
2. Many drugs affect results (eg, thiazide diuretics, oral hypoglycemics, and narcotics); see Appendix J.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. Encourage relaxation before and during blood-drawing procedure.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Resume normal activities.
2. Interpret test results and counsel appropriately for urine concentration disorders and polyuria.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

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**Clinical Alert**

To distinguish SIADH from other conditions that cause dilutional hyponatremia, other tests must be done, such as plasma osmolality, plasma sodium, and water-loading test.

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**Atrial Natriuretic Factor (ANF), ANP and BNP**

Atrial natriuretic factor (ANF) is a hormone secreted by the cardiac atria during acute and chronic cardiac volume and pressure overload. The discovery of ANF indicates that the heart is an endocrine gland and confirms speculation that there is a mechanism in or near the heart that regulates body fluid hemostasis. This hormone enhances salt and water excretion, blocks aldosterone and renal secretion, and inhibits the action of angiotensin II and vasopressin.

Recently, a hormone produced by the ventricles of the heart, brain natriuretic peptide or B-type natriuretic peptide (BNP),
has been shown to increase in response to ventricular volume expansion and pressure overload. BNP is a marker of ventricular systolic and diastolic dysfunction. This test is useful in diagnosing congestive heart failure. It is not useful for diagnosing other heart conditions. Chart 6.1 describes types of heart failures; Chart 6.2 offers a scale for grading them. Figure 6.3 illustrates the relationship of BNP to heart disease.

### Chart 6.1 Heart Failure

<table>
<thead>
<tr>
<th>Type of Heart Failure</th>
<th>Signs and Symptoms</th>
<th>Tests to Diagnose</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Left heart failure</strong></td>
<td>Shortness of breath at rest and exercise</td>
<td>History/physical exam</td>
</tr>
<tr>
<td>(congestive heart failure)</td>
<td>Persistent cough</td>
<td>Electrocardiogram</td>
</tr>
<tr>
<td></td>
<td>Weakness or fatigue</td>
<td>Chest x-ray</td>
</tr>
<tr>
<td></td>
<td>Edema in feet, ankles, legs</td>
<td>Blood tests: brain natriuretic peptide, atrial natriuretic factor</td>
</tr>
<tr>
<td></td>
<td>Weight gain</td>
<td>Pulmonary function tests</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Cardiac ultrasound</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Treadmill stress test</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Thallium stress test</td>
</tr>
</tbody>
</table>

**Right heart failure**  
An increase in left ventricular filling pressure that is reflected back in the pulmonary circulation

**High-output failure**  
Inability of the heart to supply the body with blood-borne nutrients despite adequate blood volume and normal myocardial contractility

### Chart 6.2 Grading Heart Diseases

- **Class I**—no limitation of physical activity; no fatigue, shortness of breath, or heart palitations with ordinary activities
- **Class II**—slight limitation in physical activity; with fatigue, shortness of breath, or heart palitations during ordinary activities
- **Class III**—marked limitation of physical activity; with fatigue, shortness of breath, or heart palitations with less-than-ordinary physical activity
- **Class IV**—severe to complete limitation of physical activity with fatigue, shortness of breath, or heart palitations with any exertion; symptoms occur even at rest

**Footnote**

Source: The New York Heart Association, 2001

**Figure 6.3** Relationship of BNP to heart disease (classification of the New York Heart Association) (Source: Biosite Diagnostics, San Diego, CA, USA)

**Reference Values**

- **Normal** Atrial natriuretic factor (ANF): 20–77 pg/mL or 20–77 ng/L  
  B-type natriuretic peptide (BNP): <100 pg/mL or <100 ng/L

**Procedure**

1. Obtain a plasma sample by venipuncture from a fasting patient. Use a lavender-topped KEDTA tube. If a nonfasting sample is obtained, notify laboratory.
2. Prechill the tube at 4°C before drawing sample. After drawing sample, chill tube in wet ice for 10 minutes. Place
specimen in a biohazard bag.

**Clinical Implications** Increased ANF levels occur in:

1. Congestive heart failure
2. Cardiovascular disease with elevated filling pressure
3. A symptomatic cardiac volume overload
4. Paroxysmal atrial tachycardia

**Interfering Factors** See Appendix J for drugs that affect test outcomes.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and need to fast. Assess for signs and symptoms indicating need for testing (e.g., chronic fatigue, cough, heart palpitations, high blood pressure).
2. Withhold cardiovascular medications per physician's order (e.g., β and calcium antagonists, cardiac glycosides, diuretics, vasodilators) before drawing specimen.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Medications and usual diet may be restarted per physician's order.
2. Evaluate patient outcomes and monitor appropriately for congestive heart failure.
3. In collaboration with physician, explain need for possible follow-up tests and medication therapy.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Cortisol (Hydrocortisone)**

Cortisol (hydrocortisone/compound F) is a glucocorticosteroid of the adrenal cortex and affects metabolism of proteins, carbohydrates, and lipids. Cortisol stimulates glucogenesis by the liver, inhibits the effect of insulin, and decreases the rate of glucose use by the cells. In health, the secretion rate of cortisol is higher in the early morning (6:00–8:00 a.m.) and lower in the evening (4:00–6:00 p.m.). This variation is lost in patients with Cushing's syndrome and in persons under stress.

The cortisol test evaluates adrenal hormone function. Cortisol is elevated in adrenal hyperfunction and decreased in adrenal hypofunction. Suppression and stimulation tests may also be done. Cortisol (dexamethasone) suppression test screens for Cushing's syndrome and identifies depressed persons who are likely to respond to antidepressants or electroshock therapy. It is based on the fact that ACTH production is suppressed in healthy persons after a low dose of dexamethasone but not in persons with Cushing's syndrome or in some depressed persons.

**Reference Values**

**Normal** Cortisol: 8:00 a.m.: 5–23 µg/dL or 138–635 nmol/L; 4:00 p.m.: 3–16 µg/dL or 83–441 nmol/L; Midnight: <50% of 8:00 a.m. level

- Newborns: 2–11 µg/dL or 55–304 nmol/L
- Maternal (at birth): 51.2–57.4 µg/dL or 1413–1584 nmol/L

After first week of life, cortisol levels attain adult values. **Suppression** 8:00 a.m. following administration of dexamethasone: <5 µg/dL (a.m. value) or <138 nmol/L.

**Stimulation** Baseline: at least 5 µg/dL or 138 nmol/L; After Cortrosyn administration: rise of at least 10 µg/dL or 276 nmol/L

**Procedure**

1. Obtain 5-mL venous blood samples at 8:00 a.m. and at 4:00 p.m. Serum is preferred. Heparin anticoagulant may be used. Place specimen in a biohazard bag.
2. Observe standard precautions.

**Clinical Implications**

1. Decreased cortisol levels are found in the following conditions:
   a. Adrenal hyperplasia
   b. Addison's disease
   c. Anterior pituitary hyposecretion (pituitary destruction)
   d. Hypothyroidism (hypopituitarism)
2. Increased cortisol levels are found in the following conditions:
   a. Hyperthyroidism
   b. Stress (trauma, surgery)
   c. Carcinoma (extreme elevation in the morning and no variation later in the day)
   d. Cushing's syndrome (high on rising but no variation later in the day)
   e. Overproduction of ACTH due to tumors (oat cell cancers)
   f. Adrenal adenoma
   g. Obesity

**Interfering Factors**

1. Pregnancy will cause an increased value.
2. There is no normal diurnal variation in patients under stress.
3. Drugs such as spironolactone and oral contraceptives will give falsely elevated values (see Appendix J).
4. Decreased levels occur in persons taking dexamethasone, prednisone, or prednisolone (steroids) (see Appendix J).
5. Random cortisol tests are useless and provide no pertinent information.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and blood-drawing procedure. Blood must be drawn at 8:00 a.m. and 4:00 p.m.
2. Encourage relaxation.
3. Ensure that no radioisotopes are administered within 1 day before the test.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. Interpret test results and counsel appropriately for adrenal dysfunction.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Cortisol Suppression (Dexamethasone Suppression; DST)**

See foregoing cortisol test for purpose and indications. The DST test helps to differentiate causes of elevated cortisol. Cortisol <15 µg/dL (<41.4 nmol/L) is indication of adrenal cortisol insufficiency.

**Reference Values**

*Normal*: <5 µg/dL (<138 nmol/L) or <50% of baseline (8:00 a.m. specimen)

**Procedure**

1. Obtain a 5-mL venous blood the day following administration of dexamethasone. Serum or heparinized plasma is acceptable. Observe standard precautions. Place specimen in a biohazard bag.
2. Administer late evening or bedtime; dexamethasone tablets by mouth. There is a low-dose and high-dose suppression test in which either 1.0 mg or 8.0 mg of dexamethasone is given, respectively, at 11:00 p.m. The following morning at 8:00 a.m., a blood sample is drawn to measure cortisol. (Some Cushing's disease patients have false-positive results with this low dose.)

**Clinical Implications**

1. Suppression occurs in persons with:
   a. Cushing's syndrome (>10 µg/dL or >276 nmol/L)
   b. Endogenous depression (50% of cases)
2. No suppression occurs in:
   a. Adrenal adenoma, carcinoma
   b. Ectopic ACTH-producing tumors

**Interfering Factors** False suppression can occur in the following conditions:

1. Pregnancy
2. High doses of estrogens
3. Alcoholism
4. Uncontrolled diabetes
5. Trauma, high stress, fever, dehydration
6. Phenytoin (Dilantin) (see Appendix J for other drugs)

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. Fasting is required for the 8:00 a.m. test.
2. Discontinue all medications for 24 to 48 hours before the study. Especially important are spironolactone, estrogens, birth control pills, cortisol, tetracycline, stilbestrol, and phenytoin. Check with the physician.
3. Weigh the patient and record weight.
4. Have baseline blood cortisol drawn at 8:00 a.m. and 4:00 p.m. Give 1 mg dexamethasone at 11:00 p.m. the same day. Draw blood at 8:00 a.m. the next morning.
5. Ensure that no radioisotopes are administered within 1 week before test.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. Interpret test results and counsel appropriately for Cushing's syndrome or depression.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Cortisone Stimulation (Cosyntropin, Cortrosyn Stimulation); Adrenocorticotropic Hormone (ACTH) Stimulation**

This detects adrenal insufficiency after Cortrosyn administration. Cortrosyn is a synthetic subunit of ACTH that exhibits the full corticosteroid-stimulating effect of ACTH in healthy persons. Failure to respond is an indication of adrenal insufficiency. See foregoing cortisol tests for values. This screening test is less time consuming and can be done on an
Reference Values

**Normal** Cortisol: >20 µg/dL (>552 nmol/L) rise after Cortrosyn administration

**Procedure**

1. Obtain a 4-mL fasting venous blood sample at 8:00 a.m. Observe standard precautions.
2. Administer Cortrosyn intramuscularly or intravenously as prescribed.
3. Obtain additional 4-mL blood specimens 30 and 60 minutes after administration of Cortrosyn. Serum or heparinized blood is acceptable.

**Clinical Implications**

1. Absent or blunted response to cortisol stimulation occurs in the following conditions:
   a. Addison's disease (adrenal insufficiency)
   b. Hypopituitarism (secondary adrenal insufficiency)
   c. Adrenal carcinoma, adenoma
2. Response to cortisol stimulation: adrenal hyperplasia

**Interfering Factors**

1. Prolonged steroid administration
2. Estrogens (see Appendix J)

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. Fasting during test is required. Blood specimens are obtained before and after IM injection of Cortrosyn.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. Interpret test results and monitor appropriately for adrenal insufficiency.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Clinical Alert**

In adrenal hyperplasia, there is an increase of cortisol levels of 3 to 5 times the normal; in adrenal carcinoma, there is no increase.

**Gastrin**

Gastrin, a hormone secreted by the antral G cells in stomach mucosa, stimulates gastric acid production and affects antral motility and secretion of pepsin and intrinsic factor. Gastrin values follow a circadian rhythm and fluctuate physiologically in relation to meals. The lowest values are between 3:00 a.m. and 7:00 a.m.

Measurement of serum gastrin is generally used to diagnose stomach disorders such as gastrinoma and Zollinger-Ellison syndrome in the presence of hyperacidity. (Gastric hyperacidity must be documented.)

**Reference Values**

**Normal** Adults: <25–100 pg/mL or <12–48 pmol/L Children: 10–125 pg/mL or 5–60 pmol/L Postprandial: 95–140 pg/mL or 46–67 pmol/L

**Procedure**

1. Obtain a 5-mL venous blood sample from a fasting patient. Serum is required.
2. Freeze if not tested immediately. If not fasting, this must be noted because values are different. Place specimen in a biohazard bag.
3. Observe standard precautions.

**Clinical Implications**

1. Increased gastrin levels are found in the following conditions:
   a. Stomach carcinoma (reduction of gastric acid secretion)
   b. Gastric and duodenal ulcers
   c. Zollinger-Ellison syndrome (>500 pg/mL or >240 pmol/L)
   d. Pernicious anemia
   e. Gastric carcinoma
   f. End-stage renal disease (gastrin metabolized by the kidneys)
   g. Antral G-cell hyperplasia
   h. Vagotomy without gastric resection
   i. Hyperparathyroidism
j. Pyloric obstruction
2. Decreased gastrin levels occur in the following conditions:
   a. Antrectomy with vagotomy
   b. Hypothyroidism

Interfering Factors Values will be falsely increased in nonfasting patients, elderly patients, and diabetic patients taking insulin, as well as in postgastroscopy patients and those taking H₂ secretion blockers (cimetidine), steroids, and calcium. A protein meal can elevate gastrin markedly.

Interventions
Pretest Patient Care
1. Explain test purpose and procedure.
2. Remind patient that fasting is required for 12 hours preceding the test. Water is permitted; no coffee. No radioisotopes for 1 week.
3. Note if specimen is drawn postprandial. (If after eating, note what was eaten.)
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Have patient resume normal activities.
2. Interpret test results and monitor appropriately. Follow-up testing using gastric stimulation or gastrin suppression may be indicated.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Growth Hormone (hGH); Somatotropin

Human growth hormone (somatotropin, hGH) is essential to the growth process and has an important role in the metabolism of adults. It is secreted by the pituitary gland in response to exercise, deep sleep, hypoglycemia, glucagon, insulin, and vasopressin. It also stimulates the production of RNA, mobilizes fatty acids from fat deposits, and is intimately connected with insulinism. If the pituitary gland secretes too little or too much hGH in the growth phase of life, dwarfism or gigantism will result, respectively. An excess of growth hormone during adulthood leads to acromegaly. This test confirms hypopituitarism or hyperpituitarism so that therapy can be initiated as soon as possible. Challenge or stimulation tests are generally used to detect hGH deficiency and are more informative. Much controversy surrounds the use of growth hormone stimulation tests, and the diagnosis should be considered in the context of the clinical picture.

Reference Values

Normal Men: <5 ng/mL or <226 pmol/L Women: <10 ng/mL or <452 pmol/L Children: 0–20 ng/mL or 0–904 pmol/L Newborns: 5–40 ng/mL or 226–1808 pmol/L Stimulation test (using arginine, glucagon or insulin): >5 ng/mL or >226 pmol/L (rise from baseline) >10 ng/mL or >452 pmol/L peak response from baseline Suppression test (using 100 g glucose): 0–2 ng/mL or 0–90 pmol/L or undetectable

NOTE
Because of marked fluctuations in hGH, a random specimen has limited value. Stimulation or inhibitor tests provide more information.

Procedure
1. Obtain a 5-mL venous blood sample from a fasting patient. Serum is best to use. Observe standard precautions. Place specimen in a biohazard bag.
2. Check with your laboratory for specific challenge protocols for stimulation tests such as insulin-induced hypoglycemia, arginine transfusion, glucagon infusion, L-dopa, and propranolol with exercise.

Clinical Implications
1. Increased hGH levels are associated with the following conditions:
   a. Pituitary gigantism
   b. Acromegaly
   c. Laron's dwarfism (hGH resistant)
   d. Ectopic GH secretion
   e. Uncontrolled diabetes mellitus
2. Decreased hGH levels are associated with the following conditions:
   a. Pituitary dwarfism
   b. Hypopituitarism
   c. Adrenocortical hyperfunction
3. Following stimulation testing, no response (or an inadequate response) is seen in hGH and ACTH deficiencies (hypopituitarism).
   a. Blood glucose must fall to <40 mg/dL (<2.2 mmol/L)
   b. Adrenergic signs must be observed.
4. Following suppression tests, there is no or incomplete suppression in persons with gigantism or acromegaly.
   a. Paradoxical rises in hGH may occur in patients with acromegaly.
   b. Partial suppression is sometimes seen in anorexia nervosa.
   c. In children, rebound-stimulation effect may be seen 2 to 5 hours following administration of glucose (suppression test).
Interfering Factors

1. Increased levels are associated with the use of oral contraceptives, estrogens, arginine, glucagon, levodopa, low glucose, and insulin.
2. Levels will rise to 15 times normal by the second day of starvation; levels also rise after deep sleep, stress, exercise, and anorexia.
3. Decreased levels are associated with obesity and the use of corticosteroids.
4. Many drugs interfere with test results (see Appendix J).
5. Recently administered radioisotopes interfere with test results.

Interventions

Pretest Patient Care

1. Explain test purpose and blood-drawing procedure.
2. Remind patient that fasting from food for 8 to 10 hours is required; water is permitted. For accurate levels, the patient should be free of stress and at complete rest in a quiet environment for at least 30 minutes before specimen collection.
3. Note the patient's physiologic state (eg, feeding, fasting, sleep, and/or activity) at testing in the health care record.
4. For stimulation tests, collect one tube before stimulation and at timed intervals (eg, 10, 20, 30, 45, and 60 minutes) after stimulation. For suppression tests, collect one tube before suppression and 30, 60, 90, and 120 minutes after suppression.
5. Remember that for initial testing of hGH deficiency, a vigorous exercise test is considered to be a simple, risk-free screening test, especially for children.
6. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results and monitor appropriately. A glucose challenge test may be indicated for follow-up.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Parathyroid Hormone Assay; Parathyrin; Parathormone (PTH-C-Terminal)

Parathormone (PTH), a polypeptide hormone produced in the parathyroid gland, is one of the major factors in the regulation of calcium concentration in extracellular fluid. Three molecular forms of PTH exist: intact (also called native or glandular hormone); multiple N-terminal fragments; and C-terminal fragments.

This test studies altered calcium metabolism, establishes a diagnosis of hyperparathyroidism, and distinguishes nonparathyroid from parathyroid causes of hypercalcemia. A decrease in the level of ionized calcium is the primary stimulus for PTH secretions, whereas a rise in calcium inhibits secretions. This normal relation is lost in hyperthyroidism, and PTH will be inappropriately high in relation to calcium. Acute changes in secretory activity are better reflected by the PTH, N-terminal assay. PTH and N-terminal levels are usually decreased when hypercalcemia is due to neoplastic secretions (prostaglandins). PTH and N-terminal levels may be a more reliable indication of secondary hyperparathyroidism in patients with renal failure. Creatinine level is determined concurrently with all PTH assays to determine kidney function and for meaningful interpretation of results.

Reference Values

Normal N-terminal: 8–24 pg/mL or 8–24 ng/L Intact molecule: 10–65 pg/mL or 10–65 ng/L Calcium: 8.5–10.9 mg/dL (calcium must be tested to properly interpret results) C-terminal (biomolecule): 50–330 pg/mL or 50–330 ng/L

Procedure

1. Obtain a 10-mL venous blood sample from a patient who has fasted for 10 hours. Collect the sample in chilled vials and keep on ice. Observe standard precautions. Serum or EDTA is used.
2. Immediately take specimen to the laboratory and centrifuge at 4°C after blood has clotted.

Clinical Implications

1. Increased PTH values occur with:
   a. Primary hyperparathyroidism
   b. Pseudohyperparathyroidism when there is a primary defect in renal tubular responsiveness to PTH (secondary hyperparathyroidism)
   c. Hereditary vitamin D dependency
   d. Zollinger-Ellison syndrome
   e. Spinal cord injury
2. Decreased PTH values occur in the following conditions:
   a. Hypoparathyroidism (Graves’ disease)
   b. Nonparathyroid hypercalcemia
   c. Secondary hypoparathyroidism (surgical)
   d. Magnesium deficiency
   e. Sarcoidosis
   f. Hyperthyroidism
   g. DiGeorge’s syndrome
3. Increased PTH–N-terminal values occur in the following conditions:
   a. Primary hyperparathyroidism
   b. Secondary hyperparathyroidism (more reliable than PTH-C-terminal)
4. **Decreased PTH–N-terminal values** occur in the following conditions:
   a. Hypoparathyroidism
   b. Nonparathyroidism hypercalcemia
   c. Aluminum-associated osteomalacia
   d. Severely impaired bone mineralization

5. **Increased PTH–C-terminal values** occur in the following conditions:
   a. Primary hyperparathyroidism (very specific for)
   b. Some neoplasms with elevated calcium
   c. Renal failure (even if parathyroid disease is absent)

6. **Decreased PTH–C-terminal values** occur in the following conditions:
   a. Hypoparathyroidism
   b. Nonparathyroid hypercalcemia

**Interfering Factors**

1. Elevated blood lipids and hemolysis interfere with test methods.
2. Milk-alkali syndrome may falsely lower PTH levels (Burnett’s syndrome).
3. Recently administered radioisotopes (see Appendix J) will alter results.
4. Vitamin D deficiency will decrease PTH levels.
5. Many drugs alter results; phosphates raise PTH levels up to 125%, and vitamin A and D overdoses decrease PTH levels (see Appendix J).

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. Remind patient that fasting for at least 10 hours is required. Draw blood by 8:00 a.m. because of circadian rhythm changes. Concurrently, also draw blood for testing of calcium level.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. Interpret test results and monitor appropriately for calcium imbalance and hypoparathyroidism or hyperparathyroidism.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Somatomedin C (SM-C); Insulin-like Growth Hormone**

Somatomedin C, a polypeptide hormone produced by the liver and other tissues, mediates growth hormone activity and glucose metabolism. It is carried in the blood and is bound to a protein carrier that prolongs its half-life.

This test is used to monitor the growth of children as well as to diagnose acromegaly and hypopituitarism. Normal somatomedin C results rule out a deficiency of growth hormone. Testing of somatomedin C is preferable to growth hormone tests because its levels are more constant. Somatomedin C is also a reliable nutrition index, having a low value for anorexia or malnutrition.

**Reference Values**

**Normal** See Table 6.8.

**Table 6.8 Values for Somatomedin C**

<table>
<thead>
<tr>
<th>Age (yr)</th>
<th>Male</th>
<th>Female</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ng/mL</td>
<td>nmol/L</td>
</tr>
<tr>
<td>0–5</td>
<td>0–103</td>
<td>0–13.5</td>
</tr>
<tr>
<td>6–8</td>
<td>2–118</td>
<td>0.2–15.4</td>
</tr>
<tr>
<td>9–10</td>
<td>15–148</td>
<td>2.0–19.4</td>
</tr>
<tr>
<td>11–13</td>
<td>55–216</td>
<td>7.2–28.3</td>
</tr>
<tr>
<td>14–15</td>
<td>114–232</td>
<td>14.9–30.4</td>
</tr>
<tr>
<td>16–17</td>
<td>84–221</td>
<td>11.0–28.9</td>
</tr>
<tr>
<td>18–19</td>
<td>56–177</td>
<td>7.3–23.2</td>
</tr>
<tr>
<td>20–24</td>
<td>75–142</td>
<td>9.8–18.6</td>
</tr>
<tr>
<td>25–50</td>
<td>60–122</td>
<td>7.9–16.0</td>
</tr>
</tbody>
</table>

*Note:* levels slowly decrease as person ages.

**Procedure**

1. Be aware that it is preferred that the patient be fasting. Obtain a 5-mL plasma venous blood sample using EDTA anticoagulant. Serum may also be used. Observe standard precaution. Place specimen in a biohazard bag.
2. Chill blood-drawing tubes before and place on ice immediately after obtaining specimen. Spin the sample in a
refrigerated centrifuge. Freeze if not testing immediately.

Clinical Implications

1. *Increased somatomedin C levels* are associated with the following conditions:
   a. Acromegaly (some cases), gigantism
   b. Hypoglycemia associated with non–islet cell tumors
   c. Hepatoma
   d. Wilms' tumor
   e. Precocious puberty

2. *Decreased somatomedin C levels* are associated with the following conditions:
   a. Dwarfism (short stature)
   b. Hypopituitarism
   c. Hypothyroidism
   d. Puberty delay
   e. Laron's dwarfism
   f. Cirrhosis of liver and other hepatocellular diseases
   g. Malnutrition and anorexia
   h. Diabetes mellitus (diabetic retinopathy)
   i. Emotional deprivation syndrome (maternal deprivation)

Interfering Factors

1. Somatomedin C levels are increased 2 to 3 times in pregnancy.
2. Somatomedin C levels are decreased in the following conditions:
   a. Acute illness
   b. Normal aging

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. Fasting is not required.
2. Do not administer radioisotopes within 1 week of testing.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results and monitor appropriately for abnormal growth and development.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

NOTE

Because SM-C is decreased with malnutrition, it can be used to monitor therapy for food deprivation.

FERTILITY TESTS

Fertility denotes the ability of a man and woman to reproduce; conversely, infertility denotes the lack of fertility—an involuntary reduction in the ability to produce children. When a couple has been engaging in regular, unprotected sexual intercourse for at least 1 year without conceiving, the couple is considered infertile. In about one third of cases, a male factor is the predominant cause; in another one third, the female factor predominates; and in another one third, no cause is found in either partner.

The workup for infertility starts with a complete history and physical exam for both the woman and the man, including their sexual history. A rational approach is to put each partner through a series of tests that generally uncover a vast majority of the contributing factors of infertility. These tests usually take 2 to 3 months to complete.

Standard pretest and posttest care for couples undergoing fertility testing includes the following: Provide information and support. Be sensitive to the couple's need for privacy and confidentiality. Maintain a communication network about new procedures, tests, and treatments. Help couples deal with feelings of sadness and loss. Assist couples to deal with the effects of stress and the financial burden during the diagnostic process. Assist couples in arranging work and testing schedules with the least amount of disruption for the couple. Arrange for counseling with experts who understand the different ways infertility affects someone's life.

Tests include evaluation of amenorrhea, anovulation, sperm count (angiosperm, oligospermia), hormone testing, hysterosalpingogram, laparoscopy, hysteroscopy, fertiloscopy, semen analysis, postcoital test, endometrial biopsy, and chromosome karyotype to exclude Kallmann's syndrome. Hormone testing rules pregnancy in or out (eg, chorionic gonadotropin, prolactin, luteinizing hormone [LH], follicle-stimulating hormone [FSH], thyroid-stimulating hormone [TSH], postcoital test, and antisperm antibodies). Also see estrogen testing in Chapter 3.
A postcoital examination is done to assess cervical mucus and competent sperm motility. A specimen is obtained from the endocervical canal within 2 to 12 hours of coitus and is examined for viscosity (stretching to 6 cm is normal) and for ferning effect of estrogen. The presence of >50% sperm confirms male competence.

**Chorionic Gonadotropin; Human Chorionic Gonadotropin (hCG) ß Subunit; Pregnancy Test**

The glycoprotein hormones hCG, luteinizing hormone (LH), follicle-stimulating hormone (FSH), and thyroid-stimulating (TSH) are composed of two different subunits. A subunit is similar in all of the glycoprotein hormones, and the ß subunit is unique to each hormone. Highly specific assays allow hCG to be measured in the presence of other glycoprotein hormones. The increased sensitivity of the ß-hCG test detects pregnancy as early as 6 to 10 days after implantation of the oocyte. A variety of poorly differentiated or undifferentiated neoplasms may produce ectopic chorionic gonadotropin. Assay for total hCG, both a and ß subunits, or ß-hCG may detect ectopic tumors (eg, choriocarcinoma, hydatidiform mole, germinal testicular tumors). In these neoplasms, hCG is usually the product of syncytiotrophoblastic cells.

This qualitative test detects normal pregnancy. It is quicker but less sensitive (sensitivity, 20–50 mIU/mL) than the quantitative test. This test can be expected to become positive within 3 days of implantation (ie, just after the first missed menstrual period). Cross-reactivity with LH is low, and false-positive results are rare. Occasionally, a patient with very high LH levels will give a borderline reaction. The qualitative test is usually done using urine.

The quantitative ß-hCG test is used for nonroutine detection of hCG. It is sensitive to 1 to 3 mIU/mL. This test provides the most sensitive and specific test for the detection of early pregnancy, estimation of gestational age, and diagnosis of ectopic pregnancy or threatened spontaneous abortion. This test is also useful in the workup and management of testicular tumors. High levels may be found in choriocarcinoma, embryonal cell carcinoma, and ectopic pregnancy. hCG levels are extremely useful in following germ cell neoplasms that produce hCG, especially trophoblastic neoplasms. There is little cross-reactivity with LH.

**Reference Values**

**Normal Qualitative** (for routine pregnancy tests): urine or serum negative (not pregnant)

**Quantitative** (for nonroutine detection of hCG)

- **Men:** <5.0 IU/L or mIU/mL
- **Nonpregnant women:** <5.0 IU/L or mIU/mL
- **Pregnant women:**
  - 1 week of gestation: 5–50 mIU/mL or IU/L
  - 2 weeks of gestation: 50–500 mIU/mL or IU/L
  - 3 weeks of gestation: 100–10,000 mIU/mL or IU/L
  - 4 weeks of gestation: 1080–30,000 mIU/mL or IU/L
  - 6–8 weeks of gestation: 3500–115,000 mIU/mL or IU/L
  - 12 weeks of gestation: 12,000–270,000 mIU/mL or IU/L
  - 13–16 weeks of gestation: up to 200,000 mIU/mL or IU/L
  - 17–40 weeks of gestation: gradual fall to 4000 mIU/mL or IU/L

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum is used for the test.
2. Observe standard precautions. Place specimen in a biohazard bag.
3. Urine may be used for the qualitative test. First morning specimen is recommended.

**Clinical Implications**

1. **Increased hCG values** occur in the following conditions:
   - a. Pregnancy
   - b. Successful therapeutic insemination and in vitro fertilization
   - c. Hydatidiform mole
   - d. Choriocarcinoma
   - e. Seminoma
   - f. Ovarian and testicular teratomas
   - g. Ectopic pregnancy
   - h. Certain neoplasms of the lung, stomach, and pancreas
   - i. Down syndrome (trisomy 21), mid-trimester elevation

2. **Decreased hCG values** occur in:
   - a. Threatened spontaneous abortion
   - b. Ectopic pregnancy
   - c. Trisomy 18, decrease at mid-trimester

**Interfering Factors**

1. Lipemia, hemolysis, and radioisotopes administered within 1 week of testing may affect results.
2. Test results can be positive up to 1 week after complete abortion.
3. False-negative and false-positive results can be caused by many drugs (see Appendix J).

**Clinical Alert**

Because there is great variability in hCG concentration among pregnant women, a single test determination cannot be used to accurately date the gestational age. Serial determinations may be helpful when abnormal pregnancy is suspected. Serial values do not double every 48 hours. In normal pregnancy, the hCG level doubles every 48 hours during the first 6 weeks of gestation.

**Interventions**

**Pretest Patient Care**
1. Explain test purpose and procedure.
2. Determine and record date of last menstrual period in women.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Resume normal activities.
2. Interpret test results and counsel appropriately for pregnancy or gestational problems.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Follicle-Stimulating Hormone (FSH); Luteinizing Hormone (LH)**

FSH and LH are glycoprotein pituitary hormones produced and stored in the anterior pituitary. They are under complex regulation by hypothalamic gonadotropin-releasing hormone and by gonadal sex hormones (estrogen and progesterone in females and testosterone in males). FSH acts on granulosa cells of the ovary and Sertoli’s cells of the testis, and LH acts on Leydig's (interstitial) cells of the gonads. Normally, FSH increases occur at earlier stages of puberty, 2 to 4 years before LH reaches comparable levels. In males, FSH and LH are necessary for spermatozoa development and maturation. In females, follicular formation in the early stages of the menstrual cycle is stimulated by FSH; then the midcycle surge of LH causes ovulation of the FSH-ripened ovarian follicles to occur.

This test measures the gonadotropic hormones FSH and LH and may help determine whether a gonadal deficiency is of primary origin or is due to insufficient stimulation by the pituitary hormones.

Evaluation of FSH supports other studies related to determining causes of hypothyroidism in women and endocrine dysfunction in men. In primary ovarian failure or testicular failure, FSH levels are increased. Measuring the levels of FSH and LH is of value in studying children with endocrine problems related to precocious puberty.

In the case of anovulatory fertility problems, the presence or absence of the midcycle peak can be established through a series of daily blood specimens.

**Reference Values**

*Normal* See Table 6.9.

**Table 6.9 Values for Luteinizing and Follicle-Stimulating Hormones**

<table>
<thead>
<tr>
<th></th>
<th>Luteinizing Hormone (LH)</th>
<th>Follicle-Stimulating Hormone (FSH)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(mIU/L)</td>
<td>or</td>
</tr>
<tr>
<td><strong>Female</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Follicular</td>
<td>1.37–9.9</td>
<td></td>
</tr>
<tr>
<td>Luteal</td>
<td>1.09–9.2</td>
<td></td>
</tr>
<tr>
<td>Postmenopausal</td>
<td>19.3–100.6</td>
<td></td>
</tr>
<tr>
<td><strong>Male</strong></td>
<td>1.42–15.4</td>
<td></td>
</tr>
</tbody>
</table>

Note: contact your laboratory for reference values in infants and children. Normal values may vary with method of testing and units used.

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum is needed for the test. Place specimen in a biohazard bag.
2. In women, record the date of last menstrual period.
3. Remember that it is important to measure both FSH and LH.

**Clinical Alert**

Sometimes multiple blood specimens are necessary because of episodic releases of FSH from the pituitary gland. An isolated sample may not indicate the actual activity; therefore, pooled blood specimens or multiple single blood specimens may be required.

**Clinical Implications**

1. Decreased FSH levels occur in the following conditions:
   a. Feminizing and masculinizing ovarian tumors when FSH production is inhibited because of increased estrogen secretion.
   b. Failure of hypothalamus to function properly (Kallmann’s syndrome)
   c. Pituitary LH or FSH deficiency
   d. Neoplasm of testes or adrenal glands that influence secretion of estrogens or androgens
   e. Polycystic ovarian disease
   f. Hemochromatosis
   g. Anorexia
2. Decreased FSH and LH occur in pituitary or hypothalamic failure.

3. Increased FSH levels occur in the following conditions:
   a. Turner's syndrome (ovarian dysgenesis); about 50% of patients with primary amenorrhea have Turner's syndrome.
   b. Hypopituitarism
   c. Sheehan's syndrome
   d. Precocious puberty, either idiopathic or secondary to a CNS lesion
   e. Klinefelter's syndrome
   f. Castration
   g. Alcoholism
   h. Menopause and menstrual disorders

4. Increased FSH levels occur in the following conditions:
   a. Turner's syndrome (ovarian dysgenesis); about 50% of patients with primary amenorrhea have Turner's syndrome.
   b. Hypopituitarism
   c. Sheehan's syndrome
   d. Precocious puberty, either idiopathic or secondary to a CNS lesion
   e. Klinefelter's syndrome
   f. Castration
   g. Alcoholism
   h. Menopause and menstrual disorders

5. Both FSH and LH are increased in the following conditions:
   a. Hypogonadism
   b. Complete testicular feminization syndrome
   c. Gonadal failure
   d. Congenital absence of testicle or testicles (anorchia)
   e. Menopause

6. Elevated basal LH with an LH/FSH ratio >2 and some increase of ovarian androgen in an essentially nonovulatory adult woman is presumptive evidence of Stein-Leventhal syndrome (polycystic ovary syndrome).

Interfering Factors
1. Recently administered radioisotopes
2. Hemolysis of blood sample
3. Estrogens or oral contraceptives, testosterone
4. Several drugs affect test outcomes; see Appendix J.
5. Pregnancy

Interventions

Pretest Patient Care
1. Instruct the patient regarding test purpose and procedure.
2. For women, record date of last menstrual period.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcomes and counsel appropriately.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Prolactin (hPRL)

Prolactin is a pituitary hormone essential for initiating and maintaining lactation. The gender difference in prolactin does not occur until puberty, when increased estrogen production results in higher prolactin levels in females. Circadian changes in prolactin concentration in adults are marked by episodic fluctuation and a sleep-induced peak in the early morning hours.

This test may be helpful in the diagnosis, management, and follow-up of a prolactin-secreting tumor accompanied by secondary amenorrhea or galactorrhea, hyperprolactinemia, and infertility. It is also useful in the management of hypothalamic disease and in monitoring the effectiveness of surgery, chemotherapy, and radiation treatment of prolactin-secreting tumors.

Reference Values

Normal
Nonpregnant women: 0–23 ng/mL or 0–23 µg/L
Pregnant women: 34–386 ng/mL or 34–386 µg/L by third trimester
Men: 0–20 ng/mL or 0–20 µg/L
Children: 3.2–20 ng/mL or 3.2–20 µg/L

Procedure
1. Ensure that the patient fasts for 12 hours before testing. Obtain a 5-mL venous blood sample. Serum is used.
2. Procure specimens in the morning, between 8:00 and 10:00 a.m. Draw in chilled tubes keep specimen on ice.
3. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications

1. Increased prolactin values are associated with the following conditions:
   a. Galactorrhea or amenorrhea
   b. Diseases of the hypothalamus and pituitary (acromegaly)
   c. Prolactin-secreting pituitary tumors
   d. Chiari-Frommel syndrome
   e. Ectopic production of prolactin from tumors, carcinoma, and leukemia
   f. Hypothyroidism (primary)
   g. Polycystic ovary syndrome
   h. Anorexia nervosa
   i. Insulin-induced hypoglycemia
   j. Adrenal insufficiency

2. Decreased prolactin values are found in the following conditions:
a. Sheehan's syndrome (pituitary apoplexy)
b. Idiopathic hypogonadotropic hypogonadism

NOTE
The only result of prolactin deficiency in pregnancy is the absence of postpartum lactation.

Interfering Factors
1. Increased values are associated with newborns, pregnancy, postpartum period, stress, exercise, sleep, nipple stimulation, and lactation (breast feeding).
2. Drugs (eg, estrogens, methyldopa, phenothiazines, opiates) may increase values. See Appendix J for other drugs.
3. Dopaminergic drugs inhibit prolactin secretion. Administration of L-dopa can normalize prolactin levels in galactorrhea, hyperprolactinemia, and pituitary tumor. See Appendix J for other drugs.
4. Increased levels are found in cocaine abuse, even after withdrawal from cocaine.

Clinical Alert
Levels >200 ng/mL or >200 µg/L in a nonlactating female indicate a prolactin-secreting tumor; however, a normal prolactin level does not rule out pituitary tumor.

Interventions

Pretest Patient Care
1. Explain test purpose. Fasting is required. Obtain blood specimen between 8:00 and 10:00 a.m. (3–4 hours after patient has awakened). Obtain history of leakage from the breast in nonpregnant females.
2. Have patient avoid stress, excitement, or stimulation; venipuncture itself can sometimes elevate prolactin levels.
3. If possible, discontinue all prescribed medications for 2 weeks before test.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Have patient resume normal activities.
2. Interpret test outcome and counsel regarding repeat testing to monitor treatment. Magnetic resonance imaging may be indicated.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Progesterone

Progesterone, a female sex hormone, is primarily involved in the preparation of the uterus for pregnancy and its maintenance during pregnancy. The placenta begins producing progesterone at 12 weeks of gestation. Progesterone level peaks in the midluteal phase of the menstrual cycle. In nonpregnant women, progesterone is produced by the corpus luteum. Progesterone is the single best test to determine whether ovulation has occurred.

This test is part of a fertility study to confirm ovulation, evaluate corpus luteum function, and assess risk for early spontaneous abortion. Testing of several samples during the cycle is necessary. Ovarian production of progesterone is low during the follicular (first) phase of the menstrual cycle. After ovulation, progesterone levels rise for 4 to 5 days and then fall. During pregnancy, there is a gradual increase from week 9 to week 32 of gestation, often to 100 times the level in the nonpregnant woman. Levels of progesterone in twin pregnancy are higher than in a single pregnancy. Serum progesterone levels used with ß-hCG assist in differentiating normal uterine pregnancy from abnormal uterine or ectopic pregnancy.

Reference Values

Normal
Men: <1.0 ng/mL or <3.2 nmol/L
Women: Prepubertal: 0.1–0.3 ng/mL or 0.3–1.0 nmol/L
Follicular: 0.1–0.7 ng/mL or 0.5–2.3 nmol/L
Luteal: 2–25 ng/mL or 6.4–79.5 nmol/L
First trimester: 10–44 ng/mL or 32.6–140 nmol/L
Second trimester: 19.5–82.5 ng/mL or 62.0–262 nmol/L
Third trimester: 65–290 ng/mL or 206.7–728 nmol/L

Procedure
1. Obtain a venous blood sample. Serum is needed for test. Observe standard precautions. Place specimen in a biohazard bag.
2. Remember that the test request should include gender, day of last menstrual period, and length of gestation in women.
3. Be aware that a ß-hCG may be ordered at the same time.
4. Remember that urine levels may be done, but serum is preferred.

Clinical Implications

1. Increased progesterone levels are associated with the following conditions:
   a. Congenital adrenal hyperplasia
   b. Lipid ovarian tumor
   c. Molar pregnancy
   d. Chorionepithelioma of ovary
2. Decreased progesterone levels are associated with the following conditions:
   a. Threatened spontaneous abortion
   b. Galactorrhea-amenorrhea syndrome (primary or secondary hypogonadism)
c. Short luteal phase syndrome

Interfering Factors

1. See Appendix J for drugs that affect test outcomes.
2. Critical value: levels <10 ng/mL or <32 nmol/L are associated with abnormal pregnancy outcome.
3. 5–10 ng/mL or 16–32 nmol/L: pathologic pregnancy
4. Progesterone <5 ng/mL or <16 nmol/L: nonviable

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. Note date of last menstrual period/length of gestation.
2. Do not administer radioisotopes within 1 week before the test.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results, and counsel and monitor appropriately regarding fertility and pregnancy outcome.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Testosterone, Total and Free

Testosterone is responsible for the development of male secondary sexual characteristics. It is secreted by the adrenal glands and testes in men and by the adrenal glands and ovaries in women. Excessive production induces premature puberty in men and masculinity in women. Testosterone exists in serum as both unbound (free) fractions and bound fractions to albumin: sex hormone–binding globulin (SHBG) and testosterone-binding globulin. Unbound (free) testosterone is the active portion. Testosterone levels undergo large and rapid fluctuations; levels peak in early morning in males. Females show a cyclic elevation 1 to 2 days midcycle.

Testosterone measurements in men assess hypogonadism, pituitary gonadotropin function, impotency, and cryptorchidism; these measurements are also useful in the detection of ovarian tumors and hirsutism in women. In prepubertal boys, they can assess special precocity. This test may be part of a fertility workup in association with chronic anovulation caused by polycystic ovary syndrome.

Reference Values

**Normal**
- **Total testosterone**
  - Men: 270–1070 ng/dL or 9–38 nmol/L (values in elderly men diminish moderately)
  - Women: 15–70 ng/dL or 0.52–2.4 nmol/L
  - Pregnant women: 3–4 times normal
  - Postmenopausal women: 8–35 ng/dL or 0.3–1.2 nmol/L (half of normal)
  - Children: boys: 2–20 ng/dL or 0.07–0.7 nmol/L (depends on age, sex, and onset of puberty)

- **Free testosterone**
  - Men: 50–210 pg/mL or 174–729 pmol/L
  - Women: 1.0–8.5 pg/mL or 3.5–29.5 pmol/L
  - Children: boys: 0.1–3.2 pg/mL or 0.3–11.1 pmol/L
  - Girls: 0.1–0.9 pg/mL or 0.3–3.1 pmol/L
  - Puberty: boys: 1.4–156 pg/mL or 4.9–541 pmol/L
  - Girls: 1.0–5.2 pg/mL or 3.5–18.0 pmol/L

**Procedure**

1. Obtain a 5-mL venous blood sample; serum is preferred. Observe standard precautions. Place specimen in a biohazard bag.
2. Indicate age and gender on laboratory requisition.

Clinical Implications

1. **Males:** decreased total testosterone levels occur in the following conditions:
   a. Hypogonadism (pituitary failure)
   b. Klinefelter’s syndrome
   c. Hypopituitarism (primary and secondary)
   d. Orchidectomy
   e. Hepatic cirrhosis
   f. Down syndrome
   g. Delayed puberty
2. **Males:** decreased free testosterone levels occur in hypogonadism and elderly men.
3. **Males:** increased total testosterone levels occur in the following conditions:
   a. Hyperthyroidism
   b. Syndromes of androgen resistance
   c. Adrenal tumors
   d. Precocious puberty and adrenal hyperplasia in boys
4. **Females:** increased total testosterone levels are associated with the following conditions:
   a. Adrenal neoplasms
   b. Ovarian tumors, benign or malignant (virilizing)
   c. Trophoblastic disease during pregnancy
   d. Idiopathic hirsutism
   e. Hilar cell tumor
5. **Females:** increased free testosterone levels are associated with the following conditions:
   a. Female hirsutism
   b. Polycystic ovaries
c. Virilization

Clinical Alert

1. Testosterone levels are normal in cryptorchidism, azoospermia, and oligospermia.
2. In general, there appears to be little advantage in doing urine testosterone measurements compared with (or in addition to) serum measurements; the serum test is recommended.
3. Panic value: total testosterone >200 ng/dL or >694 pmol/L in females indicates androgenic tumors of the adrenal or ovaries, especially with severe hirsutism.

Interfering Factors

1. Alcoholism in males decreases testosterone levels.
2. Estrogen therapy increases testosterone levels (see Appendix J).
3. Many drugs, including androgens and steroids, decrease testosterone levels (see Appendix J).

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. Draw blood at 7:00 a.m. for highest levels.
2. Draw multiple pooled samples at different times throughout the day if necessary for more reliable results.
3. Do not administer radioisotopes within 1 week before test.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results and counsel appropriately regarding hormone dysfunction.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

ENZYME TESTS

Acid Phosphatase; Prostatic Acid Phosphatase (PAP)

Acid phosphatases are enzymes that are widely distributed in tissues, including the bone, liver, spleen, kidney, red blood cells, and platelets. However, their greatest diagnostic importance involves the prostate gland, where acid phosphatase activity is 100 times higher than in other tissues. Immunochemical methods are highly specific for determining the prostatic fraction; however, because PAP is not elevated in early prostatic disease, this test is not recommended for screening.

This test monitors the effectiveness of treatment of cancer of the prostate. Elevated levels of acid phosphatase are seen when prostate cancer has metastasized beyond the capsule to the other parts of the body, especially the bone. Once the carcinoma has spread, the prostate starts to release acid phosphatase, resulting in an increased blood level. The prostatic fraction procedure specifically measures the concentration of prostatic acid phosphatase secreted by cells of the prostate gland. Acid phosphatase is also present in high concentration in seminal fluid. Tests for presence of this enzyme on vaginal swabs may be used to investigate rape.

Reference Values

Normal 2.5–3.7 ng/mL or 2.5–3.7 µg/L

Procedure

1. Obtain a 5-mL venous blood sample. Serum may be used, if test is done within 1 hour. EDTA plasma is preferred to stabilize acid phosphatase.
2. Remember that morning is recommended because diurnal variation exists.
3. Place specimen in a biohazard bag, transport to lab immediately, and place on ice.

Clinical Implications

1. A significantly elevated acid phosphatase value is almost always indicative of metastatic cancer of the prostate. If the tumor is successfully treated, this enzyme level will drop within 3 to 4 days after surgery or 3 to 4 weeks after estrogen administration.
2. Moderately elevated values also occur in the absence of prostate carcinoma in the following conditions:
   a. Niemann-Pick disease
   b. Gaucher's disease
   c. Prostatitis (benign prostatic hypertrophy)
   d. Urinary retention
   e. Any cancer that has metastasized to the bone
   f. Myelocytic leukemia

Interfering Factors

1. Various drugs may cause increased and decreased PAP levels.
2. Palpation of the prostate gland and prostate biopsy before testing causes increases in PAP levels.
3. Transurethral resection of the prostate (TURP) and bladder catheterization cause increased levels.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. No palpation of or procedures on the prostate gland and no rectal examinations should be performed 2 to 3 days before test.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results and counsel appropriately regarding repeat testing. When elevated values are present, retesting and biopsy are considered.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Prostate-Specific Antigen (PSA)

Prostate-specific antigen (PSA) is functionally and immunologically distinct from prostatic acid phosphatase. PSA is localized in both normal prostatic epithelial cells and prostatic carcinoma cells. PSA has proved to be the most prognostically reliable marker for monitoring recurrence of prostatic carcinoma; however, this test does not have the sensitivity or specificity to be considered an ideal tumor marker. PSA detects incidental as well as aggressive carcinomas.

The most useful approach to date may be age-specific PSA reference ranges, which are based on the concept that blood PSA concentration is dependent on patient age. The increase in PSA with advancing age is attributed to four major factors: prostate enlargement, increasing inflammation, presence of microscopic but clinically insignificant cancer, and leakage of PSA into the serum (Table 6.10).

<table>
<thead>
<tr>
<th>Age (yr)</th>
<th>PSA Range (ng/mL)</th>
<th>PSA Range (µg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>40–49</td>
<td>0.0–2.5</td>
<td>0.0–2.5</td>
</tr>
<tr>
<td>50–59</td>
<td>0.0–3.5</td>
<td>0.0–3.5</td>
</tr>
<tr>
<td>60–69</td>
<td>0.0–4.5</td>
<td>0.0–4.5</td>
</tr>
<tr>
<td>70–79</td>
<td>0.0–6.5</td>
<td>0.0–6.5</td>
</tr>
</tbody>
</table>


Testing for both PSA and PAP increases detection of early prostate cancer. PSA testing determines the effectiveness of therapy for prostate cancer and is used as an early indicator of prostate cancer recurrence. The greatest value of PSA is as a marker in the follow-up of patients at high risk for disease progression.

PSA lacks sensitivity and specificity to be used alone as a screening test for prostatic carcinoma, but in conjunction with a digital rectal exam, the detection rate of prostatic carcinoma is greatly increased.

Reference Values

Normal Men: 0–4.0 ng/mL or 0–4.0 µg/L

Procedure

1. Obtain a 5-mL venous blood sample. Serum is needed.
2. Observe standard precautions. Place specimen in a biohazard bag.
3. Record patient's age.

Clinical Implications

1. PSA increases occur in prostate cancer (80% of patients).
2. Patients with benign prostatic hypertrophy often demonstrate values between 4.0 and 8.0 ng/mL (4.0–8.0 µg/L). Results between 4.0 and 8.0 ng/mL (4.0–8.0 µg/L) may represent benign prostatic hypertrophy or possible cancer of the prostate. Results >8.0 ng/mL or >8.0 µg/L are highly suggestive of prostatic cancer.
3. Increases to >4.0 ng/mL or >4.0 µg/L have been reported in about 8% of patients with no prostatic malignancies and no benign diseases.
4. If a prostate tumor is completely and successfully removed, no antigen will be detected.

Interfering Factors
1. Transient increases in PSA occur following prostate palpation or rectal examination.
2. Increased with urinary retention.
3. Recent exposure to radioisotopes causes test interference.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. Do not schedule any prostatic examinations, including rectal examination, prostate biopsy, or TURP, for 1 week before the blood test is performed.
3. Follow guidelines in Chapter 1 for **safe, effective, informed pretest care**.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. Interpret test results, and monitor and counsel as appropriate for response to treatment and progression or remission of prostate cancer.
3. Follow guidelines in Chapter 1 for **safe, effective, informed posttest care**.

---

**Clinical Alert**

1. PSA is not a definitive diagnostic marker to screen for carcinoma of the prostate because it is also found in men with benign prostatic hypertrophy.
2. Digital rectal examination (DRE) is recommended by the American Cancer Society as the primary test for detection of prostatic tumor. Recent studies indicate that serum PSA may offer additional information. PSA should be used in conjunction with DRE.
3. The value of prostatic cancer screening remains controversial in terms of patient morbidity and longevity outcomes.

---

**Alanine Aminotransferase (Aminotransferase, ALT); Serum Glutamic-Pyruvic Transaminase (SGPT)**

ALT is an enzyme. High concentrations occur in the liver, and relatively low concentrations are found in the heart, muscle, and kidney.

This test is primarily used to diagnose liver disease and to monitor the course of treatment for hepatitis, active postnecrotic cirrhosis, and the effects of later drug therapy. ALT is more sensitive in the detection of liver disease than in biliary obstruction. ALT also differentiates between hemolytic jaundice and jaundice due to liver disease.

**Reference Values**

**Normal** Adults (adult levels are reached by 6 months): 10–35 U/L or 0.17–0.60 µkat/L (males slightly higher) Males: 10–40 U/L or 0.17–0.68 µkat/L Females: 7–35 U/L or 0.12–0.60 µkat/L Newborns: 13–45 U/L or 0.22–0.77 µkat/L ALT values are slightly higher in males and black persons. Normal values vary with testing method. Check with your laboratory for reference values.

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum is needed for the test. Observe standard precautions. Place specimen in a biohazard bag.
2. Avoid hemolysis during collection of the specimen. (ALT activity is 6 times higher in RBCs.)

**Clinical Implications**

1. **Increased ALT levels** are found in the following conditions:
   a. Hepatocellular disease (moderate to high increase)
   b. Alcoholic cirrhosis (mild increase)
   c. Metastatic liver tumor (mild increase)
   d. Obstructive jaundice or biliary obstruction (mild increase)
   e. Viral, infectious, or toxic hepatitis (30–50 times normal)
   f. Infectious mononucleosis
   g. Pancreatitis (mild increase)
   h. Myocardial infarction, heart failure
   i. Polymyositis
   j. Severe burns
   k. Trauma to striated muscle
   l. Severe shock
2. Aspartate transaminase (AST)/ALT comparison:
   a. Although the AST level is always increased in acute MI, the ALT level does not always increase unless there is also liver damage.
   b. The ALT is usually increased more than the AST in acute extrahepatic biliary obstruction.
   c. The AST/ALT ratio is high in alcoholic liver disease; the ALT is more specific than AST for liver disease, but the AST is more sensitive to alcoholic liver disease.
Clinical Alert

Critical Value

Alcohol-acetaminophen syndrome: extremely abnormal ALT/AST values are found

>9000 U/L (>153 µkat/L): this extreme level can distinguish this syndrome from alcoholic or viral hepatitis.

Interfering Factors

1. Many drugs may cause falsely increased and decreased ALT levels (see Appendix J).
2. Salicylates may cause decreased or increased ALT levels.
3. Therapeutic heparin causes increased ALT.
4. Hemolysed blood causes increases in ALT.
5. Obesity causes increases in ALT.

Clinical Alert

There is a correlation between the presence of elevated serum ALT and abnormal antibodies to the hepatitis B virus core antigen and hepatitis C antigen. Persons with elevated ALT levels should not donate blood.

Interventions

Pretest Patient Care

1. Explain test purpose and blood-drawing procedure.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results and monitor as appropriate for liver disease.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Alkaline Phosphatase (ALP), Total; 5’-Nucleotidase

Alkaline phosphatase is an enzyme originating mainly in the bone, liver, and placenta, with some activity in the kidney and intestines. It is called alkaline because it functions best at a pH of 9. ALP levels are age and gender dependent. Post puberty ALP is mainly of liver origin.

Alkaline phosphatase is used as an index of liver and bone disease when correlated with other clinical findings. In bone disease, the enzyme level rises in proportion to new bone cell production resulting from osteoblastic activity and the deposit of calcium in the bones. In liver disease, the blood level rises when excretion of this enzyme is impaired as a result of obstruction in the biliary tract. Used alone, alkaline phosphatase may be misleading.

Reference Values

Normal Females: 1–12 years: <350 U/L >15 years: 25–100 U/L
Males: 1–12 years: <350 U/L 12–14 yrs: <500 U/L >20 yrs: 25–100 U/L
Normal values are higher in pediatric patients and in pregnancy. Values increase up to 3 times in puberty. Check with your laboratory for reference values. Values may vary with method of testing.

Procedure

1. Obtain a 5-mL fasting venous blood sample. Serum is used for this test. Anticoagulants may not be used. Observe standard precautions. Place specimen in a biohazard bag.
2. Refrigerate sample as soon as possible.
3. Note age and gender on test requisition.

Clinical Implications

1. Elevated levels of ALP in liver disease (correlated with abnormal liver function tests) occur in the following conditions:
   a. Obstructive jaundice (gallstones obstructing major biliary ducts; accompanying elevated bilirubin)
   b. Space-occupying lesions of the liver such as cancer (hepatic carcinoma) and malignancy with liver metastasis
   c. Hepatocellular cirrhosis
   d. Biliary cirrhosis
   e. Intrahepatic and extrahepatic cholestasis
   f. Hepatitis, infectious mononucleosis, cytomegalovirus
   g. Diabetes mellitus (causes increased synthesis), diabetic hepatic lipidosis
   h. Chronic alcohol ingestion
   i. Gilbert's syndrome

2. Bone disease and elevated ALP levels occur in the following conditions:
   a. Paget's disease (osteitis deformsans; levels 10 to 25 times normal)
   b. Metastatic bone tumor
   c. Osteogenic sarcoma
   d. Osteomalacia (elevated levels help differentiate between osteomalacia and osteoporosis, in which there is no elevation), rickets
3. Other diseases involving elevated ALP levels include the following:
   a. Hyperparathyroidism (accompanied by hypercalcemia), hyperthyroidism
   b. Pulmonary and myocardial infarctions
   c. Hodgkin’s disease
   d. Cancer of lung or pancreas
   e. Ulcerative colitis, peptic ulcer
   f. Sarcoidosis
   g. Perforation of bowel (acute infarction)
   h. Amyloidosis
   i. Chronic renal failure
   j. Congestive heart failure
   k. Hyperphosphatasia (primary and secondary)
4. Decreased levels of ALP occur in the following conditions:
   a. Hypophosphatasia (congenital)
   b. Malnutrition, scurvy
   c. Hypothyroidism, cretinism
   d. Pernicious anemia and severe anemias
   e. Magnesium deficiency
   f. Milk alkali (Burnett’s syndrome)
   g. Celiac sprue
   h. Magnesium and zinc deficiency (nutritional)

Clinical Alert
1. This test should not be done if the total alkaline phosphatase level is normal.
2. For evaluation of the biliary tract, alternative tests such as GGT, leucine aminopeptidase (LAP), and 5’-nucleotidase studies are recommended over the ALP ISO test.
3. Alkaline phosphatase isoenzymes have little value in children and adolescents because bone and liver fractions are normally elevated.
4. In pregnancy, marked decline of the placental isoenzyme is seen with placental insufficiency and imminent fetal demise.

Interfering Factors
1. A variety of drugs produce mild to moderate increases or decreases in ALP levels. See Appendix J for drugs that affect outcomes.
2. Young children, those experiencing rapid growth, pregnant women, and postmenopausal women have physiologically high levels of ALP; this level is slightly increased in older persons.
3. After IV administration of albumin, there is sometimes a marked increase in ALP for several days.
4. ALP levels increase at room temperature and in refrigerated storage. Testing should be done the same day.
5. ALP levels decrease if blood is anticoagulated.
6. ALP levels increase after fatty meals.

Interventions
Pretest Patient Care
1. Explain test purpose and blood-drawing procedure. Fasting is required.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Have patient resume normal activities.
2. Interpret test results and monitor appropriately for liver or bone disease and evidence of tumor. Testing for 5’-nucleotidase provides supportive evidence in the diagnosis of liver disease. When ALP and 5’-nucleotidase test results are evaluated, they provide definitive diagnosis of Paget’s disease and rickets, in which high levels of ALP accompany normal (0–5 U/L) or marginally increased 5’-nucleotidase activity. 5’-Nucleotidase is increased in liver disease (eg, hepatic carcinoma, biliary cirrhosis, extrahepatic obstruction, metastatic neoplasia of liver).
3. Remember that to confirm biliary abnormality, a useful test is gamma glutamyltransferase (GGT). The GGT test is elevated in hepatobiliary disease, but not in uncomplicated bone disease.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Alkaline Phosphatase Isoenzymes (ISO)
The isoenzymes of ALP are produced in various tissues. AP-1, a_2 is produced in the liver and by proliferating blood vessels. AP-2, β_1 is produced by bone and placental tissue. The intestinal isoenzyme AP-3, β_2 is present in small quantities in group O and B individuals who are Lewis-positive secretors. Placental ALP is present in the last trimester of pregnancy.

Any patient with an elevation of serum total alkaline phosphatase is a candidate for ALP isoenzyme study. The ALP ISO is mainly used to distinguish between bone and liver elevations of alkaline phosphatase.

Reference Values
Normal AP-1, a_2: values (liver) reported as weak, moderate, or strong or 24–158 U/L (0.40–2.64 µkat/L) AP-2, β_1:
values (bone) reported as weak, moderate, or strong or 24–146 U/L (0.40–2.44 µkat/L) AP-3, \( \beta_2 \); values (intestines) reported as weak, moderate, or strong or 0–22 U/L (0–0.36 µkat/L) AP-4: values (placental) reported as weak, moderate, or strong. Placental AP-4 is found only in pregnant women.

**Procedure**

1. Obtain a 5-mL fasting venous blood sample in a plain red-topped tube or SST tube. Serum is needed. Centrifuge blood promptly, with 30 minutes after draw.
2. Observe standard precautions. Place specimen in a biohazard bag.
3. Refrigerate if not tested immediately.

**Clinical Implications**

1. Liver (AP-1, \( \alpha_2 \)) isoenzymes are elevated in hepatic and biliary diseases such as the following conditions:
   a. Cirrhosis (hepatic)
   b. Hepatic carcinoma
   c. Biliary obstruction, primary biliary cirrhosis
2. Bone (AP-2, \( \beta_1 \)) isoenzymes are elevated in the following conditions:
   a. Paget's disease
   b. Hyperparathyroidism
   c. Bone cancer, rickets (all types)
   d. Osteomalacia, osteoporosis
   e. Malabsorption syndrome
   f. Certain renal disorders (uremia bone disease or renal rickets)
3. Intestinal (AP-3, \( \beta_2 \)) isoenzymes are elevated in the following conditions:
   a. Intestinal infarction
   b. Ulcerative lesions of stomach, small intestine, and colon
   c. Individuals with blood type O or B secrete intestinal isoenzymes 2 hours after a meal.
4. Placental (AP-4) isoenzymes are increased in the following conditions:
   a. Pregnancy (late in third trimester to onset of labor)
   b. Complications of pregnancy such as hypertension and preeclampsia
5. Placental-like isoenzymes occur in some cancers (unidentified isoenzymes):
   a. Regan's isoenzyme
   b. Nagao's isoenzyme

**Interfering Factors** Same as for alkaline phosphatase.

**Interventions**

**Pretest Patient Care**

1. See [total alkaline phosphatase patient pretest care](#) on page 389.
2. Remember that the same guidelines apply to alkaline phosphatase isoenzyme testing.

**Posttest Patient Aftercare**

1. See [total alkaline phosphatase patient posttest aftercare](#) on page 389.
2. Remember that the same guidelines apply to alkaline phosphatase isoenzyme testing.

**Angiotensin-Converting Enzyme (ACE)**

Angiotensin I is produced by the action of renin on angiotensinogen. Angiotensin I–converting enzyme (ACE) catalyzes the conversion of angiotensin I to the vasoactive peptide angiotensin II. Angiotensin I is concentrated in the proximal tubules.

This test is used primarily to evaluate the severity and activity of sarcoidosis. Serial determinations may be helpful in following the clinical course of the disease with steroid treatment. It is also used in the investigation of Gaucher’s disease.

**Reference Values**

*Normal* 8–53 U/L or 0.14–0.88 µkat/L. Check with your laboratory for reference values for infants and children—they are generally higher.

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum or heparinized plasma is used.
2. Observe standard precautions. Place specimen in a biohazard bag.
3. Freeze specimen if test is not performed immediately.

**Clinical Implications**

1. *Increased ACE levels* are associated with the following conditions:
   a. Sarcoidosis (ACE levels reflect the severity of the disease, with 68% positivity in stage 1 disease, 86% in stage 2, and 92% in stage 3)
   b. Gaucher's disease
   c. Leprosy
   d. Acute and chronic bronchitis
   e. Connective tissue diseases
f. Amyloidosis
g. Pulmonary fibrosis
h. Fungal diseases and histoplasmosis
i. Untreated hyperthyroidism
j. Diabetes mellitus
k. Psoriasis

2. **Decreased ACE levels** occur in the following conditions:
   a. Following prednisone treatment for sarcoidosis (steroid therapy)
   b. Advanced lung neoplasms
   c. Starvation

**Interfering Factors**

1. This test should not be done in persons <20 years of age because they normally have a very high level of ACE.
2. About 5% of the normal adult population have elevated ACE levels.
3. ACE is inhibited by EDTA anticoagulant.
4. Some antihypertensives may cause low ACE values.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and blood-drawing procedure.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results and monitor as appropriate for sarcoidosis and amyloid disease.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Amylase and Lipase**

Amylase, an enzyme that changes starch to sugar, is produced in the salivary (parotid) glands and pancreas; much lower activities are present in the ovaries, intestines, and skeletal muscle. If there is an inflammation of the pancreas or salivary glands, much amylase enters the blood. Amylase levels in the urine reflect blood changes by a time lag of 6 to 10 hours. (See Amylase Excretion/Clearance, Chapter 3). Lipase is a glycoprotein that, in the presence of bile salts and colipase, changes fats to fatty acids and glycerol. The pancreas is the major source of this enzyme. Lipase appears in the blood following pancreatic damage at the same time amylase appears (or slightly later) but remains elevated much longer than amylase (7 to 10 days).

Amylase and lipase tests are used to diagnose and monitor treatment of acute pancreatitis and to differentiate pancreatitis from other acute abdominal disorders (80% of patients with acute pancreatitis will have elevated amylase and lipase levels; lipase stays elevated longer). Lipase assay provides better sensitivity and specificity and is best used with amylase determination.

**Reference Values**

**Normal**

- Amylase: Newborns: 6–65 U/L or 0.1–1.1 µkat/L; Adults: 25–125 U/L or 0.4–2.1 µkat/L; Elderly persons (>60 years): 24–151 U/L or 0.4–2.5 µkat/L
- Lipase: Adults: 10–140 U/L or 0.17–2.3 µkat/L; Elderly persons (>60 years): 18–180 U/L or 0.30–3.0 µkat/L

Normal values vary widely according to method of testing; check with your laboratory for reference ranges. Amylase levels are low for the first 2 months of life. Most of the activity is of salivary origin. Children up to 2 years of age have virtually no pancreatic amylase.

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum is used. (EDTA, citrate, and oxalate anticoagulant interfere with lipase testing.)
2. Observe standard precautions. Place specimen in a biohazard bag.

**Clinical Implications**

1. **Greatly increased amylase levels** occur in acute pancreatitis early in the course of the disease. The increase begins in 3 to 6 hours after the onset of pain.
2. **Increased amylase levels** also occur in the following conditions:
   a. Chronic pancreatitis, pancreatic trauma, pancreatic carcinoma, obstruction of pancreatic duct
   b. Partial gastrectomy
   c. Acute appendicitis, peritonitis
   d. Perforated peptic ulcer
   e. Cerebral trauma, shock
   f. Obstruction or inflammation of salivary duct or gland and mumps
   g. Acute cholecystitis (common duct stone)
   h. Intestinal obstruction with strangulation
   i. Ruptured tubal pregnancy and ectopic pregnancy
   j. Ruptured aortic aneurysm
   k. Macroamylasias
3. **Decreased amylase levels** occur in the following conditions:
   a. Pancreatic insufficiency
   b. Hepatitis, severe liver disease
c. Advanced cystic fibrosis  
d. Pancreatectomy  

4. Elevated lipase levels occur in pancreatic disorders (eg, pancreatitis, alcoholic and nonalcoholic; pancreatic carcinoma).

5. Increased lipase values also are associated with the following conditions:
   a. Cholecystitis
   b. Hemodialysis
   c. Strangulated or infarcted bowel
   d. Peritonitis
   e. Primary biliary cirrhosis
   f. Chronic renal failure

6. Serum lipase levels are normal in patients with elevated amylase who have peptic ulcer, salivary adenitis, inflammatory bowel disease, intestinal obstruction, and macroamylasemia. Coexistence of increased serum amylase and normal lipase levels may be a helpful clue to the presence of macroamylasemia.

### Clinical Alert

<table>
<thead>
<tr>
<th>Panic Level for Lipase</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;600 IU/L or &gt;10 µkat/L</td>
</tr>
</tbody>
</table>

### Interfering Factors

1. Amylase  
   a. Anticoagulated blood gives lower results. Do not use EDTA, citrate oxalate.  
   b. Lipemic serum interferes with test.  
   c. Increased levels are found in alcoholic patients and pregnant women and in diabetic ketoacidosis.  
   d. Many drugs can interfere with this test (see Appendix J).

2. Lipase  
   a. EDTA anticoagulant interferes with test.  
   b. Lipase is increased in about 50% of patients with chronic renal failure.  
   c. Lipase increases in patients undergoing hemodialysis.  
   d. Many drugs can affect outcomes. See Appendix J.

### Interventions

#### Pretest Patient Care

1. Explain test purpose and procedure. Amylase and lipase testing are done together in the presence of abdominal pain, epigastric tenderness, nausea, and vomiting. These findings characterize acute pancreatitis as well as other acute surgical emergencies.

2. If amylase/creatinine clearance testing is also being done, collect a single, random urine sample at the same time blood is drawn.

3. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

#### Posttest Patient Aftercare

1. Have patient resume normal activities.

2. Interpret test results and monitor as appropriate for pancreatitis or other acute abdominal conditions.

3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

### Aspartate Transaminase (Aminotransferase, AST); Serum Glutamic-Oxaloacetic Transaminase (SGOT)

Aspartate transaminase (AST) is an enzyme present in tissues of high metabolic activity; decreasing concentrations of AST are found in the heart, liver, skeletal muscle, kidney, brain, pancreas, spleen, and lungs. The enzyme is released into the circulation following the injury or death of cells. Any disease that causes change in these highly metabolic tissues will result in a rise in AST levels. The amount of AST in the blood is directly related to the number of damaged cells and the amount of time that passes between injury to the tissue and the test. Following severe cell damage, the blood AST level will rise in 12 hours and remain elevated for about 5 days.

This test is used to evaluate liver and heart disease. The ALT is usually ordered along with the AST.

### Reference Values

**Normal**  
Men: 14–20 U/L or 0.23–0.33 µkat/L  
Women: 10–36 U/L or 0.17–0.60 µkat/L  
Newborns: 47–150 U/L or 0.78–2.5 µkat/L  
Children: 9–80 U/L or 0.15–1.3 µkat/L  
Check with your laboratory. Different methods have different reference values.

#### Procedure

1. Obtain a 5-mL venous sample. Serum is used. Observe standard precautions. Place specimen in a biohazard bag.

2. Avoid hemolysis.

### Clinical Implications

1. Increased AST levels occur in MI.
   a. In MI, the AST level may be increase to 4 to 10 times the normal values.
   b. The AST level reaches a peak in 24 hours and returns to normal by post-MI day 3 to 7. Secondary rises in AST
levels suggest extension or recurrence of MI.

c. The AST curve in MI parallels that of creatinine phosphokinase (CPK).

2. Increased AST levels occur in liver diseases (10–100 times normal).
   a. Acute hepatitis and chronic hepatitis (ALT > AST)
   b. Active cirrhosis (drug induced; alcohol induced: AST > ALT)
   c. Infectious mononucleosis
   d. Hepatic necrosis and metastasis
   e. Primary or metastatic carcinoma
   f. Alcoholic hepatitis
   g. Reye’s syndrome

3. Other diseases associate with elevated AST levels include the following:
   a. Hypothyroidism
   b. Trauma and irradiation of skeletal muscle
   c. Dermatomyositis
   d. Polyomysis
   e. Toxic shock syndrome
   f. Cardiac catheterization
   g. Recent brain trauma with brain necrosis, cerebral infarction
   h. Crushing and traumatic injuries, head trauma, surgery
   i. Progressive muscular dystrophy (Duchenne’s)
   j. Pulmonary emboli, lung infarction
   k. Gangrene
   l. Malignant hyperthermia, heat angiography
   m. Mushroom poisoning
   n. Shock
   o. Hemolytic anemia, exhaustion, heat stroke

4. Decreased AST levels occur in the following conditions:
   a. Azotemia
   b. Chronic renal dialysis
   c. Vitamin B₆ deficiency

Interfering Factors

1. Slight decreases occur during pregnancy, when there is abnormal metabolism of pyridoxine.
2. Many drugs can cause elevated or decreased levels (see Appendix J). Alcohol ingestion affects results.
3. Exercise and IM injections do not affect results.
4. False decreases occur in diabetic ketoacidosis, severe liver disease, and uremia.
5. Gross hemolysis causes falsely high levels.

**Critical Value**

AST is extremely high (>20,000 U/L; >333 µkat/L) in alcohol-acetaminophen syndrome. AST > ALT, prothrombin time: 100 seconds. Creatinine: >34 mg/L or >0.30 mmol/L

**Interventions**

*Pretest Patient Care*

1. Explain test purpose and blood-drawing procedure. For diagnosis of MI, AST testing should be done on 3 consecutive days because the peak is reached in 24 hours and levels return to normal in 3 to 4 days.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

*Posttest Patient Aftercare*

1. Interpret test results and monitor appropriately for heart and liver diseases.
2. Ensure that unexplained AST elevations are further investigated with ALT and GGT tests.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Cardiac Troponin T (cTnT): Troponin I (cTnl)

Cardiac troponin is unique to the heart muscle and is highly concentrated in cardiomyocytes. These isoforms show a high degree of cardiac specificity. This protein is released with very small areas of myocardial damage as early as 1 to 3 hours after injury, and levels return to normal within 5 to 7 days. Troponin I remains increased longer than CK-MB and is more cardiac specific. Troponin T is more sensitive but less specific, being positive with angina at rest. These tests are becoming the most important addition to the clinical assessment of cardiac injury.

This test is used in the early diagnosis of small myocardial infarcts that are undetectable by conventional diagnostic methods. Cardiac troponin levels are also used later in the course of MI because they remain elevated for 5 to 7 days after injury. A single sample may be misleading; therefore, serial sampling 0, 4, 8, and 12 hours after chest pains may be ordered to rule out acute MI. See Table 6.11 for a list of cardiac markers.

**Table 6.11 Cardiac Markers**
Markers | Time of Initial Evaluation | Time of Peak Evaluation | Time to Return to Normal
---|---|---|---
CK-MB | 4–8 h | 12–24 h | 72–96 h
LDH | 2–5 days | | 10 days
Myoglobin | 2–4 h | 8–10 h | 24 h
Troponin I (cTnI) | 4–6 h | 12 h | 3–10 days
Troponin T (cTnT) | 4–8 h | 12–48 h | 7–10 days

Reference Values

**Normal** (Negative Qualitative) Troponin I: <0.35 ng/mL or <0.35 µg/L Troponin T: <0.2 ng/mL or <0.2 µg/L Total CK: 0–120 ng/mL or 0–120 µg/L CK-MB: 0–3 ng/mL or 0–3 µg/L CK index: 0–3 LDH: 140–280 U/L or 2.34–4.68 µkat/L Myoglobin: <55 ng/mL or <55 µg/L Troponin: <0.4 ng/mL or <0.4 µg/L Values may vary depending on the testing method used. Check with your laboratory for reference values.

**Procedure**

1. Obtain a 5-mL venous blood sample in a red-topped tube within hours after onset of chest pain. Observe standard precautions. Place specimen in a biohazard bag.
2. Be aware that serial samples may be ordered. Record date and time of sampling.

**Clinical Implications**

1. Positive or elevated cardiac troponin I levels indicate:
   a. Small infarcts; increases remain for 5 to 7 days.
   b. Myocardial injury during surgery
2. Positive or elevated cardiac troponin T
   a. Acute MI
   b. Perioperative MI
   c. Unstable angina
   d. Myocarditis
   e. Some noncardiac events
      1. Chronic renal failure
      2. Acute trauma involving muscle
      3. Rhabdomyolysis, polymyositis, dermatomyositis

**Interfering Factors**

1. Cardiac troponin T levels may be increased in chronic muscle or renal disease and trauma.
2. Levels are not affected by orthopedic or lung surgery.

---

**Clinical Alert**

**Critical Value**

Troponin I: >1.5 ng/mL or >1.5 µg/L

**Interventions**

**Pretest Patient Care**

1. Explain that the test is a sensitive marker for minor myocardial injury in unstable angina.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results, and counsel and monitor appropriately. Additional testing may be necessary (eg, cardiac myosin light classes, glycogen phosphorylcholine BB [GPBB]).
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Creatine Phosphokinase (CPK); Creatine Kinase (CK); CPK and CK Isoenzymes**

Creatine kinase (CPK/CK) is an enzyme found in higher concentrations in the heart and skeletal muscles and in much smaller concentrations in brain tissue. Because CK exists in relatively few organs, this test is used as a specific index of injury to myocardium and muscle. CPK can be divided into three isoenzymes: MM or CK 3, BB or CK 1, and MB or CK 2. CK-MM is the isoenzyme that constitutes almost all the circulatory enzymes in healthy persons. Skeletal muscle contains primarily MM; cardiac muscle contains primarily MM and MB; and brain tissue, GI system, and genitourinary tract contain primarily BB. Normal CK levels are virtually 100% MM isoenzyme. A slight increase in total CPK is reflected from elevated BB from CNS injury. CPK isoenzyme studies help distinguish whether the CPK originated from the heart (MB) or the skeletal muscle (MM).

The CK (CPK) test is used in the diagnosis of MI and as a reliable measure of skeletal and inflammatory muscle diseases. CK levels can prove helpful in recognizing muscular dystrophy before clinical signs appear. CK levels may rise significantly with CNS disorders such as Reye's syndrome. The determination of CK isoenzymes may be helpful in making a differential diagnosis. Elevation of MB, the cardiac isoenzyme, provides a more definitive indication of myocardial cell damage than total CK alone. MM isoenzyme is an indicator of skeletal muscle damage. Newer tests, such
as CK isoforms, allow for earlier detection of MI than is possible with CK-MB.

**Reference Values**

<table>
<thead>
<tr>
<th>Normal</th>
<th>Men: 38–174 U/L (0.63–2.90 µkat/L)</th>
<th>Women: 26–140 U/L (0.46–2.38 µkat/L)</th>
<th>Infants: 2–3 times adult values</th>
</tr>
</thead>
</table>

**Isoenzymes:**

- MM (CK<sub>3</sub>): 96%–100%
- MB (CK<sub>2</sub>): 0%–6%
- BB (CK<sub>1</sub>): 0%

**NOTE**

Normal values may vary with method of testing and reaction temperature. Check with your laboratory.

**NOTE**

Healthy African Americans have higher CK levels than do Caucasian and Hispanic persons.

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum must be used.
2. Observe standard precautions. Place specimen in a biohazard bag.
3. If a patient has been receiving multiple IM injections, note this fact on the laboratory requisition.
4. Avoid hemolysis.

**Clinical Implications**

1. **Total CK Levels**
   a. *Increased CK/CPK levels* occur in the following conditions:
      1. **Acute MI**
         a. With MI, the rise starts soon after an attack (about 4–6 hours) and reaches a peak of at least several times normal within 24 hours. CK returns to normal in 48 to 72 hours.
         b. CK and CK-MB (CK<sub>2</sub>) peaks about 1 day after onset, as does AST.
         c. Lactate dehydrogenase (LD) usually peaks 2 days after onset, when the LD<sub>1</sub>–LD<sub>2</sub> inversion (flip) is found.
         d. CK-MB, LD<sub>1</sub>, LD<sub>1</sub>–LD<sub>2</sub> ratio, total CK, and total LD classically increase with acute MI. CK-MB and LD<sub>1</sub> increase both in percentage and absolutely (each isoenzyme percentage times the respective total enzyme), peak, and then decrease.
         e. AST testing with LD and LD isoenzymes is advocated when the patient reaches medical attention 48 to 72 hours after onset of a possible acute MI.
      2. **Severe myocarditis**
      3. **After open heart surgery**
      4. **Cardioversion (cardiac defibrillation)**
      5. **Myocarditis**
   b. Other diseases and procedures that cause increased CK/CPK levels include the following:
      1. **Acute cerebrovascular disease**
      2. **Progressive muscular dystrophy** (levels may reach 20–200 times normal), Duchenne's muscular dystrophy, female carriers of muscular dystrophy
      3. **Dermatomyositis and polymyositis**
      4. **Delirium tremens and chronic alcoholism**
      5. **Electric shock, electromyography**
      6. **Malignant hyperthermia**
      7. **Reye's syndrome**
      8. **Convulsions, ischemia, or subarachnoid hemorrhage**
      9. **Last weeks of pregnancy and during childbirth**
      10. **Hypothyroidism**
      11. **Acute psychosis**
      12. **CNS trauma, extensive brain infarction**
      13. **Neoplasms of prostate, bladder, or GI tract**
      14. **Rhabdomyolysis with cocaine intoxication**
      15. **Eosinophilia-myalgia syndrome**
   c. Normal values are found in myasthenia gravis and multiple sclerosis.
   d. Decreased values have no diagnostic meaning and may be caused by low muscle mass and bed rest (overnight values can drop 20%).

2. **CK Isoenzymes**
   a. *Elevated MB (CK<sub>2</sub>) isoenzyme levels* occur in the following conditions:
      1. **Myocardial infarct** (rises 4–6 hours after MI; not demonstrable after 24–36 hours; ie, peak with rapid fall)
      2. **Myocardial ischemia, angina pectoris**
      3. **Duchenne's muscular dystrophy**
      4. **Subarachnoid hemorrhage**
      5. **Reye's syndrome**
      6. **Muscle trauma, surgery (postoperative)**
      7. **Circulatory failure and shock**
      8. **Infections of heart—myocarditis**
      9. **Chronic renal failure**
      10. **Malignant hyperthermia, hypothermia**
      11. **CO poisoning**
      12. **Polymyositis**
      13. **Myoglobulinemia**
      14. **Rocky Mountain spotted fever**
b. BB (CK₁) elevations occur in the following conditions:
   1. Reye’s syndrome
   2. Some breast, bladder, lung, uterus, testes, and prostate cancers
   3. Severe shock syndrome
   4. Brain injury, neurosurgery
   5. Hypothermia
   6. Following coronary bypass surgery
   7. Newborns

c. MM (CK₂) is elevated in most conditions in which total CK is elevated.

d. MB (CK₂) is not elevated:
   1. Exercise (total elevated)
   2. IM injections (total elevated)
   3. Strokes, cerebrovascular accident, and other brain disorders in which total CK is elevated
   4. Pericarditis
   5. Pneumonia, other lung diseases; pulmonary embolism
   6. Seizures (CK total may be very high)

Cl**inical Alert**

1. After an MI, MB appears in the serum in 6 to 12 hours and remains for about 18 to 32 hours. The finding of MB in a patient with chest pain is diagnostic of MI. In addition, if there is a negative CK-MB for ≥48 hours following a clearly defined episode, it is clear that the patient has not had an MI.
2. CK-MB, LD1, LD1/LD-2 ratio, total CK, and total LD classically increase with acute MI. CK-MB and LD1 increase both in percentage and absolutely (each isoenzyme percent times the respective total enzyme), peak, then decrease.

Interfering Factors

1. Strenuous exercise, weight lifting, and surgical procedures that damage skeletal muscle may cause increased levels of CK.
2. Alcohol and other drugs of abuse increase CK levels.
3. Athletes have a higher CK value because of greater muscle mass.
4. Multiple IM injections may cause increased or decreased CK levels (see Appendix J).
5. Many drugs may cause increased CK levels.
6. Childbirth may cause increased CK levels.
7. Hemolysis of blood sample causes increased CK levels.

Interventions

**Pretest Patient Care**

1. Explain test purpose and need for at least three consecutive blood draws following episode.
2. Note on requisition when suspected cardiac episode occurred, and dates and times of blood draws.
3. Do not allow exercise before test.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. Interpret test results and monitor as appropriate for MI, muscular dystrophy, and other causes of abnormal test outcomes.
3. Remember that high levels of CK/CK-MB may suggest other tests should be done to support diagnosis of acute MI:
   a. Total leukocyte count and differential
   b. Cardiac troponin T
   c. Myoglobin
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Galactose-1-Phosphate Uridylytransferase (GPT); Galaktokinase; Galactose-1-Phosphate**

The enzyme galactose-1-phosphate uridylytransferase is needed in the use of galactose-1-phosphate so that it does not accumulate in the body. A very rare genetic disorder resulting from an inborn (inherited or acquired during intrauterine development) error of galactose metabolism may occur.

This measurement is used to identify galactose defects, which can result in widespread tissue damage and abnormalities such as cataracts, liver disease, and renal disease. It also causes failure to thrive and mental retardation. The screening test should be done immediately to enable diet treatment if test is positive.

**Reference Values**

**Normal**

- Galactose-1-phosphate uridylytransferase: 18.5–28.5 U/g of hemoglobin (Hb) or 1.19–1.84 mU/mol Hb
- Galactose-1-phosphate (dried blood spot-screening): <0.74 mmol/L
- Galaktokinase Children 0–2 years: 11–150 mU/g Hb or 183–2500 pkat/g Hb
- Children 2–18 years: 11–54 mU/g Hb or 183–900 pkat/g Hb
- Adults: 12–40 mU/g Hb or 200–667 pkat/g Hb

**Procedure**

1. Obtain a 5-mL venous blood sample.
2. Anticoagulate with heparin.
3. Observe standard precautions. Place specimen in a biohazard bag.

**Clinical Implications** Decreased values are associated with galactosemia, a rare genetic disorder transmitted in an autosomal recessive fashion. The resulting accumulation of galactitol and/or galactose-1-phosphate can result in juvenile cataracts, liver failure, failure to thrive, and mental retardation in persons with galactose-1-phosphate uridylytransferase deficiency.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. Genetic counseling may be necessary.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results and counsel appropriately. See newborn screening in Chapter 11.
2. Instruct parents of infants and children with positive test results that the disease can effectively treated by removing galactose-containing foods, especially milk, from the diet. With dietary galactose restriction, liver and lens changes are reversible.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Hexosaminidase, Total and Isoenzyme A**

Three isoenzymes of hexosaminidase have been identified in serum: A (acid form), B (base form), and S. Hexosaminidase A is a lysosomal isoenzyme, deficiency of which characterizes patients with Tay-Sachs disease. Homozygotes have no hexosaminidase A and a large increase in hexosaminidase B and S. Heterozygotes have a moderate decrease in hexosaminidase A and a slight increase of hexosaminidase B and S.

Hexosaminidase A is used as a diagnostic test for Tay-Sachs disease and can be of help in identifying carriers among persons with no family history of Tay-Sachs. This condition is due to an autosomal recessive trait found predominantly, but not exclusively, in Ashkenazi Jews and is characterized by the appearance during infancy of psychomotor deterioration, blindness, cherry-red spot on the macula, and an exaggerated extension response to sound. In the brains of affected children, the level of ganglioside is increased 100 times owing to the deficiency of this enzyme.

**Reference Values**

**Normal Percentage of normal total hexosaminidase A** 56%–80% noncarrier: 7.2–9.88 U/L or 120–165 nkat/L <50% heterozygous: 3.30–5.39 U/L or 55–90 nkat/L 0% Tay-Sachs: 0 U/L Total hexosaminidase Noncarrier: 9.83–15.95 U/L or 164–266 nkat/L Heterozygous: 3.30–5.39 U/L or 55–90 nkat/L (carrier) Homozygous Tay-Sachs: 17.1 U/L or 285 nkat/L Leukocyte hexosaminidase total: 16.4–36.2 U/g cellular protein or 273–603 nkat/g cellular protein Leukocyte hexosaminidase A: 63%–75% of total (normal) Normal values vary with method of testing used. Check with your laboratory for reference values.

**Procedure**

1. Obtain a 5-mL venous blood sample. Allow blood to clot at +3°C and centrifuge at 3°C. The test uses serum. If the test is not performed immediately, serum must be frozen.
2. Be aware that if leukocyte hexosaminidase A is ordered also, a heparinized sample is needed. Place in ice immediately. Place specimens in biohazard bags.

**Clinical Implications**

1. **Decreased hexosaminidase A.** An almost total deficiency of the A component is diagnostic of Tay-Sachs disease or GM2 gangliosidosis. The total hexosaminidase is of no value in Tay-Sachs.
2. **Decreased hexosaminidase A and B.** In a variant of Tay-Sachs disease known as Sandhoff's disease, both A and B isoenzymes are defective, causing an absence of this enzyme. The total hexosaminidase level is also decreased in Sandhoff's disease.
3. **Increased total hexosaminidase** occurs in the following conditions:
   a. Hepatic disease (biliary obstruction)
   b. Gastric cancer
   c. Myeloma
   d. MI
   e. Vascular complications of diabetes mellitus

**Clinical Alert**

Pregnancy results in increased serum levels of total hexosaminidase and decreased hexosaminidase A, which gives a false appearance of being a carrier. Therefore, during pregnancy, the serum test should never be done. However, the leukocyte test is valid in pregnancy and should be used.

**Interfering Factors**

1. Total values are increased in pregnancy (5 times normal).
2. Oral contraceptives falsely increase values.
Clinical Alert

Critical Values for Hexosaminidase A

Less than 50% of total activity indicates Tay-Sachs carrier. If the serum Hex A test levels are ambiguous the leukocyte test should be done.

Interventions

**Pretest Patient Care**

1. Explain test purpose and procedure. Genetic counseling may occur before testing.
2. Be aware that pregnancy and/or oral contraceptives are contraindications for testing.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results and be prepared to perform genetic counseling of patient and family. The physician should be informed as soon as possible.
2. Follow guidelines in Chapter 1 for safe, effective, informed, posttest care.

Lactate Dehydrogenase (LD, LDH)

Lactate dehydrogenase is an intracellular enzyme that is widely distributed in the tissues of the body, particularly in the kidney, heart, skeletal muscle, brain, liver, and lungs. Increases in the reported value usually indicate cellular death and leakage of the enzyme from the cell.

Although elevated levels of LDH are nonspecific, this test is useful in confirming myocardial or pulmonary infarction when viewed in relation to other test findings. For example, LD remains elevated longer than CK in MI. LDK level is also helpful in the differential diagnosis of muscular dystrophy and pernicious anemia. More specific findings may be found by breaking down the LDH into its five isoenzymes. (When LD values are reported or quoted, total LDH is meant.)

**Reference Values**

**Normal**

- Newborn: 160–450 U/L
- Children: 60–170 U/L
- Adults: 140–280 U/L

Normal values vary with method of testing used. Check with your laboratory for reference values.

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum is used. Observe standard precautions.
2. Avoid hemolysis in obtaining blood sample. Place specimen in a biohazard bag.

**Clinical Implications**

1. *Increased LDH (LD)* occurs in the following conditions:
   a. High levels occur within 36 to 55 hours after MI and continue longer than elevations of SGOT or CPK (3–10 days). Differential diagnosis of acute MI may be accomplished with LDH isoenzymes.
   b. In pulmonary infarction, increased LDH occurs within 24 hours of pain onset. The pattern of normal SGOT and elevated LDH that levels off 1 to 2 days after an episode of chest pain is indicative of pulmonary infarction.
   c. *Elevated levels of LDH* are also observed in various other conditions:
      1. Congestive heart failure
      2. Liver diseases (eg, cirrhosis, alcoholism, acute viral hepatitis)
      3. Malignant neoplasms, cancer, leukemias, lymphoma
      4. Hypothyroidism
      5. Lung diseases
      6. Skeletal muscle diseases (muscular dystrophy), muscular damage
      7. Megaloblastic and pernicious anemias, hemolytic anemia, sickle cell disease
      8. Delirium tremens, seizures
      9. Shock, hypoxia, hypotension
      10. Hyperthermia
      11. Renal infarct
      12. CNS diseases
      13. Acute pancreatitis
      14. Fractures, other trauma including head
      15. Intestinal obstruction
   d. Angina and pericarditis do not produce LDH elevations.
2. *Decreased LDH levels* are associated with a good response to cancer therapy.

**Interfering Factors**

1. Strenuous exercise and the muscular exertion involved in childbirth cause increased LDH levels.
2. Skin diseases can cause falsely increased LDH levels.
3. Hemolysis of red blood cells due to freezing, heating, or shaking the blood sample will cause falsely increased LDH levels.
4. Various drugs may cause increased or decreased LDH levels (see Appendix J).
Clinical Alert

LDH is found in nearly every tissue of the body; therefore, elevated levels are of limited diagnostic value by themselves. Differential diagnoses may be accomplished with LD isoenzyme determination.

Interventions

Pretest Patient Care

1. Explain test purpose and blood-drawing procedure. Obtain recent history of MI or pulmonary infarction.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume normal activities.
2. Interpret test results and monitor for myocardial and pulmonary infarction and other diseases related to abnormal results. LD isoenzymes may be ordered.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Lactate Dehydrogenase (LDH, LD) Isoenzymes (Electrophoresis)

Electrophoresis, or separation, of LDH identifies the five isoenzymes of fractions of LDH, each with its own physical characteristics and electrophoretic properties. Fractioning the LDH activity sharpens its diagnostic value because LDH is found in many organs. LD isoenzymes are released into the bloodstream when tissue necrosis occurs. The isoenzymes are elevated in the terms of patterns established, not on the basis of the value of a single isoenzyme. The origins of the LDH isoenzymes are as follow: LD$_1$ and LD$_2$ are present in cardiac tissue and erythrocytes; LD$_3$ originates mainly from lung, spleen, pancreas, and placenta; and LD$_4$ and LD$_5$ originate from skeletal muscle and liver.

The five isoenzyme fractions of LDH show different patterns in various disorders. Abnormalities in the pattern suggest which tissues have been damaged. This test is useful in the differential diagnosis of acute MI, megaloblastic anemia (eg, folate deficiency, pernicious anemia), hemolytic anemia, and very occasionally, renal infarct. These entities are characterized by LD$_1$ increases, often with LD$_1$–LD$_2$ inversion (flip).

Reference Values

**Normal** LD$_1$: 17%–27% of total or 0.17–0.27 LDH$_2$: 29%–39% of total or 0.29–0.39 LDH$_3$: 19%–27% of total or 0.19–0.27 LDH$_4$: 8%–16% of total or 0.08–0.16 LDH$_5$: 6%–16% of total or 0.06–0.16

Procedure

1. Obtain a 5-mL venous blood sample. Serum is needed.
2. Avoid hemolysis.
3. Observe standard precautions. Place specimen in a biohazard bag. Be aware that serial determinations may be ordered (3 consecutive days).

Clinical Implications

1. Abnormal LD$_1$ and LD$_2$ patterns reflect damaged tissues (see Table 6.12).

**Table 6.12 Abnormal LD Isoenzyme Patterns**

<table>
<thead>
<tr>
<th>Disease</th>
<th>LD$_1$</th>
<th>LD$_2$</th>
<th>LD$_3$</th>
<th>LD$_4$</th>
<th>LD$_5$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Myocardial infarction</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pulmonary infarction</td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Congestive heart failure</td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Viral hepatitis</td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Toxic hepatitis</td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leukemia, granulocytic</td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Pancreatitis (extensive)</td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Megaloblastic anemia</td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Hemolytic anemia</td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Muscular dystrophy</td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
</tbody>
</table>

a. The appearance of an LD flip (ie, when LD$_1$ level is higher than LD$_2$ level) is extremely helpful in the diagnosis of MI. The presence of an LD flip 1 day following incident or with the detection of CK-MB is essentially diagnostic of MI if baseline cardiac enzymes/ isoenzymes are normal and if rises and falls are as anticipated for the diagnosis of acute MI.

b. Persistent LD$_1$–LD$_2$ flip following acute MI may represent reinfarction. When acute MI is complicated by shock, a normal pattern may be found. LD$_1$–LD$_2$ inversion commonly appears subsequent to the isomorphic pattern in instances of acute MI.

c. The LDH pattern in hemolytic, megaloblastic, and sickle cell anemia is essentially the same as in MI and other anemias. This is because red blood cells have an isoenzyme pattern similar to that of heart muscle. The time elapsed to peak values may help to differentiate these conditions.
2. LD $3$ increases occur in advanced cancer and malignant lymphoma; this level should decrease following effective therapy. LD $3$ is occasionally elevated in pulmonary infarction or pneumonia.

3. LD $5$ is increased in the following conditions:
   a. Liver disease, hepatitis
   b. Congestive heart failure, pulmonary edema
   c. Striated muscle trauma, burns

4. LD $5$ increase is more significant when LD $2$/LD $4$ ratio is increased.

5. In most cancers, one to three of the bands (LD $2$, LD $3$ and LD $4$) are frequently increased. A notable exception is in seminomas and dysgerminomas, in which LD $1$ is increased. Frequently, an increase in LD $3$ may be the first indication of the presence of cancer.

6. All LD isoenzymes are increased in systemic diseases (eg, carcinomatous collagen vascular, disseminated intravascular coagulation, sepsis) (see Table 6.12).

7. Increased total LD with normal distribution of isoenzymes may be seen in coronary artery disease (CAD) with chronic heart failure, hypothyroidism, infectious mononucleosis and other inflammatory states, uremia, and necrosis.

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Clinical Alert

1. LD isoenzyme testing should be reserved for diagnosis of complex cases. In 5% to 20% of patients with acute MI, the expected reversal of LD-1/LD-2 does not occur; in these patients, there is often simply an increase in LD-1.

2. LDH isoenzymes should be interpreted in light of clinical findings.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. Repeat testing on 3 consecutive days is likely. Obtain pertinent clinical signs and symptoms.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results and monitor appropriately for abnormal LD patterns, MI, and pulmonary infarctions.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Renin (Angiotensin); Plasma Renin Angiotensin (PRA)

Renin is an enzyme that converts angiotensinogen to angiotensin I. Derived from the liver, angiotensinogen is an $\alpha_2$-globulin in the serum. Angiotensin I is then converted in the lung to angiotensin II. Angiotensin II is a potent vasopressor agent responsible for hypertension of renal origin and is a powerful releaser of aldosterone from the adrenal cortex. Both angiotensin II and aldosterone increase blood pressure. Renin levels increase when there is decreased renal perfusion pressure. The renin-aldosterone axis regulates sodium and potassium balance and blood volume and pressure. Renal reabsorption of sodium affects plasma volume. Low plasma volume, low blood pressure, low sodium, and increased potassium induce renin release, causing increased aldosterone through stimulation of angiotensin. Potassium loss, acute blood pressure increases, and increased blood volumes suppress renin release.

This test is most useful in the differential diagnosis of hypertension, whether essential, renal, or renovascular. In primary hyperaldosteronism, the findings will demonstrate that aldosterone secretion is exaggerated and secretion of renin is suppressed. In renal vascular disease, renin is elevated.

Reference Values

Normal Renin activity—plasma (PRA) Adult (normal-sodium diet): Supine: 0.2–1.6 ng angiotensin I (AI)/mL/h or 0.2–1.6 µg AI/h/L Standing: 0.7–3.3 ng AI/mL/h or 0.7–3.3 µg AI/h/L Adult (low-sodium diet): Supine: renin levels increase 2 times normal. Standing: renin levels increase 6 times normal. Renin direct: Adult supine: 12–79 mU/L Adult standing: 13–114 mU/L

Procedure

1. Obtain a 5-mL venous blood sample. Fasting is required. Collect specimen with scrupulous attention to detail. Use EDTA as the anticoagulant to aid in preservation of any angiotensin formed before examination. Observe standard precautions.
2. Draw blood in chilled tubes and place samples on ice. Transport samples to laboratory immediately in a biohazard bag. Must be centrifuged in refrigerated centrifuge.
3. Record posture and dietary status of patient at time of blood drawing.
4. A 24-hour urine sodium should be done concurrently to aid in diagnosis.

Clinical Implications

1. Increased renin levels occur in the following conditions:
   a. Secondary aldosteronism with malignant hypertension
   b. Renovascular hypertension
   c. Reduced plasma volume due to low-sodium diet, diuretics, Addison’s disease, or hemorrhage.
   d. Chronic renal failure
   e. Salt-losing status owing to GI disease (Na and K wastage)
   f. Renin-producing tumors of kidney
g. Few patients (15%) with essential hypertension
h. Bartter's syndrome (high in renin hypertension)
i. Pheochromocytoma

2. Decreased renin levels are found in the following conditions:
   a. Primary aldosteronism (98% of cases)
   b. Unilateral renal artery stenosis
   c. Administration of salt-retaining steroids
   d. Congenital adrenal hyperplasia with 17-hydroxylase deficiency
   e. Liddle’s syndrome

Interfering Factors

1. Levels vary in healthy persons and increase under influences that tend to shrink the intravascular fluid volume.
2. Random specimens may be difficult to interpret unless dietary and salt intake of patient is regulated.
3. Values are higher when the patient is in an upright position, when the test is performed early in the day, when the patient is on a low-sodium diet, during pregnancy, and with drugs such as diuretics and antihypertensives and foods such as licorice. See Appendix J for other drugs that affect outcomes.
4. Recently administered radioisotopes interfere with test results.
5. Indomethacin and salicylates decrease renin levels.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Remember that a regular diet that contains 180 mEq (180 mmol/L) of sodium and 100 mEq (100 mmol/L) of potassium must be maintained for 3 days before the specimen is obtained. A 24-hour urine sodium and potassium should also be done to evaluate salt balance. The blood test should be drawn at the end of the 24 hour urine test.
3. Instruct the patient that it is necessary to be in a supine position for at least 2 hours before obtaining the specimen. The specimen is drawn with patient in the supine position.
4. Ensure that antihypertensive drugs, cyclic progestogens, estrogens, diuretics, and licorice are terminated at least 2 weeks and preferably 4 weeks before a renin-aldosterone workup.
5. Remember that if a standing specimen is ordered, the patient must be standing for 2 hours before testing, and blood should be drawn with the patient in the sitting position.
6. Do not allow caffeine ingestion the morning before or during the test.
7. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and counsel appropriately regarding hypertension, further testing and possible treatment.
2. Have patient resume normal activities.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Renin Stimulation/Challenge Test

**Challenge Test**

A challenge test distinguishes primary from secondary hyperaldosteronism on the basis of renin levels. The test is performed with the patient in both the recumbent and upright positions and after the patient has been maintained on a low-salt diet. In normal persons and in those with essential hypertension, renin concentration is increased by the reduction in volume due to sodium restriction and the upright position. In primary aldosteronism, volume depletion does not occur, and renin concentration remains low.

**Reference Values**

*Normal* See challenge test above.

**Procedure**

1. Admit the patient to the hospital for this test. On admission, obtain and record the patient's weight.
2. Ensure the patient follows a reduced-sodium diet supplemented with potassium for 3 days, along with diuretics (eg, furosemide, chlorothiazide), as ordered.
3. Weigh patient again on the third day, record data, and ensure that the patient remains upright for 4 hours and participates in normal activities.
4. Obtain a venous heparinized blood sample for renin at 11:00 a.m., when renin is usually at its maximum level. Place specimen on ice, and send it immediately to the laboratory in a biohazard bag.

**Interpretation of Renin Stimulation Test**

In healthy persons and most hypertensive patients, the stimulation of a low-salt diet, a diuretic, and upright posture will raise renin activity to very high levels and result in weight loss. However, in primary aldosteronism, the plasma level is expanded and remains so. In these patients, there is little if any weight loss, and the renin level is very low or undetectable. A response within the normal range can occur in the presence of aldosterone.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. The purpose of the preparation is to deplete the patient of sodium.
2. Check with individual laboratory for specific practices.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
Posttest Patient Aftercare

1. Resume normal activities.
2. Interpret test results and counsel appropriately regarding hypertension.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

?-Glutamyltransferase (?-Glutamyl Transpeptidase, GGT, ?GT)

The enzyme ?-glutamyl transpeptidase is present mainly in the liver, kidney, and pancreas. Despite the fact that the kidney has the highest level of this enzyme, the liver is considered the source of normal serum activity. ?GT has no origin in bone or placenta.

This test is used to determine liver cell dysfunction and to detect alcohol-induced liver disease. Because the GGT is very sensitive to the amount of alcohol consumed by chronic drinkers, it can be used to monitor the cessation or reduction of alcohol consumption in chronic alcoholic patients and early-risk drinkers. GT activity is elevated in all forms of liver disease. This test is much more sensitive than either the alkaline phosphatase test or the transaminase test (ie, SGOT, SGPT) in detecting obstructive jaundice, cholangitis, and cholecystitis. It is also indicated in the differential diagnosis of liver disease in children and pregnant women who have elevated levels of LDH and alkaline phosphatase. ?GT is also useful as a marker for prostatic cancer and hepatic metastasis from breast and colon.

Reference Values

Normal
Men: 7–47 U/L (0.12–1.80 µkat/L) Women: 5–25 U/L (0.08–0.42 µkat/L) Values higher in newborns and in the first 3–6 months. Values in adult males are 25% higher than in females. Values vary with method (check with your laboratory).

Procedure

1. Obtain a 5-mL venous blood sample. Serum is used.
2. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications

1. Increased ?GT levels are associated with the following conditions:
   a. Liver diseases
      1. Hepatitis (acute and chronic)
      2. Cirrhosis (obstructive and familial)
      3. Liver metastasis and carcinoma
      4. Cholestasis (especially during or following pregnancy)
      5. Chronic alcoholic liver disease, alcoholism
      6. Infectious mononucleosis
   b. ?GT levels are also increased in the following conditions:
      1. Pancreatitis
      2. Carcinoma of prostate
      3. Carcinoma of breast and lung
      4. Systemic lupus erythematosus
      5. Glycogen storage disease
   c. In MI, ?GT is usually normal. However, if there is an increase, it occurs about 4 days after MI and probably implies liver damage secondary to cardiac insufficiency.
   d. Hyperthyroidism

2. Decreased ?GT levels are found in hypothyroidism.
3. ?GT values are normal in bone disorders, bone growth, pregnancy, skeletal muscle disease, strenuous exercise, and renal failure. Children and adolescents are normal.

Interfering Factors

1. Various drugs, (eg, phenothiazines and barbiturates) affect test outcomes (see Appendix J).
2. Alcohol (ethanol)

Interventions

Pretest Patient Care

1. Explain test purpose and blood-drawing procedure. No alcohol is allowed before the test.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume normal activities.
2. Interpret test results and monitor as appropriate for liver, pancreatic, or thyroid disease and/or cancer recurrence.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Homocysteine (tHcy)

Homocysteine (tHcy) is an amino acid resulting from the synthesis of cysteine from methionine and enzyme reaction of cobalamine and folate. Large quantities of homocysteine are excreted and assimilated in the blood plasma of patients with
homocystinemia associated with:

1. Increased risk for vascular disease
2. Increased risk for venous thrombosis
3. Elevated homocysteine with a direct toxic effect on endothelium
4. Elevated in folic acid deficiency and B$_{12}$ deficiency. Folic acid deficiency is characterized by elevated plasma homocysteine; folic acid supplementation reduces plasma homocysteine. Elevated plasma homocysteine levels due to aberrant vitamin B$_{12}$ respond favorably to vitamin B$_{12}$ supplementation.
5. Increased risk for pregnancy complications and neural tube defects

This test measures the blood plasma level of homocysteine. It is useful for diagnosing individuals with potential increased risk factors for coronary artery disease and thromboses, for providing a functional assay for folic acid deficiency, and for diagnosing homocystinemia. Homocysteine is retained by persons with reduced renal function. See Chart 6.3 for testing guidelines.

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**Chart 6.3 Homocysteine Testing**

**Reasons to Test for Homocysteine**

Unexplained anemia

Peripheral neuropathy or myelopathy

Recurrent spontaneous abortions or infertility

Delayed development or failure to thrive in infants

**Who to Test**

Elderly people (>75 years of age)

Vegetarians who are not taking vitamin B$_{12}$ supplement

Patients using drugs that interfere with folate status (eg, antiepileptics, methotrexate)

**How Often to Test**

Measured every 3–5 years

In newborns at 3–5 days

In patients <40 years of age who have CVD to exclude homocystinuria

In patients who are at high risk for CVD every 3–5 years

**Footnote**

Source: Clinical Laboratory News, 2002

**Reference Values**

*Normal* 4–17 µmol/L or 0.54–2.30 mg/L for fasting specimens

**Procedure**

1. Obtain a venous blood sample. Serum or heparinized plasma is needed. Fasting is necessary.
2. Observe standard precautions. Place specimen in a biohazard bag.
3. Place on ice immediately after drawing. Centrifuge immediately and freeze within 1 hour of collection.

**Clinical Implications** Increased or elevated homocysteine levels occur in the following conditions:

1. Folic acid deficiency
2. Abnormal vitamin B$_{12}$ metabolism and deficiency
3. Homocystinuria

**Clinical Alert**

Homocysteine values and their relation to CAD are still being investigated. The methionine load test is also currently investigative and has not yet been approved as a routine test.

**Interfering Factors**
1. Penicillamine reduces plasma levels of homocysteine.
2. Nitrous oxide, methotrexate deficiency, and azauridine increase plasma levels of homocysteine.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and blood-drawing procedure.
2. Remember that the test requires fasting.
3. Evaluate renal function in patients with homocystinuria.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Allow the patient to eat and drink after blood is drawn.
2. Interpret test results and counsel appropriately.
3. Evaluate for other cardiovascular risk factors, compare test results, and monitor appropriately. Promote lifestyle changes accordingly.
4. Monitor for folic acid or vitamin B12 deficiency and provide supplements as needed.
5. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**a_1-Antitrypsin (AAT)**

a_1-Antitrypsin is a protein produced by the liver that inhibits the protease released into body fluids by dying cells. This protein deficiency is associated with pulmonary emphysema and liver disease, both at an early age. Human serum contains at least three inhibitors of protease. Two of the best known are a_1-antitrypsin and a_2-macroglobulin. Total antitrypsin levels in blood are composed of about 90% AAT and 10% a_2-macroglobulins.

This is a nonspecific method to diagnose inflammation, severe infection, and necrosis. AAT measurement is important for diagnosing respiratory disease and cirrhosis of the liver because of its direct relation to pulmonary and other metabolic disorders. Pulmonary problems such as emphysema occur when antitrypsin-deficient persons are unable to ward off the action of endoproteases. Those who are deficient in AAT develop emphysema at a much earlier age than do other emphysema patients.

**Reference Values**

*Normal (by rate nephelometry)* 110–200 mg/dL or 1.1–2.0 g/L. If result is <125 mg/dL (<1.25 g/L), phenotype should be determined to confirm homozygous and heterozygous deficiencies.

**Procedure**

1. Obtain a 7-mL serum sample. Use a red-topped tube.
2. Observe standard precautions. Place specimen in a biohazard bag.

**Clinical Implications**

1. Interpretation of AAT levels is based on the following:
   a. High levels are generally found in normal persons.
   b. Intermediate levels are found in persons with a predisposition to pulmonary emphysema.
   c. Low levels are found in persons with obstructive pulmonary disease and in children with cirrhosis of the liver.
2. *Increased AAT levels* occur in the following conditions:
   a. Acute and chronic inflammatory disorders
   b. After injections of typhoid vaccine
   c. Cancer
   d. Thyroid infections
   e. Oral contraceptive use
   f. Stress syndrome
   g. Hematologic abnormalities
3. *Decreased AAT levels* are associated with these progressive diseases:
   a. Adult, early-onset, chronic pulmonary emphysema
   b. Liver cirrhosis in infants (neonatal hepatitis)
   c. Pulmonary disease
   d. Severe hepatic damage
   e. Nephrotic syndrome
   f. Malnutrition

**Clinical Alert**

Patients with serum levels <70 mg/dL (<0.70 g/L) are likely to have a homozygous deficiency and are at risk for early lung disease.

**Interfering Factors** a_1-Antitrypsin in an acute-phase reactant, and any inflammatory process will elevate serum levels.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. Fasting is required if the patient's history shows elevated cholesterol and/or triglyceride levels.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and counsel appropriately. Advise patients with decreased levels to avoid smoking and, if possible, occupational hazards such as dust, fumes, and other respiratory pollutants.
2. Be aware that because AAT deficiencies are inherited, genetic counseling may be indicated. Follow-up AAT phenotype testing can be performed on family members to determine the homozygous or heterozygous nature of the deficiency.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

DRUG MONITORING

Drug monitoring is used for therapeutic management and toxicology. Blood, urine, and gastric contents are the most common specimens used for measuring drug levels. When a combination of drugs has been used, the levels of all drugs should be obtained. Toxic drug levels are necessary to evaluate substance abuse and intentional or accidental overdose so immediate interventions can be initiated (eg, enhanced diuresis, hemodialysis, adding soybean oil to bathwater) to avoid death or disabling conditions. Situations in which toxic drug levels may be measured include accidental overdose of opiates; suspected poisoning as in homicide or suicide; medical emergencies where person arrives at ED in coma or altered state of consciousness; suspected date rape situations where a drug may have been used; accidental poisoning in children; child abuse cases where results may be used to determine parental custody; illicit drug use such as heroine, cocaine, or opiates; suspected solvent vapor abuse such as sniffing of paints; exposure to anticoagulants as in rodent poisons; exposure to iron and heavy metals; intentional ingestion as in chronic overdose secondary to chronic pain; and in vehicle accident cases where alcohol is the most commonly abused. See Appendix L for discussion of these drug investigation studies.

Therapeutic Drug Management

Therapeutic drug management (formerly called therapeutic drug monitoring) is a reliable and practical approach to improving drug therapy in both instruction and maintenance in individual patients. Determination of drug levels is especially important when the potential for drug toxicity is significant or when an inadequate or undesirable response follows the use of a standard dose. Therapeutic drug management provides an easier and more rapid estimation of appropriate drug-dosage requirements than does observation of the drug effects themselves. For some drugs, monitoring is routinely useful (eg, digoxin); for others, it can be helpful in certain situations (eg, antibiotics). The plasma level of drugs needed to control the patient's symptoms is called the therapeutic concentration; at a steady state, the rate of drug administration is equal to the rate of drug elimination, and the concentration of the drug remains constant. Monitoring at intervals minimizes the possibility of the development of dose-related side effects. If single-drug therapy is not effective, therapeutic monitoring allows the clinician to select supplementary medication and monitor its effect on the primary drug. It should be noted that for many medications, therapeutic serum levels have not been established. Therapy is guided by clinical response and adverse reactions of the medications. Appropriate therapy may be monitored or managed by other methods, including outcome of liver function tests, CBC with differential, platelet counts, serum electrolytes, serum albumin, and renal function tests.

Indications for Testing

1. Verify correct drug dosage and level; drug source, dose, or regimen is changed
2. When noncompliance (nonadherence) is suspected, and patient motivation to maintain medication is poor

Reference Values

Normal See Table 6.13 for therapeutic maintenance and toxic levels.

<table>
<thead>
<tr>
<th>Name of Drug</th>
<th>Therapeutic Level</th>
<th>Toxic Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetaminophen</td>
<td>10–20 µg/mL, based on relief of symptoms</td>
<td>Should be &gt;200 µg/mL at 4 hours after ingestion or &gt;50 µg/mL at 12 hours after ingestion</td>
</tr>
<tr>
<td>Amikacin</td>
<td>Infections: 20–30 µg/mL (34–52 µmol/L)</td>
<td>Peak: &gt;35 µg/mL (&gt;60 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>Serious infections: 20–25 µg/mL</td>
<td>Trough: &gt;10 µg/mL (&gt;17 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>UTI: 15–20 µg/mL</td>
<td></td>
</tr>
<tr>
<td></td>
<td><em>Trough</em></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serious infection: 1–4 µg/mL (2–7 µmol/L)</td>
<td>Life-threatening infections: 4–8 µg/mL (7–14 µmol/L)</td>
</tr>
<tr>
<td>Amiodarone (see Fig. 6.4)</td>
<td>0.5–2.5 mg/L (0.8–3.9 µmol/L)</td>
<td>&gt;3.5 mg/L</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>Meningitis: <em>Peak</em>: 15–25 µg/mL</td>
<td>&gt;40 µg/mL</td>
</tr>
<tr>
<td></td>
<td><em>Trough</em>: 5–15 µg/mL</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Other infections: <em>Peak</em>: 10–20 µg/mL</td>
<td></td>
</tr>
<tr>
<td></td>
<td><em>Trough</em>: 5–10 µg/mL</td>
<td></td>
</tr>
<tr>
<td>Desipramine</td>
<td>50–300 ng/mL</td>
<td>Possibly toxic: &gt;300 ng/mL</td>
</tr>
<tr>
<td>Digoxin</td>
<td>CHF: 0.8–2 ng/mL (1.0–2.6 nmol/L)</td>
<td>Toxic: &gt;1000 ng/mL</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&gt;2.0 ng/mL (&gt;2.6 nmol/L)</td>
</tr>
<tr>
<td>Drug</td>
<td>Normal Serum Levels</td>
<td>Toxic Levels</td>
</tr>
<tr>
<td>----------------------</td>
<td>----------------------------------------------</td>
<td>----------------------------------------</td>
</tr>
<tr>
<td>Arrhythmias:</td>
<td>1.5–2.5 ng/mL (2.0–3.2 mmol/L)</td>
<td></td>
</tr>
<tr>
<td>Adverse reactions:</td>
<td>Nausea, vomiting, anorexia, green/yellow</td>
<td></td>
</tr>
<tr>
<td></td>
<td>visual distortion (commonly reported symptoms in patients requiring hospitalization)</td>
<td></td>
</tr>
<tr>
<td>Digitoxin (see Fig. 6.5)</td>
<td>18–35 ng/mL (24–46 nmol/L)</td>
<td>&gt;35 ng/mL (46 nmol/L)</td>
</tr>
<tr>
<td>Disopyramide (Norpace)</td>
<td>Atrial arrhythmias: 2.8–3.2 µg/mL (8.3–9.4</td>
<td>&gt;7 µg/mL (2.1 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>µmol/L</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ventricular: 3.3–7.5 µg/mL (9.7–22.2 µmol/L)</td>
<td></td>
</tr>
<tr>
<td>Epinephrine *</td>
<td>31–95 pg/mL</td>
<td>A toxic level has not been established</td>
</tr>
<tr>
<td>Ethasuximide (Zarontin)</td>
<td>40–100 µg/mL (284–710 µmol/L)</td>
<td>Panic value: &gt;150 µg/mL (1062 µmol/L)</td>
</tr>
<tr>
<td>Ethchlorvynol</td>
<td>2–9 µg/mL or 14–55 µmol/L</td>
<td>A toxic level has not been established</td>
</tr>
<tr>
<td>Flecainide</td>
<td>0.2–1 µg/mL</td>
<td>&gt;1.0 µg/mL</td>
</tr>
<tr>
<td>Flucytosine (Ancobon)</td>
<td>25–100 µg/mL</td>
<td></td>
</tr>
<tr>
<td>Fluoxetine</td>
<td>100–800 ng/mL (289–2312 nmol/L)</td>
<td>Fluoxetine + norfluoxetine &gt;2000 ng/mL (5780 nmol/L)</td>
</tr>
<tr>
<td></td>
<td>Norfluoxetine 100–600 ng/mL (289–1735 nmol/L)</td>
<td></td>
</tr>
<tr>
<td>Flurazepam *</td>
<td>0–4 ng/mL (0–9 nmol/L)</td>
<td>&gt;200 ng/mL (578 nmol/L)</td>
</tr>
<tr>
<td>Fosphenytoin</td>
<td>10–20 µg/mL</td>
<td>30–50 µg/mL</td>
</tr>
<tr>
<td></td>
<td>Lethal: &gt;100 µg/mL</td>
<td></td>
</tr>
<tr>
<td>Gabapentin *</td>
<td>Minimum effective serum level: 2 µg/mL</td>
<td>25 µg/mL</td>
</tr>
<tr>
<td>Gentamicin (Garamycin)</td>
<td>Peak:</td>
<td>Toxic level is based on panic or life-threatening values.</td>
</tr>
<tr>
<td></td>
<td>Serious infections: 6–8 µg/mL (12–17 µmol/L)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Life-threatening: 8–10 µg/mL (17–21 µmol/L)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>UTI: 4–6 µg/mL (8–12 µmol/L)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Trough: 0.5–1 µg/mL (1–2 µmol/L)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Life-threatening: 1–2 µg/mL (2–4 µmol/L)</td>
<td></td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>20–70 µg/mL, based on symptom relief</td>
<td></td>
</tr>
<tr>
<td>Lidocaine</td>
<td>1.5–5.0 µg/mL (6.14–21.4 µmol/L)</td>
<td>Potentially toxic: &gt;6 µg/mL (25 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>Life-threatening: &gt;2 µg/mL (2–4 µmol/L)</td>
<td></td>
</tr>
<tr>
<td>Lithium</td>
<td>Acute mania: 0.6–1.2 mEq/L (0.6–1.2 mmol/L)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Protection against future episodes in patients with bipolar disorder:</td>
<td></td>
</tr>
<tr>
<td>Lorazepam</td>
<td>50–240 ng/mL</td>
<td></td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Depends on low or high dose therapy</td>
<td>Low dose toxic therapy: &gt;9.1 ng/mL</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High dose toxic therapy: &gt;450 ng/mL</td>
</tr>
<tr>
<td>Mexiletine</td>
<td>0.5 µg/mL</td>
<td>Potentially toxic: &gt;9.1 ng/mL (20 mmol/L)</td>
</tr>
<tr>
<td>Oxcarbazepine *</td>
<td>Active metabolite (10-hydrox-carbazepine)</td>
<td>&gt;2 µg/mL (&gt;9 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>(Trileptal)</td>
<td></td>
</tr>
<tr>
<td>For trigeminal neuralgia</td>
<td>50–110 µmol/L; therapeutic serum levels have not been established for treatment of epilepsy.</td>
<td>Toxic levels not established</td>
</tr>
<tr>
<td>Procaainamide</td>
<td>4–10 µg/mL (17–42 µmol/L)</td>
<td>&gt;14 µg/mL (&gt;60 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>NAPA: 10–30 µg/mL (42–127 µmol/L)</td>
<td>Combined: &gt;30 µg/mL (&gt;127 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>Combined: &gt;30 µg/mL (127 µmol/L)</td>
<td></td>
</tr>
<tr>
<td>Phenytoin Children and adults:</td>
<td>Total phenytoin: 10–20 µg/mL (40–70 µmol/L)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Neonates: 8–15 µg/mL</td>
<td>Lethal: &gt;100 µg/mL (400 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>Free phenytoin: 1–2.0 µg/mL (4–8 µmol/L)</td>
<td></td>
</tr>
<tr>
<td>Salicylates</td>
<td>Antiplatelet, antipyresis, analgesia: 100 µg/mL</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Anti-inflammatory: 150–300 µg/mL</td>
<td></td>
</tr>
<tr>
<td>Temazepam</td>
<td>26 ng/mL after 24 hours</td>
<td>Information not available</td>
</tr>
<tr>
<td>Theophylline</td>
<td>Asthma: 10–20 µg/mL (56–111 µmol/L)</td>
<td>&gt;20 µg/mL (&gt;111 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>Neonatal apnea: 6–13 µg/mL (33–72 µmol/L)</td>
<td>&gt;10 µg/mL (&gt;56 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>Pregnancy: 3–12 µg/mL (17–67 µmol/L)</td>
<td>&gt;30 µg/mL (168 µmol/L)</td>
</tr>
<tr>
<td>Drug</td>
<td>Hypnotic: 1–5 µg/mL</td>
<td>&gt;10 µg/mL</td>
</tr>
<tr>
<td>--------------</td>
<td>---------------------</td>
<td>----------</td>
</tr>
<tr>
<td>Thioental</td>
<td>Anesthesia: 7–130 µg/mL</td>
<td>Coma: 30–100 µg/mL</td>
</tr>
<tr>
<td>Valproic acid</td>
<td>50–120 µg/mL (wide therapeutic range)</td>
<td>&gt;200 µg/mL</td>
</tr>
<tr>
<td>Vancomycin</td>
<td>Peak: 25–40 µg/mL (17–27 µmol/L)</td>
<td>&gt;80 µg/mL (&gt;54 µmol/L)</td>
</tr>
<tr>
<td></td>
<td>Trough: 5–10 µg/mL (3.4–6.8 µmol/L)</td>
<td></td>
</tr>
</tbody>
</table>

*Therapeutic serum levels have not been established for epilepsy.

---


**Blood, Saliva, and Breath Alcohol Content (BAC; Ethanol [Ethyl Alcohol, ETOH])**

Ethanol is absorbed rapidly from the GI tract, with peak blood levels usually occurring within 40 to 70 minutes of ingestion on an empty stomach. Food in the stomach decreases alcohol absorption. Ethanol is metabolized by the liver to acetaldehyde. Once peak blood ethanol levels are reached, disappearance is linear; a 70-kg man metabolizes 7 to 10 g/h of alcohol (15 ± 5 mg/dL/h). Symptoms of intoxication in the presence of low alcohol levels could indicate a serious acute medical problem requiring immediate attention.

Quantitation of alcohol level may be performed for medical or legal purposes, to diagnose alcohol intoxication, and to determine appropriate therapy. Alcohol level must be tested as a possible cause of unknown coma because alcohol intoxication mimics diabetic coma, cerebral trauma, and drug overdose. This test is also used to screen for alcoholism and to monitor ethanol treatment for methanol intoxication.

**Reference Values**

**Normal**

Negative: no alcohol detected <10 mg/dL or <2 mmol/L is considered negative; <20 mg/dL or <4.34 mmol/L is considered negative for the U.S. Department of Transportation (DOT); >40 mg/dL or >8.68 mmol/L is considered positive for the U.S. DOT; >80 mg/dL or >17.4 mmol/L is positive under most state drunk driving laws.

**Procedure**

1. Obtain a 5-mL venous blood sample from the arm in living persons. From dead persons, take samples from the aorta. Observe standard precautions.
   a. Use a non–alcohol-based solution (eg, povidone-iodine) for cleansing the venipuncture site.
   b. Sodium fluoride or oxalate anticoagulant is recommended. Serum can also be used.
   c. Keep blood sample tightly stoppered. Do not open.

2. A 20-mL sample of urine or gastric contents can also be used. Place specimen in a biohazard bag.

3. A breath analyzer measures ethanol content at the end of expiration following a deep inspiration. (See Appendix K for information on breath alcohol analyzers.)

**Clinical Implications**

1. At levels of 50 to 100 mg/dL (10.8–21.7 mmol/L), certain signs and symptoms are reported (eg, flushing, slowing of reflexes, impaired visual acuity).
2. At levels >100 mg/dL (>21.7 mmol/L), CNS depression is reported. In many states, this is the cutoff level for driving under the influence of alcohol.
3. Blood levels >300 mg/dL (>64.8 mmol/L) are associated with coma.
4. Death has been reported at levels >400 mg/dL (>86.4 mmol/L).
5. Properly collected urine samples will have an alcohol content similar to that of blood. Saliva samples will have an
alcohol content 1.2 times that of blood.

Interfering Factors

1. Increased blood ketones, as in diabetic ketoacidosis, can falsely elevate blood or breath test results.
2. Ingestion of other alcohols, such as isopropanol or methanol, may affect results.

Clinical Alert

1. Panic value is >300 mg/dL (>64.8 mmol/L). Report and initiate overdose treatment at once.
2. Symptoms of intoxication in the presence of low blood alcohol could indicate a serious medical problem requiring immediate medical attention.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. Proper collection, handling, and storage of the blood alcohol specimen is essential when the question of sobriety is raised.
2. Advise patient of legal rights in cases involving question of sobriety.
3. A witnessed, signed consent form may have to be obtained.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and monitor as appropriate for toxic levels.
2. If alcohol levels are high, initiate treatment at once.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Lipoprotein Tests/Lipoprotein Profiles

Lipoprotein measurements are diagnostic indicators for hyperlipidemia and hypolipidemia. Hyperlipidemia is classified as types I, Iia, Iib, III, IV, and V. Lipids are fatty substances made up of cholesterol, cholesterol esters (liquid compounds), triglycerides, nonesterized fatty acids, and phospholipids. Lipoproteins are unique plasma proteins that transport otherwise insoluble lipids. They are categorized as chylomicrons, ß-lipoproteins (low-density lipoproteins [LDLs]), pre-ß-lipoproteins (very-low-density lipoproteins [VLDLs]), and a-lipoproteins (high-density lipoproteins [HDLs]). Apolipoprotein A is mainly composed of HDL, chylomicrons, and VLDL. Apolipoprotein B is the main component of LDL. Lipids provide energy for metabolism, serve as precursors of steroid hormones (adrenals, ovaries, testes) and bile acids, and play an important role in cell membrane development. A lipid profile usually includes cholesterol, triglycerides, LDL, and HDL levels.

Cholesterol

Cholesterol testing evaluates the risk for artherosclerosis, myocardial occlusion, and coronary arterial occlusion. Cholesterol relates to coronary heart disease (CHD) and is an important screening test for heart disease. It is part of the lipid profiles. Elevated cholesterol levels are a major component in the hereditary hyperlipoproteinemias. Cholesterol determinations are also frequently a part of thyroid function, liver function, renal function, and diabetes mellitus studies. It is also used to monitor effectiveness of diet, medications, lifestyle changes (eg, exercise), and stress management.

Reference Values

Normal Normal values vary with age, diet, sex, and geographic or cultural region. Adults, fasting: Desirable level: 140–199 mg/dL or 3.63–5.15 mmol/L Borderline high: 200–239 mg/dL or 5.18–6.19 mmol/L High: >240 mg/dL or >6.20 mmol/L Children and adolescents (12–18 years): Desirable level: <170 mg/dL or <4.39 mmol/L Borderline high: 170–199 mg/dL or 4.40–5.16 mmol/L High: >200 mg/dL or >5.18 mmol/L

Procedure

1. Obtain a 5-mL venous blood sample. Fasting is required. Serum is needed.
2. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications

1. Total blood cholesterol levels are the basis for classifying CHD risk.
   a. Levels >240 mg/dL or >6.20 mmol/L are considered high and should include follow-up lipoprotein analysis. Borderline high levels (200–239 mg/dL or 5.18–6.19 mmol/L) in the presence of CHD or two other CHD risk factors should also include lipoprotein analysis/profiles.
   b. CHD risk factors include male gender, family history, and premature CHD (MI or sudden death before age 55 years in a parent or sibling), smoking (>10 cigarettes per day), hypertension, low HDL cholesterol levels (<35 mg/dL or <0.91 mmol/L confirmed by repeat measurement), diabetes mellitus, history of definite cerebrovascular or occlusive peripheral vascular disease, and severe obesity (>30% overweight).
   c. In public screening programs, all patients with cholesterol levels >200 mg/dL or >5.18 mmol/L should be referred to their physicians for further evaluation. Before initiating any therapy, the level should be retested.
2. Elevated cholesterol levels (hypercholesterolemia) occur in the following conditions:
   a. Type II familial hypercholesterolemia
   b. Hyperlipoproteinemia types I, IV, and V
   c. Cholestasis
d. Hepatocellular disease, biliary cirrhosis
e. Nephrotic syndrome glomerulonephritis
f. Chronic renal failure
g. Pancreatic and prostatic malignant neoplasms
h. Hypothyroidism
i. Poorly controlled diabetes mellitus
j. Alcoholism
k. Glycogen storage disease (von Gierke’s disease)
l. Werner’s syndrome
m. Diet high in cholesterol and fats (“dietary affluence”)
n. Obesity

3. Decreased cholesterol levels (hypocholesterolemia) occur in the following conditions:
a. Hypo-a-lipoproteinemia
b. Severe hepatocellular disease
c. Myeloproliferative diseases
d. Hyperthyroidism
e. Malabsorption syndrome, malnutrition
f. Megaloblastic or sideroblastic anemia (chronic anemias)
g. Severe burns, inflammation
h. Conditions of acute illness, infection
i. Chorionic obstructive lung disease
j. Mental retardation

Interfering Factors
1. Estrogens decrease plasma cholesterol levels; pregnancy increases these levels.
2. Certain drugs increase or decrease cholesterol levels.
3. Seasonal variations in cholesterol levels have been observed; levels are higher in fall and winter and lower in spring and summer.
4. Positional variations occur; levels are lower when sitting versus standing and lower when recumbent versus sitting.
5. Plasma (EDTA) values are 10% lower than serum.

Interventions

Pretest Patient Care
1. Explain test purpose and procedure. An overnight fast before testing is recommended, although nonfasting specimens may be taken. Pretest, a normal diet should be consumed for 7 days. The patient should abstain from alcohol for 48 hours before testing. Prolonged fasting with ketosis increases values.
2. Document drugs the patient is taking.
3. Encourage the patient to relax.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test results and counsel appropriately. Cholesterol levels are influenced by heredity, diet, body weight, and physical activity. Some lifestyle changes may be necessary to reduce elevated levels.
2. Remember that cholesterol levels >200 mg/dL (or >5.18 mmol/L) should be retested and the results averaged. If the two results differ by >10%, a third test should be done.
3. Be aware that once hyperlipidemia has been established, the diet should be lower in animal fats and should replace saturated fats with polyunsaturated fats. Fruits, vegetables (especially greens), and whole-grain products should be increased. Patients with diabetes, as well as others, should seek counsel from a dietitian regarding diet management if necessary. Therapy for hyperlipidemia should always begin with diet modification.
4. Remember that the American Heart Association and National Cholesterol Education Programs have excellent resources for providing diet and lifestyle management information.
5. Be aware that at least 6 months of dietary therapy should be tried before initiating cholesterol-reducing drug therapy.
6. Perform a comprehensive lipoprotein analysis if cholesterol levels are not lowered within 6 months after start of therapy.

Clinical Alert
1. Cholesterol measurement should not be done immediately after MI. A 3-month wait is suggested.
2. >300 mg/dL or >7.8 mmol/L: there is a strong relationship to coronary heart disease, but only a fraction of those with CAD have cholesterol increased.

High-Density Lipoprotein Cholesterol (HDL-C)

HDL-C is a class of lipoproteins produced by the liver and intestines. HDL is composed of phospholipids and one or two apolipoproteins. It plays a role in the metabolism of the other lipoproteins and in cholesterol transport from peripheral tissues to the liver. LDL and HDL may combine to maintain cellular cholesterol balance through the mechanism of LDL moving cholesterol into the arteries and HDL removing it from the arteries. Decreased HDL levels are atherogenic, whereas elevated HDL levels protect against atherosclerosis by removing cholesterol from vessel walls and transporting it to the liver where it is removed from the body. There is a strong relationship of HDL cholesterol and CAD.
HDL-C, the good cholesterol, is used to assess CAD risk and monitor persons with known low HDL levels. HDL-C levels are inversely proportional to CHD risk and are a primary independent risk factor. When a slightly increased cholesterol is due to high HDL therapy, this is not indicated.

Reference Values

**Normal**
- Men: 35–65 mg/dL or 0.91–1.68 mmol/L
- Women: 35–80 mg/dL or 0.91–2.07 mmol/L

- <25 mg/dL or <0.65 mmol/L: CHD risk at dangerous level 2 times the risk
- 26–35 mg/dL or 0.67–0.91 mmol/L: average CHD risk: 1.5 times
- 36–44 mg/dL or 0.93–1.14 mmol/L: moderate CHD risk: 1.2 times the risk
- 45–59 mg/dL or 1.16–1.53 mmol/L: below-average CHD risk
- >75 mg/dL or >1.94 mmol/L of HDL: no risk (associated with longevity)

**Procedure**
1. Obtain a 5-mL venous blood sample. Fasting is necessary. The HDL is precipitated out from the total cholesterol for analysis.
2. Calculate a cholesterol/HDL-C ration from these values.

**Clinical Implications**
1. **Increased HDL-C values** occur in the following conditions:
   a. Familial hyper-a-lipoproteinemia (HDL excess)
   b. Chronic liver disease (cirrhosis, alcoholism, hepatitis)
   c. Long-term aerobic or vigorous exercise
2. **Decreased HDL-C values** are associated with increased risk for CHD and premature CHD and occur in the following conditions:
   a. Familial hypo-a-lipoproteinemia (Tangier disease), Apo C-III deficiency
   b. a-ß-Lipoproteinemia
   c. Hypertriglyceridemia (familial)
   d. Poorly controlled diabetes mellitus
   e. Hepatocellular diseases
   f. Cholestasis
   g. Chronic renal failure, uremia, nephrotic syndrome
   h. In the United States, 3% of men have low HDL levels for unknown reasons, even though cholesterol and triglyceride values are normal, and they are at risk for premature CAD.

**Interfering Factors**
1. Increased HDL level is associated with estrogen therapy, moderate intake of alcohol and other drugs (especially androgenic and related steroids), and insulin therapy.
2. Decreased HDL levels are associated with the following:
   a. Certain drugs such as steroids, antihypertensive agents, diuretics, beta blockers, triglycerides, and thiazides
   b. Stress and recent illness
   c. Starvation and anorexia
   d. Obesity, lack of exercise
   e. Smoking
   f. Hypertriglyceridemia (>400 mg/dL or >10.36 mmol/L) (retest making sure the patient is properly fasting)

**Interventions**

**Pretest Patient Care**
1. Explain test purpose. A 9–12 hour fast is recommended. Alcohol should not be consumed for at least 24 hours before test.
2. Ensure that patient is on a stable diet for 3 weeks.
3. If possible, withhold all medication for at least 24 hours before testing. Check with physician.
4. Encourage relaxation.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
1. Interpret test results and counsel appropriately (see cholesterol patient aftercare, page 423).
2. Remember that low HDL levels can be raised by diet management, exercise, weight loss, and smoking cessation.
   Many resources are available through the American Heart Association and other organizations.
3. Drug therapy may be necessary if other methods fail to raise HDL levels.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Clinical Alert**
Cholesterol and HDL-C levels should not be done immediately after MI. A 3-month wait is suggested.

**Clinical Alert**
The cholesterol/HDL ratio provides more information than does either value alone. The higher the cholesterol/HDL ratio, the greater the risk for developing atherosclerosis. This ratio should be reported with total cholesterol values, along with the % HDL-C.
Very-Low-Density Lipoprotein (VLDL); Low-Density Lipoprotein (LDL)

Sixty to 70% of the total serum cholesterol is present in the LDL. LDLs are the cholesterol-rich remnants of the VLDL lipid transport vehicle. Because LDL has a longer half-life (3–4 days) than its precursor VLDL, LDL is more prevalent in the blood. It is mainly catabolized in the liver and possibly in nonhepatic cells as well. The VLDLs are major carriers of triglycerides. Degradation of VLDL is a major source of LDL. Circulating fatty acids form triglycerides in the liver, and these are packaged with apoprotein and cholesterol to be exported into the blood as VLDLs.

This test is specifically done to determine CHD risk. LDL, "the bad cholesterol," is closely associated with increased incidence of atherosclerosis and CHD. The test of choice is LDL because it has a longer half-life and it is easier to measure.

Reference Values

**Normal**
- **Adults:** Desirable: <130 mg/dL or <3.4 mmol/L Borderline high-risk: 140–159 mg/dL or 3.4–4.1 mmol/L High-risk: >160 mg/dL or >4.1 mmol/L Children and adolescents: Desirable: <110 mg/dL or <2.8 mmol/L Borderline high-risk: 110–129 mg/dL or 2.8–3.4 mmol/L High-risk: >130 mg/dL or >3.4 mmol/L

**Procedure**

1. Use the following equation for VLDL calculated (estimation): triglycerides divided by 5.
2. Calculate LDL cholesterol levels by using the Friedwald's formula:
   \[
   \text{LDL cholesterol} = \text{total cholesterol} - \text{HDL cholesterol} - \frac{(\text{triglycerides})}{5}
   \]
3. Remember that the formula is valid only if the cholesterol and triglyceride values are from a fasting specimen and the triglyceride value is >400 mg/dL or >10.4 mmol/L.
4. Lipoprotein analysis measures fasting levels of total cholesterol, total triglycerides, and HDL cholesterol. Calculate LDL cholesterol from these values.
5. Remember that there is a nondirect test for LDH that may be ordered if triglycerides are >400 mg/dL or >10.4 mmol/L.

**Clinical Implications**

1. **Increased LDL levels** are caused by the following conditions:
   a. Familial type 2 hyperlipidemia, familial hypercholesterolemia
   b. Secondary causes include the following:
      1. Diet high in cholesterol and saturated fat
      2. Hyperlipidemia secondary to hypothyroidism
      3. Nephrotic syndrome
      4. Multiple myeloma and other dysglobulinemias
      5. Hepatic obstruction or disease
      6. Anorexia nervosa
      7. Diabetes mellitus
      8. Chronic renal failure
      9. Porphyria
      10. Premature CHD
2. **Decreased LDL levels** occur in the following conditions:
   a. Hypolipoproteinemia
   b. Tangier disease
   c. Type I hyperlipidemia
   d. Apo C-II deficiency
   e. Hyperthyroidism
   f. Chronic anemias
   g. Severe hepatocellular disease
   h. Reye's syndrome
   i. Acute stress (burns, illness)
   j. Inflammatory joint disease
   k. Chronic pulmonary disease

**Interfering Factors**

1. Increased LDLs are associated with pregnancy and certain drugs such as steroids, progestins, and androgens (see Appendix J).
2. Not fasting may cause false elevation.
3. Decreased LDLs are found in women taking oral estrogen therapy.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose. A 9–12 hour fast is recommended. Alcohol should not be consumed for at least 24 hours before test.
2. Remember that patient should ideally be on a stable diet for 3 weeks.
3. If possible, withhold all medication for at least 24 hours before testing. Check with physician.
4. Encourage relaxation.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
Posttest Patient Aftercare

1. Interpret test results and counsel appropriately about results and need for further testing.
2. If patient has high LDH levels, repeat the test in 2 to 8 weeks and average the values to establish an accurate baseline from which to devise a treatment plan (Table 6.14).

<table>
<thead>
<tr>
<th>Table 6.14 Stages of Treatment for High LDH Levels</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Initiation Level</strong></td>
</tr>
<tr>
<td><strong>DIETARY TREATMENT</strong></td>
</tr>
<tr>
<td>Without CHD or two other risk factors</td>
</tr>
<tr>
<td>With CHD or two other risk factors</td>
</tr>
<tr>
<td><strong>DRUG TREATMENT</strong></td>
</tr>
<tr>
<td>Without CHD or two other risk factors</td>
</tr>
<tr>
<td>With CHD or two other risk factors</td>
</tr>
</tbody>
</table>

3. A comprehensive history and physical exam, together with analysis of test results, determines whether high LDL cholesterol is secondary to another disease or drug or is the result of a familial lipid disorder. The patient's total coronary risk profile, clinical status, age, and gender are considered when prescribing a cholesterol-lowering treatment program (see Table 6.15 for LDL-C/HDL-C ratios).

<table>
<thead>
<tr>
<th>Table 6.15 LDL-C/HDL-C Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Risk Level</strong></td>
</tr>
<tr>
<td>Low</td>
</tr>
<tr>
<td>Average</td>
</tr>
<tr>
<td>Moderate</td>
</tr>
<tr>
<td>High</td>
</tr>
</tbody>
</table>

Clinical Alert

Another method for assessing CAD/CHD risk is by calculating the LDH/HDL ratio (LDL-C–HDL-C).

NOTE

Patients need a lower initiation level and goal if they are at high risk because of existing CHD or any two of the following risk factors: male gender, family history of premature CHD, smoking, hypertension, low HDL cholesterol, diabetes mellitus, cerebrovascular or peripheral vascular disease, or severe obesity.

Apolipoprotein A and B (Apo A-I, Apo B)

Hypolipoproteins/apolipoproteins are surface proteins of lipoprotein particles and are important in the study of atherosclerosis. Apolipoprotein A is the main (90%) component of HDL. Apolipoprotein B is the main component of LDL and VLDL and is important in regulating cholesterol synthesis and metabolism.

This test is used to evaluate the risk for CAD. APO A-I deficiencies are often associated with premature cardiovascular disease. Apo B plays an important role in LDL catabolism. The ratio of Apo A to Apo B correlates more closely with increased risk for CAD than do cholesterol levels or the LDL/HDL ratio. The lower the ratio, the higher the risk.

Reference Values

**Normal** Apo A-I: Men: 90–155 mg/dL (0.90–1.55 g/L) Women: 94–172 mg/dL (0.94–1.72 g/L) Apo B: Men: 55–100 mg/dL (0.55–1.00 g/L) Women: 45–110 mg/dL (0.45–1.10 g/L) Apo A-I/Apo B ratio: Men: 0.80–1.33 Women: 0.94–2.63

Procedure

1. Obtain a 5-mL venous blood sample. Serum is needed.
2. Do not freeze the specimen. Place specimen in a biohazard bag.
3. Fasting for 12 hours is needed.

Clinical Implications

1. *Increased* Apo A-I is associated with familial (inherited) hyper-a-lipoproteinemia.
2. *Decreased* Apo A-I is associated with the following conditions:
   a. Tangier disease (extremely low) hypo-a-lipoproteinemia
b. ß-Lipoproteinemia
c. Apo C-II deficiency
d. Apo A-I Melano disease
e. Apo A-I–C-III deficiency
f. Hypertriglyceridemia (familial)
g. Poorly controlled diabetes
h. Premature CHD
i. Hepatocellular disease
j. Nephrotic syndrome and renal failure

3. Increased Apo B is associated with the following conditions:
   a. Hyperlipoproteinemia types IIa, IIb, and V
   b. Premature CHD Fredrickson type IIa
   c. Diabetes mellitus
   d. Hypothyroidism
   e. Nephrotic syndrome, renal failure
   f. Hepatic disease and obstruction
   g. Dysglobulinemia
   h. Porphyria
   i. Cushing's syndrome
   j. Werner's syndrome

4. Decreased Apo B occurs with the following conditions:
   a. ß-Lipoproteinemia
   b. Hypo-a-lipoproteinuria (Tangier disease)
   c. Hypo-ß-lipoproteinemia
d. Type I hyperlipidemia
e. Apo C-II deficiency
f. Hypothyroidism
g. Malnutrition/malabsorption
h. Reye's syndrome

Interfering Factors

1. Decreased Apo A-I is associated with a diet high in polyunsaturated fats, smoking, and some drugs (see Appendix J).
2. Decreased Apo B is associated with a diet high in polyunsaturated fats and low-cholesterol diets, and many drugs.
3. Increased apolipoprotein levels can be caused by various drugs.
4. Apolipoproteins are acute-phase reactants and should not be measured in ill patients (e.g., acute stress, burns, major illness, inflammatory diseases)

Clinical Alert

An adverse Apo A-I/Apo B ratio in early life is a potential marker for CHD risk. Apo A-I values less than 90 mg/dL or 0.90 g/L indicate increased CAD risk. Apo B values above 110 mg/dL or 1.10 g/L indicate increased CAD risk.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. A 12-hour fast is required, but water may be taken. Smoking is prohibited.
2. Alcohol is prohibited.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume normal activities.
2. Interpret test results and counsel appropriately regarding CAD risk and potential lifestyle changes.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Triglycerides

Triglycerides account for >90% of dietary intake and comprise 95% of fat stored tissues. Because they are insoluble in water, they are the main plasma glycerol ester. Normally stored in adipose tissue as glycerol, fatty acids, and monoglycerides, the liver reconverts these to triglycerides. Of the total, 80% of triglycerides are in VLDL, and 15% are in LDL.

This test evaluates suspected atherosclerosis and measures the body's ability to metabolize fat. Elevated triglycerides, together with elevated cholesterol, are atherosclerotic disease risk factors. Because cholesterol and triglycerides can vary independent of each other, measurement of both values is more meaningful. Triglyceride level is needed to calculate the LDL-C and is also used to evaluate turbid samples of blood and plasma.

Reference Values

Normal Desired: <150 mg/dL or <1.70 mmol/L Borderline high: 150–199 mg/dL or 1.70–2.25 mmol/L High: 200–499 mg/dL or 2.26–5.64 mmol/L Very high: >500 mg/dL or >5.65 mmol/L See Table 6.16 for values.

Table 6.16 Values for Triglycerides
### Procedure

1. Obtain a 5-mL venous blood sample. Serum is used, but now many labs use EDTA anticoagulant plasma levels, which are slightly lower. Fasting for 12 to 14 hours is required.
2. Observe standard precautions. Do not use glycerinated tubes. Place specimen in a biohazard bag.

### Clinical Implications

1. **Increased triglycerides** occur with the following conditions:
   - a. Hyperlipoproteinemia type I, Iib, III, IV, and B
   - b. Liver disease, alcoholism (can be extremely high with alcoholism)
   - c. Nephrotic syndrome, renal disease
   - d. Hypothyroidism
   - e. Poorly controlled diabetes mellitus
   - f. Pancreatitis
   - g. Glycogen storage disease (von Gierke's disease)
   - h. Myocardial infarction (elevated levels may persist for several months)
     - i. Gout
     - j. Werner's syndrome
   - k. Down syndrome
   - l. Anorexia nervosa

2. **Decreased triglyceride** levels occur with the following conditions:
   - a. Congenital a-ß-lipoproteinemia
   - b. Malnutrition, malabsorption syndromes
   - c. Hyperthyroidism, hyperparathyroidism
   - d. Brain infarction
   - e. Chronic obstructive lung disease

### NOTE

Certain levels of triglycerides are associated with certain disorders:

1. Desirable: <150 mg/dL (<1.70 mmol/L)—not associated with a disease state
2. Borderline: 150–500 mg/dL (1.70–5.65 mmol/L)—associated with peripheral vascular disease and may be a marker for genetic forms of hyperlipoproteinemias that need specific therapy
3. Hypertriglyceridemia: >500 mg/dL (>5.6 mmol/L)—associated with risk for pancreatitis
4. >1000 mg/dL (>11.3 mmol/L)—associated with type I or V hyperlipidemia and substantial risk for pancreatitis
5. >5000 mg/dL (>56.5 mmol/L)—associated with eruptive xanthoma, corneal arcus, lipemia retinalis, and enlarged liver and spleen

### Interfering Factors

1. A transient increase occurs following a heavy meal or alcohol ingestion.
2. Transient decrease occurs after strenuous exercise, permanent decrease with weight loss.
3. Increased values are associated with pregnancy and oral contraceptive use.
4. Values may be increased in acute illness, colds, or flu.
5. Many drugs cause increases and decreases (see Appendix J).
6. Values are increased with obesity, physical inactivity, and smoking.

### Interventions

#### Pretest Patient Care

1. Explain test purpose and procedure. Fasting for at least 12 hours overnight is required, but water may be ingested.
2. Ask the patient to follow a normal diet for 1 week pretest. No alcohol is permitted for at least 24 to 48 hours before testing.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

#### Posttest Patient Aftercare

1. Interpret test results (Chart 6.4) and counsel appropriately. Weight reduction, a low-fat diet, and an exercise program can reduce high triglyceride levels.
2. Advise that triglycerides are not a strong predictor of CHD and, as such, are not an independent risk factor if <250 mg/dL (<2.8 mmol/L). However, increased levels may increase cardiovascular disease risk.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Chart 6.4 Example of Lipid Test Outcomes, Interpretation, and Intervention

A 63-year-old woman, on annual exam, had the following findings: moderately overweight, height 5'6" (167 cm), weight 170 pounds (77 kg), BP 126/72, pulse 72 regular, moderately active lifestyle, and family (father and mother) history of coronary artery and vascular diseases. Fasting lipid panel was ordered and done in the clinic July 10, 2002. Results were:

Results of First Testing *
Cholesterol (C) 320 mg/dL (8.3 mmol/L)
Triglycerides 414 mg/dL (4.68 mmol/L)
HDL (good) cholesterol (C) 38 mg/dL (0.99 mmol/L)
LDL (bad) cholesterol (C) Unable to calculate
Cholesterol/HDL ratio 8.4 H
*For reference ranges, see table below.

Because the triglycerides were high, glucose Hb A1c was ordered, and the results were within normal limits. A screening TSH was then done, with results of 6.4 MIU/L (normal, 0.40–5.50), and treatment with oral levothyroxine (Levoxyl), 50 mg (synthetic thyroid) every day for 2 months.

On September 10, 2002, a fasting lipid panel and TSH tests were repeated. Results are reported in the following table:

<table>
<thead>
<tr>
<th>Chemistry</th>
<th>ABN/CRIT Units</th>
<th>Ref. Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cholesterol</td>
<td>333 H mg/dL</td>
<td>&lt; 200</td>
</tr>
<tr>
<td>Triglycerides</td>
<td>496 H mg/dL</td>
<td>35–160</td>
</tr>
<tr>
<td>HDL (good) C</td>
<td>44 mg/dL</td>
<td>&gt;40</td>
</tr>
<tr>
<td>LDL (bad) C</td>
<td></td>
<td>70–130</td>
</tr>
<tr>
<td>Chol/HDL ratio</td>
<td>7.6 H</td>
<td>3.0–6.0</td>
</tr>
</tbody>
</table>
*Unable to calculate owing to triglycerides greater than 400 mg/dL.

Elevated triglycerides were treated by administering gemfibrozil, 600 mg bid for 3 months and a low-cholesterol diet. Levothyroxine was continued to maintain TSH levels WNL.

Repeat lipid panels were done at a community testing at a local drugstore on December 15, 2003, with the following results given to the physician at the next follow-up office visit. Gemfibrozil was discontinued and atorvastatin (Lipitor), 200 mg 1 tablet daily at 8 p.m. was begun for 2 months and then follow-up lipid panel was ordered.

Results of Second Testing
Total blood cholesterol 289 mg/dL (7.5 mmol/L)
HDL (good) cholesterol (C) 30 mg/dL (0.78 mmol/L)
Total C to HDL ratio 9.6
LDL (bad) cholesterol (C) 92 mg/dL (2.4 mmol/L)
Triglycerides 100 mg/dL (1.13 mmol/L)

<table>
<thead>
<tr>
<th>Lipids</th>
<th>Excellent Protection</th>
<th>Moderate Risk</th>
<th>High Risk</th>
<th>Very High Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Cholest. (mg/dL)</td>
<td>Age (yr) M F M F M F M F</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>60+</td>
<td>=213 =227 214–240 228–252 252–262 257–278 &gt;262 &gt;275</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LDL (bad) Cholest. (mg/dL)</td>
<td>Age (yr) M F M F M F M F</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>60+</td>
<td>=143 =149 144–165 150–175 186–190 176–198 &gt;190 &gt;198</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HDL (good) Cholest. (mg/dL)</td>
<td>Age (yr) M F M F M F M F</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>20–39</td>
<td>=51 =63 51–37 63–45 &lt;37 &lt;45 — —</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>40–59</td>
<td>=52 =69 52–37 69–49 &lt;37 &lt;49 — —</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;60+</td>
<td>=60 =74 60–40 74–50 &lt;40 &lt;50 — —</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triglyc. (mg/dL)</td>
<td>Age (yr) M F M F M F M F</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>40–59</td>
<td>=121 =98 122–170 99–140 171–231 144–190 &gt;231 &gt;190</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>
Triglyc. (mg/dL)       
40–59 =121 =98  122–170 99–140  171–231 144–190 >231 >190
60+ =110 =110  111–154 111–146  147–206 147–200 >206 >206

Total Cholest./HDL ratio      
20–39 =3.6 =2.8  3.7–5.1 2.9–3.8  5.2–6.1 3.7–4.2 >61 >42
40–59 =4.2 =3.0  4.3–6.0 3.1–4.9  6.1–7.4 4.1–4.9 >74 >2.9
60+ =4.0 =3.2  4.1–6.0 3.3–4.8  6.1–6.9 4.9–5.5 >6.9 >5.5

These test results were given to physician at the next follow-up office visit.

On February 20, 2003, the lipid panel results were:

Total cholesterol 190 mg/dL (4.9 mmol/L)  
Triglycerides 207 mg/dL (2.34 mmol/L)  
HDL 45 mg/dL (1.17 mmol/L)  
LDL 104 mg/dL (2.7 mmol/L)

Because the patient had arthritis, she was treated with rofecoxib (Vioxx), 25 mg every day, and ranitidine, 150 mg twice daily. Liver studies were indicated. Results of the ALT were 28 (normal, 10–60 U/L or 0.17–1.02 µkat/L). As a result of these testings, Lipitor and arthritis medications were continued until the next physician visit in 6 months.

Clinical Alert
Test normal or reference values may vary somewhat from different laboratories owing to testing methods used. It is important to compare test results/outcomes with the normal reference values for the specific laboratory.

Clinical Alert
1. Panic values of >500 mg/dL (>5.6 mmol/L) indicate hypertriglyceridemia in the presence of diagnosed pancreatitis.
2. Values of >1000 mg/dL (>11.3 mmol/L) present a substantial risk for pancreatitis.
3. Chylomicronemia, although associated with pancreatitis, is not accompanied by increased atherogenesis. Chylomicrons are not seen in normal fasting serum but instead are found as exogenous triglycerides in healthy persons after a fatty meal has been eaten. After refrigeration, chylomicrons float to the surface of a blood sample.

Lipoprotein Electrophoresis
Lipoproteins are composed of hydrophobic lipids bound to protein, which produces a liquid-soluble complex. Chylomicrons primarily transport dietary triglycerides from the intestines. They are proteins derived from dietary sources, and if significantly increased, they can extend into the pre-ß area. In hyperchylomicronemia, chylomicrons represent dietary fat in transport. The standing plasma contains a cream layer over a clear layer in type I hyperlipidemia (where chylomicrons are elevated), but not in type IV (where both chylomicrons and triglycerides are elevated). VLDLs transport cholesterol and triglycerides that have been synthesized in the liver. LDLs are the major cholesterol-transporting lipoproteins. Atherosclerotic plaque cholesterol is derived from LDLs, and LDL elevations are associated with an increased CAD risk. Conversely, HDLs provide protection against atherosclerosis by reversing cholesterol transport mechanisms. Levels of plasma HDL cholesterol are inversely proportional to the risk for heart disease.

Lipoprotein electrophoresis evaluates hyperlipidemia and determines abnormal serum lipoprotein distribution and concentration. Quantitation is not available with this procedure. Visual estimates of stain density in comparison to normal patterns are usually done. Serum cholesterol and triglyceride levels should also be done at the same time.

Reference Values
Normal For 12- to 14-hour fasting specimen: Chylomicrons: 0%–2% about 90% triglycerides ß or LDL: 33%–52% (mass fraction of total lipoprotein)—cholesterol, triglyceride phospholipid Pre-ß or VLDL: 7%–28% (mass fraction of total lipoprotein)—triglyceride, phospholipid cholesterol a or HDL: 10%–30% (mass fraction of total lipoprotein)—protein, phospholipid, cholesterol Plasma appearance: clear

Procedure
1. Obtain a 5-mL sample of serum or plasma. Fasting 12–14 hours is required. Do not freeze.
2. Observe standard precautions. Place specimen in a biohazard bag. To aid in the classification, the blood sample is refrigerated overnight, and the serum or plasma is observed for any creamy layers, turbidity, or color change.

Clinical Implications
1. Patients may be phenotyped (ie, physical appearance or classification makeup) using Frederickson's classification system. Triglyceride, cholesterol, and lipoprotein levels are considered in this system.
2. Lipoproteins are decreased in the following conditions:
   a. ß-Lipoproteinemia
   b. Tangier disease
3. Lipoproteins are increased in the following conditions:
   a. Hyper-ß-lipoproteinemia
   b. Hypercholesterolemia
   c. Hyper-a-lipoproteinemia
   d. Hyper pre-ß-lipoproteinemia

Interfering Factors

1. Lipid phenotypes are affected by stress or dietary changes.
2. Phenotyping is invalid in the presence of secondary disorders, such as diabetes mellitus, renal failure, or nephritis.
3. Certain drugs may alter electrophoretic mobilizing of lipoproteins.
4. Heparinized blood is not acceptable; test results are not reliable during heparin therapy.

NOTE
A clear distinction must be made between primary (inherited) and secondary (liver disease, alcoholism, metabolic diseases) causes.

Interventions

Pretest Patient Care

1. Explain test purpose and blood-drawing procedure. A 12-hour fast is required before blood is drawn.
2. Ask the patient to follow a normal diet for 2 weeks before test.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and counsel appropriately regarding dietary and drug therapy. The National Cholesterol Education Program and other organizations have many resources available. (National Cholesterol Education Program, National Institutes of Health, 9000 Rockville Pike, Bethesda, MD 20814).
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

NOTE
This test has been largely replaced with the lipid profile panel.

Free Fatty Acids; Fatty Acid Profile

Free fatty acids are formed by lipoprotein and triglyceride breakdown. The amount of free fatty acids and triglycerides present in blood comes from dietary sources or fat deposits or is synthesized by the body. Carbohydrates can be converted to fatty acids and then stored in fat cells as triglycerides. Fatty acid and carbohydrate metabolism is altered in the fat breakdown process (eg, when fasting). Unusually high levels are associated with untreated diabetes.

Specific fatty acid measurement can be useful for monitoring nutritional status in the presence of malabsorption, starvation, and long-term parenteral nutrition. It is also valuable for the differential diagnosis of polyneuropathy when Refsum's disease is suspected. In this disease, the enzyme that degrades phytic acid is lacking. Free fatty acids are also useful in detecting pheochromocytoma and glucagon thyrotropin and adrenocorticotropic-secreting tumors.

Reference Values

**Normal** Adults: 8–25 mg/dL or 0.28–0.89 mmol/L. Children: <31 mg/dL or obese adults: <31 mg/dL or <1.0 mmol/L. **Fatty acid profile** Linoleate: >25% of total fatty acids Arachnidate: 0%–6% Oleic: 26%–35% Linoleic: 8%–16% Steric: 10%–14% Phytic acid Normal: 0.3% Borderline: 0.3%–0.5%

Procedure

1. Obtain a 5-mL blood sample and place on ice. Serum or EDTA plasma may be used.
2. Fasting is required.
3. The blood serum should be separated from blood cells within 45 minutes of collection and should be placed on ice. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications

1. **Increased free fatty acid values** are associated with the following conditions:
   a. Poorly controlled diabetes mellitus
   b. Pheochromocytoma
   c. Hyperthyroidism
   d. Huntington's chorea
   e. von Gierke's disease
   f. Alcoholism
   g. Acute myocardial infarction
   h. Reye's syndrome
2. **Increased phytic acid** occurs in the following conditions:
   a. Refsum's disease (>50%; repeat the test to confirm)
b. ß-Lipoproteinemia

3. **Decreased fatty acids** are found in:
   a. Cystic fibrosis
   b. Malabsorption (acrodermatitis enteropathica)
   c. Zinc deficiency (linoleate and arachnidate low)

**Interfering Factors**

1. Values are elevated by strenuous exercise, anxiety, hypothermia, certain drugs (see Appendix J), and long-term fasting.
2. Values are decreased by long-term IV or parenteral nutrition therapy and certain drugs (see Appendix J).
3. Prolonged fasting or starvation (rise as much as 3 times normal) affects levels.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and blood-drawing procedure. Fasting is required, but water may be taken.
2. Do not test patients receiving heparin therapy. For free fatty acids, no alcohol may be taken within 24 hours.
3. Discontinue strenuous exercise before the test. Encourage relaxation.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Resume normal activities.
2. Interpret test results and monitor appropriately.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

See Figure 6.6 for a complete laboratory test.

---

**FIGURE 6.6** Example of Laboratory Test Results (54 year old female, annual physical examination). Includes lipid panel, liver panel, and basic metabolic panel.

**THYROID FUNCTION TESTS**

Laboratory determinations of thyroid function are useful in distinguishing patients with euthyroidism (normal thyroid gland function) from those with hyperthyroidism (increased function) or hypothyroidism (decreased function).

**Patient Care for Thyroid Testing**

**Pretest Patient Care**

1. Explain test purpose and blood specimen collection procedure. To understand the thyroid function tests, it is necessary to understand the following basic concepts. The thyroid gland takes iodine from the circulating blood, combines it with the amino acid tyrosine, and converts it to the thyroid hormones thyroxine (T\(_4\)) and triiodothyronine (T\(_3\)). Iodine composes about two thirds of the weight of the thyroid hormones. The thyroid gland stores T\(_3\) and T\(_4\) until they are released into the bloodstream under the influence of TSH from the pituitary gland. Only a small amount of the hormones is not bound to protein. However, it is the free portion of the thyroid hormones that is the true determinant of the thyroid status of the patient.
2. Assess for signs and symptoms of thyroid disease and note thyroid and iodine medications. Fasting is required for some tests.
3. Be aware that a typical thyroid panel includes the following tests:
   a. T\(_3\) uptake (T\(_3\) U)
   b. Free T\(_4\)
   c. Total T\(_4\)
   d. T\(_3\) total
   e. Free thyroxine index (FTI, T\(_7\))
   f. TSH
4. Remember that the most useful laboratory tests to confirm or exclude hyperthyroidism are total T\(_4\), free thyroxine
index (FTI), total T₄, and the ultrasensitive TSH. The most useful tests to detect hypothyroidism are total T₄, FTI, and TSH (thyrotropin). A thyrotropin-releasing hormone (TRH) stimulation test can be valuable in establishing the thyroid status in some patients with equivocal signs of thyroid dysfunction and borderline laboratory values. It should be kept in mind that values obtained for the assessment of thyroid function can be influenced by factors other than disease, such as age, current illness, binding capacity of serum proteins, and some drugs.

5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results, counsel and monitor appropriately for abnormal thyroid function and disease. Follow-up testing may be required.
2. Remember that thyroid antibody testing can also be done for diagnosis of autoimmune thyroid testing.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Calcitonin

Calcitonin, a hormone secreted by the C cells (parafollicular) of the thyroid gland, inhibits bone resorption by regulating the number and activity of osteoblasts. Calcitonin is secreted in direct response to high blood calcium levels and helps to prevent abrupt changes in calcium levels and the excessive loss of calcium.

Measurement of calcitonin is used to diagnose familial medullary thyroid carcinoma (MTC) and postoperatively to detect recurrence or metastasis of thyroid carcinoma. This test is done to measure increases in immunoreactive calcitonin after stimulation with calcium and/or pentagastrin. Early detection of elevated calcitonin leads to diagnosis of tumor or abnormally secreting C cells before cancer spreads. (Doubling of serum levels correlates with recurrence.) Calcitonin levels are also used in the investigation of families (of a patient with MTC) to detect early subclinical cases of MTC that may exist as C-cell hyperplasia or microscopic MTC.

Reference Values

**Normal**
- Men: <19 pg/mL or <19 ng/L
- Women: <14 pg/mL or <14 ng/L

**Calcium infusion (2.4 mg/kg):**
- Men: <190 pg/mL or <190 ng/L
- Women: <130 pg/mL or <130 ng/L

**Pentagastrin injection (0.5 µg/kg):**
- Men: <110 pg/mL or <110 ng/L
- Women: <35 pg/mL or <35 ng/L

Procedure

1. Obtain a 5-mL venous blood specimen in green-topped tube. Fasting is necessary.
2. Heparinize and chill the blood immediately. If testing is not performed immediately, blood should be frozen. Place specimen in a biohazard bag.

Clinical Implications

1. **Increased levels of calcitonin** are associated with the following conditions:
   - Medullary thyroid cancer (MTC)
   - C-cell hyperplasia
   - Chronic renal failure
   - Pernicious anemia
   - Zollinger-Ellison syndrome
   - Cancer of lung (oat cell lung marker), breast, or pancreas (ectopic calcitonin)
   - Carcinoid syndrome
   - Alcoholic cirrhosis
   - Patients with pancreatitis and thyroiditis
   - Hypercalcemia of any etiology

2. In a small proportion of patients who do have medullary cancer, the fasting level of calcitonin is normal. In these instances, a provocative test using calcium or pentagastrin should be done.
   - Very high levels (ie, 5- to 30-fold increase over basal levels) are evidence of MTC but are not diagnostic.
   - These stimulation tests are not needed if the basal calcitonin test is diagnostically high.
   - In patients with elevated calcitonin levels who do not have MTC, the response is not as vigorous.

Interfering Factors

1. Levels are normally increased in pregnancy at term and in newborns.
2. Gross lipemia and hemolysis interfere with test.

**Clinical Alert**

1. Screening families of patients with proven medullary cancer of the thyroid with the calcitonin test is recommended because the tumor has both sporadic and familial incidence.
2. If the calcitonin test is normal in family members, it is advisable to repeat the calcium provocative test periodically (over a period of months or years).
3. Some patients do not respond to the stimulation test who have medullary thyroid carcinoma.

Interventions

**Pretest Patient Care**

1. Explain test purpose and procedure.
2. Remind patient that fasting from food overnight is required. Water is permitted.
3. Be aware that if the provocative tests using calcium and pentagastrin are to be done, the patient is to be fasting,
a. Inject pentagastrin 0.5 µg/kg IV push. Draw blood samples before the injection to determine baseline value of calcitonin. Draw a blood sample 1.5, 2, and 5 minutes after the injection.
b. Inject calcium, 2.0 mg/kg, after baseline sample is drawn. Draw a blood sample 5 and 10 minutes after injection.

4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

NOTE
A combined calcium and pentagastrin test may be more effective and reliable than either test by itself.

Posttest Patient Aftercare
1. Interpret test outcome and monitor side effects of injection.
2. Be aware that the patient may experience transient nausea or fatigue after injection and may experience chest pain for a short time.
3. Resume normal activities when symptoms abate.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Free Thyroxine (FT₄)

Free thyroxine (FT₄) comprises a small fraction of total thyroxine. The FT₄ is unbound to protein and available to the tissues, and it is the metabolically active form of this hormone. This fraction constitutes about 5% of the circulatory T₄.

FT₄ has diagnostic value in situations in which total hormone levels do not correlate with the thyrometabolic state, and there is suspected abnormality in thyroxine-binding globulin (TBG) levels. It provides a more accurate picture of the thyroid status in persons with abnormal TBG levels in pregnancy and in those who are receiving estrogens, hydrogen, phenytoin, or salicylates.

Reference Values
Normal 0.7–2.0 ng/dL or 10–26 pmol/L For patients taking levothyroxine (Synthroid), up to 5.0 ng/dL or 64 pmol/L

Procedures
1. Obtain a 5-mL venous blood sample. Accurate results can be obtained with as little as 0.5 mL of blood in pediatric cases. Serum is needed for this test.
2. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications
1. Increased FT₄ levels are associated with the following conditions:
   a. Graves' disease (hyperthyroidism)
   b. Hypothyroidism treated with thyroxine
   c. Euthyroid sick syndrome
2. Decreased FT₄ levels are associated with the following conditions:
   a. Primary hypothyroidism
   b. Secondary hypothyroidism (pituitary)
   c. Tertiary hypothyroidism (hypothalamic)
   d. Hypothyroidism treated with triiodothyronine

Interfering Factors
1. Values are increased in infants at birth and rise even higher after 2 to 3 days of life.
2. Many drugs affect test outcomes (see Appendix J).
3. Heparin causes falsely elevated FT₄ values.
4. Levels can fluctuate in patients with severe or chronic illness.
5. Levels fluctuate in pregnancy (low in late pregnancy)

Interventions
Pretest Patient Care
1. See patient care for thyroid testing. The same protocols prevail in FT₄ testing.
2. Follow Chapter 1 guidelines for safe, effective informed pretest care.

Posttest Patient Aftercare
1. See patient care for thyroid testing. The same protocols prevail in FT₄ testing.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Free Triiodothyronine (FT₃)

This is one of the determinations used to evaluate thyroid function and measure that fraction of the circulatory T₃ that exists in the free state in the blood, unbound to protein. FT₃ is done to rule out T₃ toxicosis, to evaluate thyroid
replacement therapy, and to clarify protein-binding abnormalities.

Reference Values

**Normal** Adults: 260–480 pg/dL or 4.0–7.4 pmol/L

**Procedure**

1. Obtain a 5-mL venous blood sample.
2. Observe standard precautions. Place specimen in a biohazard bag.

**Clinical Implications**

1. *Increased FT₃ values* are associated with the following conditions:
   a. Hyperthyroidism
   b. T₃ toxicosis
   c. Peripheral resistance syndrome
2. *Decreased FT₃ values* are associated with the following conditions:
   a. Hypothyroidism (primary and secondary)
   b. Third trimester of pregnancy

**NOTE**

In nonthyroidal illness, a low FT₃ level is a nonspecific finding.

**Interfering Factors**

1. Recently administered radioisotopes and some drugs (see Appendix J)
2. High altitude: FT₃ levels are higher

**Interventions**

**Pretest Patient Care**

1. See patient care for thyroid testing. The same protocols prevail for FT₃.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. See patient care for thyroid testing. The same protocols prevail in FT₃ testing.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Free Thyroxine Index (FTI, T₄)**

The free thyroxine index (FTI) is a mathematical calculation used to correct the estimated total T₄ for the amount of TBG present. To perform this calculation, two results are needed: the T₄ value and the T₃ uptake ratio. The product of these two values is the FTI. The FTI is useful in the diagnosis of hyperthyroidism and hypothyroidism, especially in patients with known or suspected abnormalities in thyroxine-binding protein levels. In such cases, blood levels and clinical signs may seem contradictory unless both T₄ and TBG are considered as interrelated parameters of thyroid status. Measurement of FT₄ also gives a more accurate picture of the thyroid status when the TBG is abnormal in pregnant women or persons being treated with estrogen, androgens, phenytoin, or salicylates.

**Reference Values**

**Normal** Adults: 1.5–4.5 index (these are arbitrary units) Check with your laboratory for their normal values.

**Procedure**

1. Make a calculation based on results of T₃ uptake and T₄ total, as follows:

   \[ \text{FTI} = \frac{T₄ \text{ total} \times T₃ \text{ U(%)}}{100} \]

2. Remember that the FTI permits meaningful interpretation by balancing out most nonthyroidal factors. In recent years, this parameter has lost popularity and is of dubious value.

**Clinical Implications** Application of the equation of the FTI includes the information presented in Table 6.17. This is a mathematical calculation that does not involve the patient.

**Table 6.17 Application of Equation to Determine Thyroxine Uptake**

<table>
<thead>
<tr>
<th>Status</th>
<th>TBG</th>
<th>T₃ Uptake</th>
<th>FTI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Euthyroid Normal</td>
<td>35%</td>
<td>9.0</td>
<td>3.1</td>
</tr>
<tr>
<td>Euthyroid Low</td>
<td>52%</td>
<td>4.0</td>
<td>2.1</td>
</tr>
<tr>
<td>Euthyroid High</td>
<td>13%</td>
<td>16.0</td>
<td>2.8</td>
</tr>
<tr>
<td>Hypothyroid High</td>
<td>24%</td>
<td>4.0</td>
<td>0.9</td>
</tr>
<tr>
<td>Hyperthyroid Low</td>
<td>46%</td>
<td>13.0</td>
<td>6.0</td>
</tr>
</tbody>
</table>
Interfering Factors

1. Levels fluctuate in pregnancy.
2. See Appendix J for drugs that affect test outcomes.

Interventions

**Pretest Patient Care**

1. Inform the patient about the test purpose and method of calculation.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Be prepared to counsel parent if treatment required.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Neonatal Thyroid-Stimulating Hormone (TSH)

Signs of congenital hypothyroidism are minimal at birth. Congenital hypothyroidism has an incidence of 1:3600 to 1:5000 in the United States.

This measurement is used as a confirmatory test or in conjunction with neonatal T4 for infants with positive T4 screens or low blood serum T4 levels, and for screening in all U.S. states. See newborn screening in Chapter 11 for more information.

**Reference Values**

**Normal** Newborn screen: <20 µU/mL or <20 mU/L by third day of life TSH surges at birth, peaking at 30 minutes of life at a level of 25–160 µU/mL or 25–160 mU/L. It declines and reaches adult levels by the first week to 10 days of life.

**Procedure**

1. Cleanse the infant's heel with an antiseptic and puncture with a sterile disposable lancet. Collect this whole blood specimen 3 to 7 days after birth.
2. Be aware that if bleeding is slow, it helps to hold the leg dependent for a short time before blotting the blood on the filter paper. Do not use pipettes or capillary tubes to collect blood.
3. Completely fill in the circles on the filter paper. This can best be done by placing one side of the filter paper against the infant's heel and watching for the blood to appear on the front side of the paper and to fill the circle completely. The filter paper is a special filter paper card obtained from the laboratory.
4. Air dry the filter paper for 1 hour, fill in all information, and send to the laboratory immediately. Do not expose samples to extreme heat or light.

**Clinical Implications** An elevated neonatal TSH test is associated with neonatal hypothyroidism, a confirmatory test.

**Interventions**

**Pretest Patient Care**

1. Inform the parents about the test purpose and method of specimen collection.
2. See patient care for thyroid testing on page 437.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Be prepared to counsel parent or parents regarding steps to take if the TRH test is abnormal and type of treatment required. See newborn screening in Chapter 11.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Neonatal Thyroxine (T4); Neonatal Screen for Hypothyroidism

Normal brain growth and development cannot take place without adequate thyroid hormone. Congenital hypothyroidism (cretinism) is characterized by low levels of T4 and elevated levels of TSH. Screening for congenital hypothyroidism is now done in all 50 states. If hypothyroidism is undetected, growth and mental retardation occur, and in some cases, death occurs.

This is a screening test of T4 activity to detect neonatal hypothyroidism. Specimens should be obtained after the first 24 hours of protein feeding or within the first week of life. Thyroxine is obtained from whole blood blotted on filter paper using a radioimmunoassay technique.

**Reference Values**

**Normal** Peaks in 24 hours then decreases. Neonates (1–3 days): 12–22 µg/dL or 152–292 nmol/L Neonates (1–2 weeks): 10–17 µg/dL or 126–214 nmol/L

**Procedure**

1. Cleanse the infant's heel with an antiseptic and puncture the skin with a sterile disposable lancet. To help blood flow, warm the foot or massage the leg.
2. Be aware that if bleeding is slow, it helps to hold the leg dependent for a short time before blotting the blood on the
filter paper. Wipe away the first drop of blood.

3. Completely fill in the circles on the filter paper. This can best be done by placing one side of the filter paper against the infant’s heel and watching for the blood to appear on the front side of the paper and to fill the circle completely. Do not damage filter paper. Apply a sterile dressing to the wound.

4. Air dry for 1 hour, fill in all requested information, and send to the laboratory immediately. Protect specimen from extreme heat and light.

Clinical Implications

1. Low values are associated with hypothyroidism.
2. A number of nonthyroid conditions can result in depressed T₄ levels (eg, low birth weight, prematurity, twinning, fetal distress, deficient TBG levels).

Interventions

Pretest Patient Care

1. Refer to neonatal TSH testing for care. The same protocols prevail for neonatal T₄.
2. Be aware that T₄ is usually collected at the same times as the phenylketonuria (PKU) specimen.
3. Remember that the optimal collection time is 3–7 days after birth; the baby must be on protein feeding for at least 24 hours. For low-weight or premature babies, the recommended time is 4–10 days old.

Posttest Patient Aftercare

1. Refer to neonatal TSH testing for care. The same protocols prevail for neonatal T₄. Also, see newborn screening in Chapter 11.
2. Remember that if baby is released early, the baby must be brought back for testing.

Thyroglobulin (Tg)

Thyroglobulin is composed of glycoprotein and the iodinated secretions of epithelial cells of the thyroid. These iodinated secretions contain both the precursors of T₃ and T₄ and the hormones themselves.

This test is helpful in the differential diagnosis of hyperthyroidism and in monitoring the course of differentiated or metastatic thyroid cancer. It is not useful in the diagnosis of thyroid cancer. Levels decrease following successful initial treatment, and in recurrence of metastases, the level will again rise. Lack of sensitivity and specificity limits the value of this test.

Reference Values

<table>
<thead>
<tr>
<th>Normal Adults</th>
<th>3–42 ng/mL or 3–42 µg/L Newborns (48 hours): 36–48 ng/mL or 36–48 µg/L</th>
</tr>
</thead>
</table>

NOTE

87% of normal adults have serum values of Tg <10 ng/mL or <10 µg/L. Athyrotic patients have values <5 ng/mL or <5 µg/L.

Procedure

1. Obtain a 5-mL venous blood sample. Serum is needed.
2. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications

1. Increased thyroglobulin levels are associated with the following conditions:
   a. Untreated and metastatic differentiated thyroid cancers (not MTC)
b. Hyperthyroidism (not good correlation with elevated \( T_4 \))
c. Subacute thyroiditis, thyrotoxicosis
d. Benign adenoma (some cases)
e. Occurrence of metastases after initial treatment (thyroid carcinoma)

2. Decreased thyroglobulin levels are associated with the following conditions:
   a. Thyrotoxicosis factitia
   b. Infants with goitrous hypothyroidism

Interfering Factors

1. Newborns have high Tg levels that drop to adult levels by 2 years of age
2. Autoantibodies to Tg cause decreased values. Thyroglobulin antibody test may have to be done to confirm decreased levels.

Interventions

Pretest Patient Care

1. See patient care for thyroid testing on page 437.
2. Ensure that patient is off thyroid medication for 6 weeks before specimen collection. The TSH should be elevated before testing for thyroglobulin.
3. Determination of Tg levels may be substituted for \(^{131}\)I scans in patients at low risk for thyroid cancer.

Posttest Patient Aftercare

1. Resume thyroid medication and normal activities.
2. Monitor as appropriate for metastatic thyroid cancer.
3. Refer to patient aftercare instructions for thyroid testing on page 439. The same protocols prevail for Tg testing.

Thyroid-Stimulating Hormone (Thyrotropin; TSH)

The thyroid is unique among the endocrine glands because it has a large store of hormone and a slow rate of normal turnover. Stimulation of the thyroid gland by the TSH, which is produced by the anterior pituitary gland, causes the release and distribution of stored thyroid hormones. TSH stimulates secretion of \( T_4 \) and \( T_3 \). TSH secretion is physiologically regulated by \( T_3 \) and \( T_4 \) (feedback inhibition) and is stimulated by thyrotropin-releasing hormone (TRH) from the hypothalamus. TSH is the single most sensitive test for primary hypothyroidism. If there is clear evidence of hypothyroidism and the TSH is not elevated, then an implication of possible hypopituitarism exists.

This measurement is used in the diagnosis of primary hypothyroidism when there is thyroid gland failure owing to intrinsic disease, and it is used to differentiate primary from secondary hypothyroidism by determining the actual circulatory level of TSH. TSH levels are high in primary hypothyroidism. Low TSH levels occur in hyperthyroidism.

TSH measurements with sufficient sensitivity to distinguish low levels from normal levels have become the preferred test for hyperthyroidism. The high-sensitivity TSH test is useful for diagnosing sick euthyroid patients and in differentiating mild hyperthyroidism from Graves' disease. With the new, sensitive assays, a TRH stimulation test is no longer necessary.

Reference Values

**Normal**

- Adults: 0.4–4.2 mIU/L (SI units are the same)
- Neonates: 3–20 µIU/L by day 3 of life

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum is needed.
2. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications

1. **Increased TSH levels** are seen in the following conditions:
   a. Adults and neonates with primary hypothyroidism
   b. Thyrotropin-producing tumor (eg, ectopic TSH secretion from lung, breast tumors)
   c. Hashimoto's thyroiditis
   d. Thyrotoxicosis due to pituitary tumor
   e. TSH antibodies (rare)
   f. Hypothyroid patients receiving insufficient thyroid replacement hormone or thyroid hormone resistance

2. **Decreased TSH levels** are associated with the following conditions:
   a. Primary hyperthyroidism
   b. Secondary and tertiary hypothyroidism
   c. Treated Graves' disease
   d. Euthyroid sick disease
   e. Overreplacement of thyroid hormone in treatment of hypothyroidism
Clinical Alert

Critical Values

<0.1 mIU/L is an indication of primary hyperthyroidism or exogenous thyrotoxicosis.

Risk exists for atrial fibrillation at TSH levels <0.1 mIU/L (major risk factor for stroke).

Interfering Factors

1. Values are normally high in neonatal cord blood. There is hypersecretion of TSH in newborns up to 2 to 3 times normal. The TSH level approaches normal by the first week of life.
2. Values are suppressed during treatment with thyroxine and corticosteroids. See Appendix J for other drugs.
3. Values are abnormally increased with lithium, potassium iodide, amphetamine abuse, and iodine-containing drugs.
4. Radioisotopes administered within 1 week before test invalidate the result.
5. Values may be decreased in the first trimester of pregnancy.
6. Values are increased in elderly patients (>80 years old); upper limit for these patients is 10.0 mIU/L.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results and counsel as appropriate for hypothyroidism or hyperthyroidism.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Thyroxine-Binding Globulin (TBG)

Almost all of the thyroid hormones in the blood are protein bound: albumin, thyroid-binding prealbumin, and most important, thyroxine-binding globulin (TBG). Variations in TBG levels have a major effect on bound and free (metabolically active) forms of T_4 and T_3. Before considering this test, TSH, FTI, and total T_4 should be measured.

The TBG measurement is useful to distinguish between hyperthyroidism causing high T_4 levels and euthyroidism with increased binding by TBG, increased T_4, and normal levels of free hormones; to identify hereditary deficiency or increase of TBG; and to work up thyroid disease in hypothyroid populations, when the mean TBG concentration is significantly higher than the mean level in normal thyroid populations. In hyperthyroid populations, the mean TBG level concentration is lower than the mean level in normal thyroid populations.

Reference Values

<table>
<thead>
<tr>
<th>Group</th>
<th>Reference Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal Infants:</td>
<td>3–6 mg/dL or 30–60 mg/L</td>
</tr>
<tr>
<td>Men: 1.2–2.5 mg/dL or 12–25 mg/L Women: 1.4–3.0 mg/dL or 14–30 mg/L On oral contraceptives: 1.5–5.5 mg/dL or 15–55 mg/L Third trimester of pregnancy: 4.7–5.9 mg/dL or 47–59 mg/L</td>
<td></td>
</tr>
</tbody>
</table>

Procedure

1. Obtain a 5-mL venous blood specimen. Serum is needed.
2. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications

1. The TBG test is increased in the following conditions:
   a. Genetically determined high TBG
   b. Hypothyroidism (some cases)
   c. Infectious hepatitis
   d. Acute intermittent porphyria
   e. Estrogen-producing tumors (endogenous or exogenous)
   f. Late-stage HIV infections
2. The TBG test is decreased in the following conditions:
   a. Genetic deficiency of TBG
   b. Nephrotic syndrome
   c. Major illness, surgical stress
   d. Ovarian hypofunction
   e. Acromegaly
   f. Chronic liver disease
   g. Marked hypoproteinemia, malnutrition

Interfering Factors

1. Many drugs increase (eg, estrogens, oral contraceptives) or decrease (eg, nicotinic acid, phenytoin, and steroids) values (see Appendix J).
2. Neonates have higher values.
3. Recently administered radioisotopes affect results.
4. Pregnancy increases levels.
5. Prolonged heroin use or methadone increases levels.

Interventions

Pretest Patient Care

1. See patient care for thyroid testing (see page 437).
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. See patient care for thyroid testing (see page 439).
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Thyroxine (T₄), Total

Thyroxine is the thyroid hormone with four atoms of iodine; hence, it is called T₄. The combination of the serum T₄ and T₃ uptake as an assessment of TBG helps to determine whether an abnormal T₄ value is due to alterations in serum TBG or to changes in thyroid hormone levels. Deviations of both tests in the same direction usually indicate that an abnormal T₄ level is due to abnormalities in thyroid hormone. Deviations of the two tests in opposite directions provide evidence that an abnormal T₄ may relate to alterations in TBG.

Thyroxine, one of the thyroid function panel tests, is a direct measurement of the concentration of T₄ in the blood serum. Total T₄ level is a good index of thyroid function when the TBG is normal. The increase in TBG levels normally seen in pregnancy and with estrogen therapy increases total T₄ levels. The decrease of TBG levels in persons receiving anabolic steroids, in chronic liver disease, and in nephroses decreases the total T₄ value. This test is commonly done to rule out hyperthyroidism and hypothyroidism. The T₄ test also can be used as a guide in establishing maintenance doses of thyroid in the treatment of hypothyroidism. In addition, it can be used in hyperthyroidism to follow the results achieved with antithyroid drug administration.

Reference Values

Normal
- Adults: 5.4–11.5 µg/dL or 57–148 nmol/L
- Children: 6.4–13.3 µg/dL or 83–172 nmol/L
- Neonates: 11.8–22.6 µg/dL or 152–292 nmol/L
If testing is done by radioimmunoassay, it is reported as T₄ RIA.

Procedure

1. Obtain a 5-mL venous blood sample. Serum is used. If the patient is already receiving thyroid treatment, it must be discontinued 1 month before the test.
2. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications

1. Increased T₄ values are found in the following conditions:
   a. Hyperthyroidism (Graves’ disease, goiter)
   b. Clinical status that increases TBG
   c. Thyrotoxicosis factitia
   d. Acute thyroiditis
   e. Hepatitis, liver disease
   f. Lymphoma
2. Decreased T₄ values are found in the following conditions:
   a. Hypothyroidism
   b. Disorders of decreased TBG
   c. Hypoproteinemia
   d. Treatment with triiodothyronine
   e. Nephrotic syndrome

Interfering Factors

1. Total thyroxine levels increase during the second or third month of pregnancy as a result of increased estrogen production. Normal range: 5.5–16.0 µg/dL or 71–206 nmol/L.
2. Total thyroxine levels increase with the use of drugs such as estrogens, heroin, and methadone and excess iodine (see Appendix J).
3. Contrast agents used for x-rays and other diagnostic procedures affect results.
4. Values are decreased with salicylates and anticonvulsants, steroids.

Clinical Alert

Panic Range

>20 µg/dL or >258 nmol/L: Thyroid storm is possible.
<2.0 µg/dL or <26 nmol/L: Myxedema coma is possible.
Interventions

**Pretest Patient Care**

1. Explain test purpose and procedure. $T_4$ is usually the first test used in the diagnosis of hypothyroidism or hyperthyroidism, along with the TSH.
2. Have patient avoid strenuous exercise.
3. Do not administer radiopaque contrast for 1 week before testing.
4. If patient is on thyroid therapy, discontinue treatment for 1 month before testing to determine baseline values.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. See patient care for thyroid testing on page 439.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

$T_4$ values are higher in neonates due to elevated TBG.

Values rise abruptly in the first few hours after birth and decline gradually until the age of 5 years.

**Triiodothyronine ($T_3$), Total**

$T_3$ has three atoms of iodine, compared with four atoms in $T_4$. $T_3$ is more active metabolically than $T_4$, but its effect is shorter. There is much less $T_3$ than $T_4$ in the serum, and it is bound less firmly to TBG.

This measurement is a quantitative determination of the total $T_3$ concentration in the blood and is the test of choice in the diagnosis of $T_3$ thyrotoxicosis. *It is not the same as the $T_3$ uptake test that measures the unsaturated TBG in serum.* It can also be very useful in the diagnosis of hyperthyroidism. $T_3$ thyrotoxicosis refers to a variant of hyperthyroidism in which a thyrotoxic patient has elevated $T_3$ values and normal $T_4$ values. This test is not reliable in diagnosing hypothyroidism.

**Reference Values**

*Normal*

- Adults: 80–200 ng/dL or 1.2–3.1 nmol/L
- Adolescents (12–23 years): 82–213 ng/dL or 1.3–3.28 nmol/L
- Children (1–14 years): 105–245 ng/dL or 1.6–3.8 nmol/L
- Pregnancy: 116–247 ng/dL or 1.8–3.8 nmol/L

If radioimmunoassay is used, the result is reported as $T_3$ RIA.

**Procedure**

1. Obtain a 5-mL venous blood sample. Serum is needed.
2. Observe standard precautions. Place specimen in a biohazard bag.

**Clinical Implications**

1. *Increased $T_3$ values* are associated with the following conditions:
   a. Hyperthyroidism
   b. $T_3$ thyrotoxicosis (Graves' disease)
   c. Daily dosage of >25 µg of $T_3$ (Cytomel)
   d. Acute thyroiditis
   e. TBG elevation from any cause
   f. Daily dosage of >300 µg of $T_4$
   g. Early thyroid failure
   h. Thyrotoxicosis factitia
   i. Iodine deficiency goiter

2. *Decreased $T_3$ values* are associated with the following conditions:
   3. Hypothyroidism; however, some clinically hypothyroid patients will have normal levels.
   4. Starvation and state of nutrition subacute nonthyroid illness
   5. TBG decrease from any cause

**Interfering Factors**

1. Values are increased in pregnancy and with the use of drugs such as estrogens, methadone, and heroin (see Appendix J).
2. Values are decreased with the use of drugs such as anabolic steroids, androgens, large doses of salicylates, and phenytoin, nicotinic acid (see Appendix J).
3. Fasting causes $T_3$ level to decrease.

Clinical Alert

Panic values of <50 ng/dL (<0.77 nmol/L) or >300 ng/dL (>4.62 nmol/L)
Interventions

Pretest Patient Care

1. Care is the same as for T₄ testing (see page 451).
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Care is the same as for T₄ testing (see page 451).
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Triiodothyronine Uptake (T₃ U)

This test is an indirect measurement of unsaturated thyroxine-binding globulin (TBG) in blood. This determination, expressed in arbitrary terms, is inversely proportional to the TBG. For this reason, low T₃ U levels are indicative of situations that result in elevated levels of TBG uptake. For example, in hypothyroidism, when insufficient T₄ is available to produce saturation of TBG, unbound TBG (UTBG) is elevated, and T₃ U values are low. Similarly, in pregnant patients or those receiving estrogen, TBG levels are increased proportionately more than are T₄ levels, resulting in high levels of UTBG, which are reflected in low T₃ U results. This test should not be ordered alone; it is useful only when T₄ is done. It is also used to calculate the T₇ or FTI.

Reference Values
Normal 0.9–1.10 (ratio between patient specimen and the standard control) 25%–35% uptake (these are arbitrary units)

Procedure

1. Obtain a 5-mL venous blood sample. Serum is needed.
2. Observe standard precautions. Place specimen in a biohazard bag.

Clinical Implications
See Table 6.18 for implications of clinical conditions on test results.

Table 6.18 Implications of Conditions for T₃ (T₃ U) Testing

<table>
<thead>
<tr>
<th>Clinical Condition</th>
<th>T₄</th>
<th>T₃ U</th>
<th>FT₄ I</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hyperthyroid</td>
<td></td>
<td>Increased</td>
<td>Increased</td>
</tr>
<tr>
<td>Hypothyroid</td>
<td></td>
<td>Decreased</td>
<td>Decreased</td>
</tr>
<tr>
<td>Increased TBG, as in pregnancy</td>
<td></td>
<td>Increased</td>
<td>Decreased Normal</td>
</tr>
<tr>
<td>Decreased TBG, as in nephrotic syndrome</td>
<td></td>
<td>Decreased</td>
<td>Increased Normal</td>
</tr>
</tbody>
</table>

Interfering Factors

1. Decreased T₃ U levels occur in normal pregnancy, and with drugs such as estrogens, antiovulatory drugs, methadone, and heparin.
2. Increased T₃ U levels occur with drugs such as dicumarol, heparin, androgens, anabolic steroids, phenytoin, and large doses of salicylates.

Interventions

Pretest Patient Care

1. See patient care for thyroid testing (see page 437).
2. Pretest care is the same as for T₄ testing (see page 451).

Posttest Patient Aftercare

1. See patient care for thyroid testing (see page 439).
2. Posttest care is the same as for T₄ testing (see page 451).

Clinical Alert

1. This test has nothing to do with the actual T₃ blood level despite its name, which is sometimes confusingly abbreviated to the T₃ test. It is emphasized that the T₃ U and the true T₃ are entirely different tests. The T₃ U gives only an indirect measurement of overall binding.
2. This test should be used only in conjunction with the T₄ test to calculate the free thyroxine index (FTI).
3. Some methods of determining T₃ U have a direct relation with T₄. Check the reference values of your laboratory.

BIBLIOGRAPHY

Apple FS: Clinical and analytical standardization issues confronting cardiac troponin I. Clinical Chemistry 45(1): 18, 1999


Trundle D: Troublesome questions about thyroid tests. Med Lab Observ 28(10), 1996


Ward K: Exercise caution when testing the physically active. Med Lab Observ Aug 1995


Winter W: Diabetes mellitus: Pathophysiology, etiologies, complications, management and laboratory evaluations AACC, 1999 (Self-study course)

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OVERVIEW OF MICROBIOLOGIC STUDIES
Diagnostic Testing and Microbes

Microorganisms that “cause infectious disease” are defined as pathogens. Organisms that are pathogenic under one set of conditions may, under other conditions, reside within or on the surface of the body without causing disease. When organisms are present but do not cause harm to the host, they are considered commensals. When organisms multiply and cause tissue damage, they are considered pathogens, with the potential for causing or increasing a pathogenic process (Table 7.1). Many newly discovered organisms are clinically relevant. Some of these organisms, formerly
considered insignificant contaminants or commensals, have taken on roles as causative agents for opportunistic diseases in patients with human immunodeficiency virus (HIV) infection or other immunodeficiency syndromes or diseases associated with a compromised health state. Consequently, virtually any organism recovered in pure culture from a body site must be considered a potential pathogen.

### Table 7.1 Some Common Pathogens Detectable in Body Tissues and Fluids by Diagnostic Methods

<table>
<thead>
<tr>
<th>Nasopharynx and Oropharynx</th>
<th>Sputum</th>
<th>Feces</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ß-Hemolytic streptococci</strong></td>
<td>Blastomyces dermatitidis</td>
<td>Candida albicans</td>
</tr>
<tr>
<td><strong>Bordetella pertussis</strong></td>
<td>Bordetella pertussis</td>
<td>Campylobacter jejuni</td>
</tr>
<tr>
<td><strong>Mycoplasma spp.</strong></td>
<td>Candida albicans</td>
<td>Clostridium botulinum</td>
</tr>
<tr>
<td><strong>Moraxella catarrhalis</strong></td>
<td>Coccidiodes immitis</td>
<td>Entamoeba histolytica</td>
</tr>
<tr>
<td><strong>Herpes simplex virus</strong></td>
<td>Influenza viruses</td>
<td>Escherichia coli</td>
</tr>
<tr>
<td><strong>Pseudomonas spp.</strong></td>
<td>Streptococcus pneumoniae</td>
<td>Pseudomonas spp.</td>
</tr>
<tr>
<td><strong>Candida albicans</strong></td>
<td>Pseudomonas spp.</td>
<td>Salmonella spp.</td>
</tr>
<tr>
<td><strong>Corynebacterium diphtheriae</strong></td>
<td>Haemophilus influenzae</td>
<td>Shigella spp.</td>
</tr>
<tr>
<td><strong>Haemophilus influenzae</strong></td>
<td>ß-Hemolytic streptococci</td>
<td>Staphylococci</td>
</tr>
<tr>
<td><strong>Neisseria meningitidis</strong></td>
<td>Histoplasma capsulatum</td>
<td>Vibrio cholerae</td>
</tr>
<tr>
<td><strong>Streptococcus pneumoniae</strong></td>
<td>Klebsiella spp.</td>
<td>Vibrio comma</td>
</tr>
<tr>
<td><strong>Staphylococcus aureus</strong></td>
<td>Mycobacterium spp.</td>
<td>Vibrio parahaemolyticus</td>
</tr>
<tr>
<td><strong>Enterobacteriaceae</strong></td>
<td>Yersinia pestis</td>
<td>Yersinia enterocolitica</td>
</tr>
<tr>
<td><strong>Cryptococcus neoformans</strong></td>
<td>Francisella tularensis</td>
<td>Clostridium difficile</td>
</tr>
<tr>
<td><strong>Respiratory syncytial virus</strong></td>
<td>Staphylococcus aureus</td>
<td>Rotavirus</td>
</tr>
<tr>
<td><strong>Influenza viruses</strong></td>
<td>Mycoplasma spp.</td>
<td>Hepatitis A, B, and C</td>
</tr>
<tr>
<td><strong>Parainfluenza viruses</strong></td>
<td>Legionella spp.</td>
<td>Giardia lamblia</td>
</tr>
<tr>
<td><strong>Urine</strong></td>
<td>Cryptosporidium spp.</td>
<td>Crypto sporidium spp.</td>
</tr>
<tr>
<td><strong>Streptococcus agalactiae</strong></td>
<td>Pseudomonas aeruginosa</td>
<td></td>
</tr>
<tr>
<td><strong>Escherichia coli</strong>, other Enterobacteriaceae</td>
<td>Staphylococcus aureus</td>
<td></td>
</tr>
<tr>
<td><strong>Enterococcus spp.</strong></td>
<td>Enterococci, Enterobacteriaceae</td>
<td></td>
</tr>
<tr>
<td><strong>Neisseria gonorrhoeae</strong></td>
<td>Salmonella and Shigella spp.</td>
<td></td>
</tr>
<tr>
<td><strong>Mycobacterium tuberculosis</strong></td>
<td>Trichomonas vaginalis</td>
<td></td>
</tr>
<tr>
<td><strong>Skin</strong></td>
<td>Candida albicans and other yeasts</td>
<td></td>
</tr>
<tr>
<td><strong>Bacteroides spp.</strong></td>
<td>Staphylococcus epidermidis</td>
<td></td>
</tr>
<tr>
<td><strong>Clostridium spp.</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Enterobacteriaceae</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Fungi</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Pseudomonas spp.</strong></td>
<td></td>
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</tr>
<tr>
<td><strong>Streptococcus pyogenes</strong></td>
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</tr>
<tr>
<td><strong>Varicella zoster virus</strong></td>
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<tr>
<td><strong>Cerebrospinal Fluid</strong></td>
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<tr>
<td><strong>Vaginal Discharge</strong></td>
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<tr>
<td><strong>Urethral Discharge</strong></td>
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</tr>
</tbody>
</table>

### Basic Concepts of Infectious Disease

Infectious processes demonstrate observable physiologic responses to the invasion and multiplication of the offending microorganisms. Once an infectious disease is suspected, appropriate cultures should be done or nonculture techniques should be used, such as serologic testing for antigens and antibodies, monoclonal antibodies, and DNA probes. Proper specimen collection and appropriate blood and skin tests are necessary to detect and diagnose the presence of the...
Opportunity for infection depends on host resistance, organism volumes, and the ability of the organism to find a portal of entry and to overcome host defenses, invade tissues, and produce toxins. Organisms may become seated in susceptible persons through inhalation, ingestion, direct contact, inoculation, breaks in natural skin or mucous membrane barriers, changes in organism volumes, alterations in normal flora balances, or changes in other host defense mechanisms.

Host Factors

The development of an infectious disease is influenced by the patient's general health, normal defense mechanisms, previous contact with the offending organism, past clinical history, and type and location of infected tissue. Mechanisms of host resistance are detailed in the following lists:

1. Primary host defenses
   a. Anatomic barriers
      1. Intact skin surfaces
      2. Nose hairs
      3. Respiratory tract cilia
      4. Coughing and flow of respiratory tract fluids and mucus
      5. Swallowing and gastrointestinal (GI) tract peristalsis
   b. Physiologic barriers
      1. High or low pH and oxygen tension (prevents proliferation of organisms)
      2. Chemical inhibitors to bacterial growth (eg, proteases)
      3. Bile acids
      4. Active lysozymes in saliva and tears
      5. Fatty acids on skin surfaces

2. Secondary host defenses (physiologic barriers)
   a. Responses of complement, lysozymes, opsonins, and secretions
   b. Phagocytosis
   c. Immunoglobulin A (IgA), IgG, and IgM antibody formation
   d. Cell-mediated immune responses

3. Factors decreasing host resistance
   a. Age: the very young and the very old are more susceptible
   b. Presence of chronic disease (eg, cancer, cardiovascular disease, diabetes)
   c. Use or history of certain therapeutic modalities, such as radiation, chemotherapy, corticosteroids, antibiotics, or immunosuppressants
   d. Toxins, including alcohol, street drugs, legitimate therapeutic drugs, venom or toxic secretions from a reptile or insect, or other nonhuman bites or punctures
   e. Others, including excessive physical or emotional stress states, nutritional state, and presence of foreign material at the site

COLLECTION AND TRANSPORT OF SPECIMENS

General Principles

The health care professional is responsible for collecting specimens for diagnostic examinations. Because procedures vary, check institutional protocols for specimen retrieval, transport, preservation, and reporting of test results.

Specimens for bacterial culture should be representative of the disease process. Also, sufficient material must be collected to ensure an accurate examination. As an example, serous drainage from a diabetic foot ulcer with possible osteomyelitis may yield inaccurate results. In this case, a bone biopsy or purulent drainage of infected tissue would be a better specimen. Likewise, if there is a lesion of the skin and subcutaneous tissue, material from the margin of the lesion rather than the central part of the lesion would be more desirable. If a purulent sputum sample cannot be obtained to aid in the diagnosis of pneumonia, blood cultures, pleural fluid examination, and bronchoalveolar lavage (BAL) specimens are also acceptable.

It is imperative that material be collected where the suspected organism is most likely to be found, with as little contamination from normal flora as possible. For this reason, certain precautions must be followed routinely:

1. Observe standard precautions. Clean the skin starting centrally and going out in larger circles. Repeat several times, using a clean swab or wipe each time. If 70% alcohol is used, it should be applied for 2 minutes. Tincture of iodine requires only 1 minute of cleansing.
2. Bypass areas of normal flora; culture only for a specific pathogen.
3. Collect fluids, tissues, skin scrapings, and urine in sterile containers with tight-fitting lids. Polyester-tipped swabs in a collection system containing an ampule of Stuart's transport medium ensure adequacy of the specimen for 72 hours at room temperature.
4. Place the specimen in a biohazard bag.
Clinical Alert

1. Without routine precautions for collecting and handling specimens, the patient's condition may be incorrectly diagnosed, laboratory time may be wasted, effective treatment may be delayed, or pathogenic organisms may be transmitted to health care workers and other patients.

2. It is important to report all identified diseases, conditions, and outbreaks according to state and federal guidelines.

3. Traditional surveillance requires that a disease be diagnosed before it is reported. The newer reporting system from the CDC requires no waiting for a definite diagnosis, but relies on collecting syndrome information of signs and symptoms reported to physicians and clinics. The syndrome system compares early reports of fever, headaches, diarrhea, vomiting, rashes, and normal seasonal, daily, and required fluctuations in known illness. The computerized network is designated to provide early warning of infectious disease terrorism attack by monitoring visits to physician's offices, emergency rooms, and drugstores (for remedies). Both systems are used. The collaborative team approach relies on asking patient what signs or symptoms are present before doing testing for infectious and reportable diseases.

4. In addition, the Environmental Protection Agency (EPA) monitors air quality in major cities. This surveillance system is designed to provide 24-hour notice of release into the air of deadly pathogens (eg, smallpox, anthrax).

Footnote

Sources of Specimens

Microbiologic specimens may be collected from many sources, such as blood, pus or wound exudates or drainage, urine, sputum, feces, genital discharges or secretions, cerebrospinal fluid (CSF), and eye or ear drainage. During specimen collection, these general procedures should be followed:

1. Label specimens properly with the following information (institutional requirements may vary):
   a. Patient's name, age, sex, address, hospital identification number, and physician's full name
   b. Specimen source (eg, throat, conjunctiva)
   c. Time of collection
   d. Specific studies ordered
   e. Clinical diagnosis; suspected microorganisms
   f. Patient's history
   g. Patient's immune state
   h. Previous and current infections
   i. Previous or current antibiotic therapy
   j. Isolation status—state type of isolation (eg, contacts, respiratory, wound)
   k. Other requested information pertinent to testing

2. Avoid contaminating the specimen; maintain aseptic or sterile technique as required:
   a. Special supplies may be required:
      1. For anaerobes, sterile syringe aspiration of pus or other body fluid
      2. Anaerobic transport containers for tissue specimens
   b. Sterile specimen containers
   c. Precautions to take during specimen collection include:
      a. Care to maintain clean outside container surfaces
      b. Use of appropriately fitting covers or plugs for specimen tubes and bottles
      c. Replacement of sterile plugs and caps that have become contaminated
      d. Observation of standard precautions

3. Ensure the preservation of specimens by delivering them promptly to the laboratory. Many specimens may be refrigerated (not frozen) for a few hours without any adverse effects. Note the following exceptions:
   a. Urine culture samples must be refrigerated.
   b. CSF specimens should be transported to the laboratory as soon as possible. If this is problematic, the culture should be incubated (meningococci do not withstand refrigeration).

4. Transport specimens quickly to the laboratory to prevent desiccation of the specimen and death of the microorganisms.
   a. For anaerobic cultures, no more than 10 minutes should elapse between time of collection and culture.
   b. Feces suspected of harboring Salmonella or Shigella organisms should be placed in a special transport medium, such as Cary-Blair, if culturing of the specimen will be delayed greater than 30 minutes.

5. Ensure that specimen quantity is adequate. With few exceptions, the quantity of the specimen should be as large as possible. When only a small quantity is available, swabs should be moistened with sterile saline just before collection, especially for nasopharyngeal cultures.

6. Handle specimen collection in the following way:
   a. Submit entire fluid specimen collected. Do not submit fluids on swabs.
   b. Whenever possible, specimens should be collected before antibiotic regimens are instituted; for example, complete all blood culture sampling before starting antibiotic therapy.
   c. Collection must be geared to the rise in symptoms such as fever. (The practitioner should be familiar with the clinical course of the suspected disease.)

Transport of Specimens by Mail

Several kits containing transport media are available for use when there is a significant delay between collection and
culturings. Culture swabs (containing transport medium) are available for bacterial, viral, and anaerobic collection of specimens. Some laboratories provide Cary-Blair and polyvinyl alcohol (PVA) fixative transport vials for stool collection for culture and ova and parasite examination. Depending on the request, some specimens may have to be shipped in a Styrofoam box with refrigerant packs. This is especially true for specimens to be tested for viral examination. It is prudent to consult the reference laboratory to which specimens will be sent for information on proper collection and shipment.

According to the Code of Federal Regulations (49 CFR), a viable organism or its toxin or a diagnostic specimen (volume < 50 mL) must be placed in a secure, closed, watertight container that is then enclosed in a second secure, watertight container. Biohazard labels should be placed on the outside of the container.

Specimens that are to be transported within an institution should be placed in a sealed biohazard bag. Ideally, the requisition should accompany the specimen but not be sealed inside the bag.

### Diagnosis of Bacterial Disease

Bacteriologic studies attempt to identify the specific organism causing an infection (Table 7.2). This organism may be specific to one disease, such as *Mycobacterium tuberculosis* for tuberculosis (TB), or it may cause a variety of infections, such as those associated with *Staphylococcus aureus*. Antibiotic susceptibility studies then determine the responses of the specific organism to various classes and types of antibiotics. An antibiotic that inhibits bacterial growth is the logical choice for treating the infection.

#### Table 7.2 Bacterial Diseases and Their Laboratory Diagnosis

<table>
<thead>
<tr>
<th>Disease</th>
<th>Causative Organism</th>
<th>Source of Specimen</th>
<th>Diagnostic Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anthrax</td>
<td><em>Bacillus anthracis</em></td>
<td>Blood, sputum, skin</td>
<td>Blood, sputum, skin smear and culture</td>
</tr>
<tr>
<td>Brucellosis (undulant fever)</td>
<td><em>Brucella melitensis, Brucella abortus, Brucella suis</em></td>
<td>Blood, bone marrow, CSF, tissue, lymph node, urine</td>
<td>Culture, serology</td>
</tr>
<tr>
<td>Bubonic plague</td>
<td><em>Yersinia pestis</em></td>
<td>Buboes (enlarged and inflamed lymph nodes), blood, sputum</td>
<td>Skin, blood, and sputum smear; culture</td>
</tr>
<tr>
<td>Chancre</td>
<td><em>Haemophilus ducreyi</em></td>
<td>Genital lesion</td>
<td>Lesion smear and culture; biopsy</td>
</tr>
<tr>
<td>Cholera</td>
<td><em>Vibrio cholerae</em></td>
<td>Feces</td>
<td>Stool smear and culture</td>
</tr>
<tr>
<td>Psittacosis</td>
<td><em>Chlamydia psittaci</em></td>
<td>Blood, sputum, lung tissue</td>
<td>Culture, smear, serologic tests</td>
</tr>
<tr>
<td>Diphtheria</td>
<td><em>Corynebacterium diphtheriae</em></td>
<td>Nasopharynx</td>
<td>Nasopharyngeal smear and culture</td>
</tr>
<tr>
<td>Erysipeloïd</td>
<td><em>Erysipelothrix rhusiopathiae</em></td>
<td>Lesion, blood</td>
<td>Smear and culture</td>
</tr>
<tr>
<td>Gonorrhrea</td>
<td><em>Neisseria gonorrhoeae</em></td>
<td>Cervix, blood</td>
<td>Smear and culture</td>
</tr>
<tr>
<td>Granuloma inguinale (donovanosis)</td>
<td><em>Calymmatobacterium granulomatis</em></td>
<td>Groin lesion</td>
<td>Smear</td>
</tr>
<tr>
<td>Gastritis, gastric ulcer</td>
<td><em>Helicobacter pylori</em></td>
<td>Gastric tissue biopsy</td>
<td>Culture, biopsy</td>
</tr>
<tr>
<td>Relapsing fever</td>
<td><em>Borrelia recurrentis</em></td>
<td>Peripheral blood</td>
<td>Direct examination</td>
</tr>
<tr>
<td>Lyme disease</td>
<td><em>Borrelia burgdorferi</em></td>
<td>Blood, CSF, skin lesion</td>
<td>Serologic test</td>
</tr>
<tr>
<td>Legionnaire's disease</td>
<td><em>Legionella pneumophila</em></td>
<td>Sputum</td>
<td>Culture, direct fluorescent antibody; serologic test, urine antigen test</td>
</tr>
<tr>
<td>Leprosy (Hansen's disease)</td>
<td><em>Mycobacterium leprae</em></td>
<td>Skin scrapings</td>
<td>Skin smear, biopsy</td>
</tr>
<tr>
<td>Lymphogranuloma venereum</td>
<td><em>Chlamydia trachomatis</em></td>
<td>Genital, conjunctiva, urethra, urine</td>
<td>Culture, DNA probe, amplified test</td>
</tr>
<tr>
<td>Listeriosis</td>
<td><em>Listeria monocytogenes</em></td>
<td>Stool, blood, CSF, amniotic fluid, placenta, vagina</td>
<td>Smears and culture, serologic test</td>
</tr>
<tr>
<td>Pneumonia</td>
<td><em>Haemophilus influenzae</em>, <em>Klebsiella pneumoniae</em>, <em>Staphylococcus aureus</em>, <em>Streptococcus pneumoniae</em></td>
<td>Bronchoscopy, secretions, sputum, blood, lung aspirate or biopsy, pleural fluid</td>
<td>Smear and culture</td>
</tr>
<tr>
<td>Strep throat, scarlet fever, impetigo</td>
<td><em>Streptococcus pyogenes</em></td>
<td>Throat, lesion</td>
<td>Culture, serologic test</td>
</tr>
<tr>
<td>Tetanus</td>
<td><em>Clostridium tetani</em></td>
<td>Wound</td>
<td>Smear and culture</td>
</tr>
<tr>
<td>Toxic shock syndrome</td>
<td><em>Staphylococcus aureus</em></td>
<td>Tissue</td>
<td>Culture</td>
</tr>
<tr>
<td>Tuberculosis</td>
<td><em>Mycobacterium tuberculosis</em></td>
<td>Sputum, gastric washings, urine, CSF</td>
<td>Smear and culture of sputum, gastric washings, urine, and CSF; skin test</td>
</tr>
<tr>
<td>Tularemia</td>
<td><em>Francisella tularensis</em></td>
<td>Skin, lymph node, ulcer tissue biopsy, sputum, bone marrow</td>
<td>Serologic test, smear and culture</td>
</tr>
</tbody>
</table>
Typhoid  
*Salmonella typhi*
- Blood (after first week of infection); feces (after second week of infection)
  - Culture and serologic test

Whooping cough  
*Bordetella pertussis*
- Nasopharyngeal swab
  - Culture, fluorescent antibody test, PCR, serologic test

Nocardiosis  
*Nocardia asteroides*
- Sputum, lesion
  - Smear and culture

Mycoplasma  
*Mycoplasma pneumoniae*
- Sputum, nasopharyngeal and throat swabs
  - Serology, culture, PCR

Some questions that need to be asked when searching for bacteria as the cause of a disease process include the following: (1) Are bacteria responsible for this disease? (2) Is antimicrobial therapy indicated? Most bacteria-related diseases have a febrile course. From a practical standpoint during evaluation of the febrile patient, the sooner a diagnosis can be reached and the sooner a decision can be made concerning antimicrobial therapy, the less protracted the period of recovery.

Anaerobic bacterial infections are commonly associated with localized necrotic abscesses: they may yield several different strains of bacteria. Because of this, the term *polymicrobial disease* is sometimes used to refer to anaerobic bacterial diseases. This view is in sharp contrast to the "one organism, one disease" concept that characterizes other infections, such as typhoid fever, cholera, or diphtheria. Isolation and identification of the different strains of anaerobic bacteria through sensitivity studies is desirable so that appropriate therapy may be given.

**Studies of the Susceptibility of Bacteria to Antimicrobial Agents**

The susceptibility test detects the type and amount of antibiotic or chemotherapeutic agent required to inhibit bacteria growth. Often, culture and susceptibility tests are ordered together. Susceptibility studies also may be indicated when an established regimen or treatment is to be altered.

A common and useful test for evaluating antibiotic susceptibility is the disk diffusion method. A set of antibiotic-impregnated disks on agar is inoculated with a culture derived from the specific bacteria being tested. After a suitable period of incubation, the degree of bacterial growth within the different antibiotic zones on the disks is determined and measured. Growth zone diameters, measured in millimeters, are correlated to the minimum inhibitory concentration (MIC) to determine whether the organism is truly susceptible to the antibiotic. Another method is a broth dilution test. The organism is grown in the presence of doubling dilutions of the antibiotic. The lowest concentration of the antibiotic that inhibits the organism's growth is the MIC. Many commercial systems are based on this method.

**Clinical Implications**

1. The terms *sensitive* and *susceptible* imply that an infection caused by the bacterial strain tested will respond favorably in the presence of the indicated antimicrobial agent.
2. The terms *intermediate, partially resistant, and moderately susceptible* mean that the bacterial strain tested is not completely inhibited by therapeutic concentrations of a test drug.
3. *Indeterminate* means that the bacteria has an MIC that approaches achievable blood and tissue concentrations. It implies clinical efficacy in body sites where the antibiotic is physiologically concentrated. The intermediate category also includes a buffer zone, which should prevent major errors due to technical factors.
4. The term *resistant* implies that the organism is not inhibited by the antibiotic.
5. Some antimicrobial agents act in a *bactericidal* manner, meaning that they kill the organism. Others act in a *bacteriostatic* manner, meaning that they inhibit growth of the organism but do not necessarily kill it.
   a. Bactericidal agents
      1. Aminoglycoside
      2. Cephalosporins
      3. Metronidazole
      4. Penicillins
      5. Quinolones
      6. Rifampin
      7. Vancomycin
   b. Bacteriostatic agents
      1. Chloramphenicol
      2. Erythromycin
      3. Sulfonamides
      4. Tetracycline
6. Emergence of strains of penicillin-resistant *Neisseria gonorrhoeae*, methicillin (or oxacillin)-resistant *S. aureus*, amikacin-resistant *Pseudomonas* spp. or other gram-negative rods, and vancomycin-resistant *Enterococcus* spp. present challenges to the clinician in regard to treatment. Many hospitals screen for methicillin-resistant *S. aureus* (MRSA) and vancomycin-resistant *Enterococcus* (VRE) species so as to isolate patients infected with these organisms.

**Diagnosis of Mycobacterial Infections**

The genus *Mycobacterium* contains several species of bacteria that are pathogenic to humans (Table 7.3). For example, *M. tuberculosis* is spread from person to person through inhalation of airborne respiratory secretions containing mycobacteria expelled during coughing, sneezing, or talking. In patients with the acquired immunodeficiency syndrome (AIDS), *Mycobacterium avium–intracellulare* (MAI) complex is acquired through the GI tract, often through ingestion of
contaminated water or food.

<table>
<thead>
<tr>
<th>Causative Organism</th>
<th>Source of Specimen</th>
<th>Diagnostic Test</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Mycobacterium tuberculosis</em></td>
<td>Sputum, urine, CSF, tissue, bone marrow</td>
<td>Culture and smear; skin test; DNA probe</td>
</tr>
<tr>
<td><em>Mycobacterium avium–intracellulare</em></td>
<td>Sputum, stool, CSF, tissue, blood, semen, lymph nodes</td>
<td>Culture and smear; DNA probe</td>
</tr>
<tr>
<td><em>Mycobacterium kansasii</em></td>
<td>Skin, joint, lymph nodes, sputum, tissue</td>
<td>Culture and smear</td>
</tr>
<tr>
<td><em>Mycobacterium leprae</em></td>
<td>CSF, skin, bone marrow, lymph nodes</td>
<td>Histopathologic examination of lesion</td>
</tr>
<tr>
<td><em>Mycobacterium marinum</em></td>
<td>Joint lesion</td>
<td>Culture and smear</td>
</tr>
<tr>
<td><em>Mycobacterium xenopi</em></td>
<td>Surgical wound, bone, joint, tissue, sputum</td>
<td>Culture and smear</td>
</tr>
<tr>
<td><em>Mycobacterium chelonei</em></td>
<td>Surgical wound, sputum, tissue</td>
<td>Culture and smear</td>
</tr>
</tbody>
</table>

The disease progression of mycobacteriosis, particularly in patients with AIDS, is rapid (a few weeks). This short time span has required new methods for rapid recovery and identification of mycobacteria so that antibiotic therapy can be instituted promptly. These newer techniques involve the use of instruments that shorten the growth period for mycobacteria to 1 to 2 weeks. Isotopic nucleic acid probes are available for culture identification of *M. tuberculosis*, MAI complex, *Mycobacterium kansasii*, and *Mycobacterium gordonae*. Polymerase chain reaction (PCR) techniques, which use DNA technology to detect mycobacteria directly in clinical specimens, is also available to clinical laboratories.

A disturbing problem that has arisen since the resurgence of TB among persons with AIDS is the appearance of multidrug-resistant *M. tuberculosis* strains.

### Collection of Specimens

1. Sputum and bronchial aspirates and lavages are the best samples for diagnosis of pulmonary infection. Purulent sputum (5 to 10 mL) from the first productive cough of the morning should be expectorated into a sterile container. If the specimen is not processed immediately, it should be refrigerated. Pooled specimens collected over several hours are not acceptable. For best results, three specimens should be collected over several days. A prerequisite of good specimen collection is the use of sterile, sturdy, leak-proof containers placed into biohazard bags.

2. If the patient is unable to produce sputum, an early-morning gastric sample may be aspirated and cultured. This specimen must be hand-delivered to the laboratory to be processed or neutralized immediately.

3. Patients with suspected renal disease should provide early-morning urine specimens collected for 3 days in a row. Pooled 24-hour urine collections are not recommended. Unless processed immediately, the specimen should be refrigerated.

4. If TB meningitis is suspected, at least 10 mL of CSF should be obtained.

5. Sterile body fluids, tissue biopsy samples, and material aspirated from skin lesions are acceptable specimens for mycobacterial cultures. Tissue should be placed in a neutral transport medium to avoid desiccation. Swab specimens are not suitable for mycobacterial culture.

6. Feces are commonly the first specimens from which MAI complex can be isolated in a patient with disseminated disease. An acid-fast stain is usually performed directly. Culture is performed only if the smear tests positive.

7. MAI complex organisms can also be isolated from the blood of immunosuppressed patients.

### Diagnosis of Rickettsial Disease

Rickettsiae are small, gram-negative coccobacilli that structurally resemble bacteria but are one tenth to one half as large. Polychromatic stains (Giemsa stain) are better than simple stains of the Gram stain for demonstrating rickettsiae in cells.

Rickettsiosis is the general name given to any disease caused by rickettsiae (Table 7.4). These organisms are considered to be obligate intracellular parasites; that is, they cannot exist anywhere except inside the bodies of living organisms. Diseases caused by rickettsiae are transmitted by *arthropod vectors*, such as lice, fleas, ticks, or mites. Rickettsial diseases are divided into the following general groups:

<table>
<thead>
<tr>
<th>Disease</th>
<th>Geographic Distribution</th>
<th>Natural Cycle</th>
<th>Transmission to Humans</th>
<th>Serologic Diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Typhus epidemic</td>
<td><em>Rickettsia prowazekii</em></td>
<td>Worldwide</td>
<td>Body louse</td>
<td>Human infected louse feces into broken skin</td>
</tr>
<tr>
<td>Endemic (murine)</td>
<td><em>Rickettsia typhi</em></td>
<td>Worldwide</td>
<td>Flea</td>
<td>Rodents</td>
</tr>
</tbody>
</table>
Spotted fever, Rocky Mountain spotted fever

Rocky Mountain spotted fever, caused by *Rickettsia rickettsii*, is characterized by an acute febrile illness, severe headache, and possibly pneumonia or hepatitis. It can cause encephalitis in children and has been isolated in breast milk and in the placenta of infected mothers, making it possible for a fetus to be infected in utero. Both complement fixation and fluorescent antibody tests can detect antibodies to the organism. *R. rickettsii* displays an antigenic variation during an infection. Phase I antibodies are preponderant during the chronic phase, whereas phase II antibodies predominate during the acute phase. A diagnosis is made when the phase I titer in a convalescent serum specimen is four times greater than that in an acute serum specimen.

Early diagnosis of rickettsial infection is usually based on observation of clinical symptoms such as fever, rash, and exposure to ticks. Biopsy specimens of skin tissue from a patient with suspected Rocky Mountain spotted fever can be tested with an immunofluorescent stain and diagnosed 3 to 4 days after symptoms appear. Signs and symptoms include:

1. Fever
2. Skin rashes
3. Parasitism of blood vessels
4. Prostration
5. Stupor and coma
6. Headache
7. Ringing in the ears
8. Dizziness

North Asian tick-borne rickettsiosis

North Asian tick-borne rickettsiosis is caused by *Rickettsia sibirica* and is characterized by fever, headache, and rash. It is typically transmitted by ticks, and serologic tests—IFA or latex agglutination—are used to diagnose the disease.

Buotonneuse fever

Buotonneuse fever, caused by *Rickettsia conorii*, is characterized by fever, headache, and rash. It is typically transmitted by ticks, and serologic tests—IFA or latex agglutination—are used to diagnose the disease.

Queensland tick typhus

Queensland tick typhus is caused by *Rickettsia australis* and is characterized by fever, headache, and rash. It is typically transmitted by ticks, and serologic tests—IFA or latex agglutination—are used to diagnose the disease.

Rickettsial pox

Rickettsial pox is caused by *Rickettsia akari* and is characterized by fever, headache, and rash. It is typically transmitted by mites, and serologic tests—IFA or latex agglutination—are used to diagnose the disease.

Scrub typhus

Scrub typhus is caused by *Rickettsia tsutsugamushi* and is characterized by fever, headache, and rash. It is typically transmitted by trombiculid mites, and serologic tests—IFA or latex agglutination—are used to diagnose the disease.

Ehrlichiosis

Ehrlichiosis is caused by *Ehrlichia canis* and *Ehrlichia sennetsu* and is characterized by fever, headache, and rash. It is typically transmitted by ticks, and serologic tests—IFA or latex agglutination—are used to diagnose the disease.

Q fever

Q fever, caused by *Coxiella burnetii*, is characterized by an acute febrile illness, severe headache, rigors, and possibly pneumonia or hepatitis. It can cause encephalitis in children and has been isolated in breast milk and in the placenta of infected mothers, making it possible for a fetus to be infected in utero. Both complement fixation and fluorescent antibody tests can detect antibodies to the organism. *C. burnetii* displays an antigenic variation during an infection. Phase I antibodies are preponderant during the chronic phase, whereas phase II antibodies predominate during the acute phase. A diagnosis is made when the phase I titer in a convalescent serum specimen is four times greater than that in an acute serum specimen.

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1. Fever
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3. Parasitism of blood vessels
4. Prostration
5. Stupor and coma
6. Headache
7. Ringing in the ears
8. Dizziness

1. Typhus-like fevers
2. Spotted fever
3. Scrub typhus
4. Q fever
5. Other rickettsial diseases

Q fever, caused by *Coxiella burnetii*, is characterized by an acute febrile illness, severe headache, rigors, and possibly pneumonia or hepatitis. It can cause encephalitis in children and has been isolated in breast milk and in the placenta of infected mothers, making it possible for a fetus to be infected in utero. Both complement fixation and fluorescent antibody tests can detect antibodies to the organism. *C. burnetii* displays an antigenic variation during an infection. Phase I antibodies are preponderant during the chronic phase, whereas phase II antibodies predominate during the acute phase. A diagnosis is made when the phase I titer in a convalescent serum specimen is four times greater than that in an acute serum specimen.
**NOTE**

Rickettsial diseases are often characterized by an incubation period of 10 to 14 days, followed by an abrupt onset of the signs and symptoms listed, in a patient with a history of arthropod bites. Cultures of rickettsia are performed only in reference laboratories. Rickettsial infections usually are diagnosed by serologic methods, using acute and convalescent serum specimens. A fourfold rise in serum antibody titer is preferable, but a single titer greater than 1:64 is highly suggestive of infection (see Chap. 8).

### Diagnosis of Parasitic Disease

About 70 species of animal parasites commonly infect the human body (Table 7.5). More than half of these can be detected by examination of stool specimens because the parasites inhabit the GI tract and its environs. Of the parasites that can be diagnosed by stool examinations, about one third are single-celled protozoa, and two thirds are multicellular worms. Only six or seven types of intestinal protozoa are clinically important, but almost all of the worm classes are potentially pathogenic.

<table>
<thead>
<tr>
<th>Disease</th>
<th>Causative Organism</th>
<th>Source of Specimen</th>
<th>Diagnostic Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amebiasis</td>
<td>Entamoeba histolytica</td>
<td>Stool, liver</td>
<td>Stool smear, rectal biopsy, serologic test</td>
</tr>
<tr>
<td>Ascariasis</td>
<td>Ascaris lumbricoides</td>
<td>Stool, sputum</td>
<td>Ova and parasite examination, antigen test, rectal biopsy, serologic test</td>
</tr>
<tr>
<td>Cestodiasis of intestine (tapeworm disease)</td>
<td>Taenia saginatus, Taenia solium, Dipylidium caninum, Hymenolepis nana, Hymenolepis diminuta</td>
<td>Stool</td>
<td>Ova and parasite examination, Scotch tape test for Enterobius vermicularis</td>
</tr>
<tr>
<td>Chagas’ disease</td>
<td>Trypanosoma cruzi</td>
<td>Blood, spinal fluid</td>
<td>Giemsa- or Wright-Giemsa–stained smear</td>
</tr>
<tr>
<td>Cryptosporidiosis</td>
<td>Cryptosporidium parvum</td>
<td>Stool, lung, gallbladder</td>
<td>Ova and parasite examination, antigen test, direct fluorescent antibody test</td>
</tr>
<tr>
<td>Cysticercosis</td>
<td>Taenia solium larvae</td>
<td>Muscle and brain</td>
<td>Muscle and brain cyst biopsy</td>
</tr>
<tr>
<td>Echinococcosis</td>
<td>Echinococcus granulosus</td>
<td>Sputum and urin, liver, spleen</td>
<td>Ova and parasite examination, direct microscopic examination, serologic test, Casoni's skin test; liver and bone biopsy</td>
</tr>
<tr>
<td>Enterobiasis (pinworm disease)</td>
<td>Enterobius vermicularis</td>
<td>Stool</td>
<td>Scotch tape test</td>
</tr>
<tr>
<td>Filariasis</td>
<td>Wuchereria bancrofti, Brugia malayi, Loa loa</td>
<td>Blood</td>
<td>Blood smear, lymph node biopsy, serologic test</td>
</tr>
<tr>
<td>Giardiasis</td>
<td>Giardia lamblia</td>
<td>Stool, duodenal aspirate or biopsy</td>
<td>Ova and parasite examination, antigen test, direct fluorescent antibody test, microscopic examination of Entero test</td>
</tr>
<tr>
<td>Hookworm</td>
<td>Ancylostoma duodenale, Necator americanus</td>
<td>Stool</td>
<td>Ova and parasite examination</td>
</tr>
<tr>
<td>Isospora</td>
<td>Isospora belli</td>
<td>Stool</td>
<td>Ova and parasite examination</td>
</tr>
<tr>
<td>Kala-azar</td>
<td>Leishmania donovani</td>
<td>Liver, bone marrow, blood</td>
<td>Giemsa- or Wright-Giemsa–stained smear and culture, lymph node and spleen biopsy</td>
</tr>
<tr>
<td>Malaria</td>
<td>Plasmodium falciparum, Plasmodium malariae, Plasmodium vivax, Plasmodium ovale</td>
<td>Blood, bone marrow</td>
<td>Giemsa- or Wright-Giemsa–stained smear</td>
</tr>
<tr>
<td>Acanthamoebiasis</td>
<td>Acanthamoeba culbertsoni</td>
<td>CSF, corneal biopsy or scraping</td>
<td>Smear and tissue culture</td>
</tr>
<tr>
<td>Naegleriiosis</td>
<td>Naegleria fowleri</td>
<td>CSF</td>
<td>Smear</td>
</tr>
<tr>
<td>Sarcocystis</td>
<td>Sarcocystis hominis or Sarcocystis suihominis</td>
<td>Stool</td>
<td>Ova and parasite examination</td>
</tr>
<tr>
<td>Blastocystis</td>
<td>Blastocystis hominis</td>
<td>Stool</td>
<td>Ova and parasite examination</td>
</tr>
<tr>
<td>Onchocerciasis</td>
<td>Onchocerca volvulus</td>
<td>Skin</td>
<td>Skin biopsy</td>
</tr>
<tr>
<td>Paragonimiasis</td>
<td>Paragonimus westermani</td>
<td>Sputum, stool</td>
<td>Ova and parasite examination; serologic test, skin test</td>
</tr>
<tr>
<td>Scabies</td>
<td>Sarcoptes scabiei</td>
<td>Skin</td>
<td>Skin smear, direct examination</td>
</tr>
<tr>
<td>Schistosomiasis of intestine and bladder</td>
<td>Schistosoma mansoni, Schistosoma japonicum, Schistosoma haematobium</td>
<td>Stool, urine</td>
<td>Ova and parasite examination, serologic test; skin test; rectal, bladder, and liver biopsy</td>
</tr>
<tr>
<td>Strongyloidiasis</td>
<td>Strongyloides stercoralis</td>
<td>Stool, duodenal aspirate</td>
<td>Ova and parasite examination, serologic test</td>
</tr>
<tr>
<td>Toxoplasmosis</td>
<td>Toxoplasma gondii</td>
<td>Blood, tissue, CSF</td>
<td>Serologic test, tissue smear, biopsy</td>
</tr>
</tbody>
</table>
Diagnosis of parasites begins with ova and parasite examination. Other diagnostic options include sigmoidoscopy, biopsies, barium radiologic studies, and serologic tests. Collection of fecal specimens for parasites should be done before administration of barium sulfate, mineral oil, bismuth, antimalarial drugs, and some antibiotics (eg, tetracycline). For ova and parasite examination, ideally, one specimen should be collected every other day for a total of three specimens. At the most, these specimens should be gathered within 10 days.

For detection of Giardia, other diagnostic tests such as the Entero-Test capsule (string test) and duodenal aspiration or biopsy may be necessary. The Entero-Test consists of a gelatin capsule containing a coiled length of nylon yarn. The capsule is swallowed, the gelatin dissolves, and the weighted string is carried into the duodenum. After about 4 hours, the string is withdrawn, and the accompanying mucus is examined microscopically for Giardia. Duodenal fluid also can be submitted by the physician to be examined for Giardia and Strongyloides stercolaris. The specimen should contain no preservatives and should be examined for organisms within 1 hour after collection.

Cryptosporidium parvum has long been recognized as an animal parasite but is also capable of infecting humans, especially physically compromised patients. Organisms have been recovered from the gallbladder, the lungs, and the stool.

Collection of Specimens

1. Multiple specimens may be necessary to detect a parasitic infection.
2. Most parasites found in humans are identified in blood or feces but may also be evident in urine, sputum, tissue fluids, or biopsy tissues.
3. Fecal specimens should not be contaminated with water or urine. All specimens should be labeled with the patient's name, clinician's name, identification number (if applicable), and date and time collected. Various commercial collection systems are available to allow collection of specimens at home, in a nursing institution, or in a hospital setting. Clear instructions should be communicated and given in writing to the patient to ensure proper collection. See Chapter 4, Stool Studies, for more information.
4. When sputum is collected for ova and parasites, it should be "deep sputum" from the lower respiratory tract. It should be collected early in the morning, before the patient eats or brushes the teeth, and immediately delivered to the laboratory. See Appendix E, Guidelines for Specimen Transport and Storage, for more information.

Clinical Considerations

1. General considerations
   a. Eosinophilia is considered a definite indicator of parasitic infection. Protozoa also may produce associated eosinophilia.
   b. Protozoa and helminths, particularly larvae, may be found in organs, tissues, and blood.
2. Specimen-related considerations
   a. Hepatic puncture can reveal visceral leishmaniasis. Liver biopsy may yield toxocarial larvae and schistosomal worms and eggs. Hepatic abscess material from the peripheral area may reveal more organisms than the necrotic center.
   b. Bone marrow may be positive for trypansomiasis and malaria when blood samples produce negative results. Bone marrow specimens are obtained through puncture of the sternum, iliac crest, vertebral processes, trochanter, or tibia.
   c. Puncture or biopsy samples from a lymph node may be examined for the presence of trypansomiasis, leishmaniasis, toxoplasmosis, and filariasis.
   d. Mucous membrane lesion or skin samples may be obtained through scraping, needle aspiration, or biopsy.
   e. CSF may contain trypanosomes and Toxoplasma organisms.
   f. Sputum may reveal Paragonimus westermani (lung fluke) eggs. Occasionally, the larvae and hookworm of S. stercolaris or Ascariis lumbricoides may be expectorated during pulmonary migration. In pulmonary echinococcosis (hydatid disease), hydatid cyst contents may be found in sputum.
   g. Specimens taken from cutaneous ulcers should be aspirated below the ulcer bed rather than at the surface. A few drops of saline may be introduced by needle and syringe to aspirate the intracellular leishmanial organisms.
   h. Conjunctival scrapings or biopsy specimens can be examined histologically or cultured for the presence of Acanthamoeba. This organism is rare but can cause keratitis among contact lens wearers.
   i.Films for blood parasites are usually prepared when the patient is admitted. Samples should be taken at 6- to 18-hour intervals for at least 3 successive days.

Diagnosis of Fungal Disease

Fungal diseases, also known as mycoses, are believed to be more common now than in the past because of increased use of antibacterial and immunosuppressive drugs (Table 7.6). Fungi prefer the debilitated host, the person with chronic
Of more than 200,000 species of fungi, approximately 200 species are generally recognized as being pathogenic for humans. Fungi live in soil enriched by decaying nitrogenous matter and are capable of maintaining a separate existence through a parasitic cycle in humans or animals. The systemic mycoses are not communicable in the usual sense of human-to-human or animal-to-animal transfer. Humans become accidental hosts through inhalation of spores or by introduction of spores into tissues through trauma. Altered susceptibility may result in fungal lesions; this frequently occurs in patients who have a debilitating disease, diabetes, or impaired immunologic responses due to steroid or antimetabolite therapy. Prolonged administration of antibiotics can result in a fungal superinfection.

Fungal diseases may be classified according to the type of tissues involved:

1. **Dermatophytoses** include superficial and cutaneous mycoses, such as athlete’s foot, ringworm, and “jock itch.” Species of *Microsporum*, *Epidermophyton*, and *Trichophyton* are the causative organisms.
2. **Subcutaneous mycoses** involve the subcutaneous tissues and muscles.
3. **Systemic mycoses** involve the deep tissues and organs and are the most serious of the three groups.

Amphotericin B, introduced into practice in 1958, was for many years the only drug available to treat invasive fungal infections. Now ketoconazole, fluconazole, itraconazole, and lipid formulations of amphotericin B provide alternative choices when treatment of fungal disease is warranted.

### Collection of Hair and Skin Specimens

1. Clean the suspected area with 70% alcohol to remove bacteria. Use sterile techniques and standard precautions.
2. Scrape the peripheral erythematous margin of putative “ringworm” lesions with a sterile scalpel or wooden spatula and place the scrapings in a covered sterile container.
3. Clip samples of infected scalp or beard hair and place in a covered sterile container.

<table>
<thead>
<tr>
<th>Disease</th>
<th>Causative Organism</th>
<th>Source of Specimen</th>
<th>Diagnostic Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aspergillosis</td>
<td><em>Aspergillus fumigatus</em>, <em>Aspergillus flavus</em>, <em>Aspergillus terreus</em></td>
<td>Sputum, tissue, ear, corneal scraping</td>
<td>Smear and culture, serologic test, chest x-ray, computed tomography</td>
</tr>
<tr>
<td>Blastomycosis</td>
<td><em>Blastomyces dermatitidis</em></td>
<td>Skin lesion, sputum, bone, joint</td>
<td>Smear and culture, serologic test, skin biopsy</td>
</tr>
<tr>
<td>Candidiasis</td>
<td><em>Candida albicans</em></td>
<td>Mucous membrane, sputum, blood, tissue, urine, CSF</td>
<td>Smear and culture</td>
</tr>
<tr>
<td>Coccidioidomycosis</td>
<td><em>Coccidioides immitis</em></td>
<td>Sputum, bone, skin, joint, CSF</td>
<td>Smear and culture, serologic skin test, biopsy</td>
</tr>
<tr>
<td>Cryptococcosis</td>
<td><em>Cryptococcus neoformans</em></td>
<td>CSF, sputum, urine</td>
<td>Smear and culture, serologic test, antigen detection</td>
</tr>
<tr>
<td>Histoplasmosis</td>
<td><em>Histoplasma capsulatum</em></td>
<td>Sputum, urine, blood, bone marrow</td>
<td>Smear and culture, serologic test, biopsy</td>
</tr>
<tr>
<td>Mucormycosis</td>
<td>Members of order Mucorales (<em>Absidia, Rhizopus, Mucor</em>)</td>
<td>Nose, pharynx, stool, CSF, sputum, ear</td>
<td>Smear and culture, serologic test, biopsy</td>
</tr>
<tr>
<td>Paracoccidioidomycosis</td>
<td><em>Paracoccidioides</em></td>
<td>Lung tissue, sputum, bone, CSF</td>
<td>Smear and culture, serologic test, biopsy</td>
</tr>
<tr>
<td>Pseudallescheriasis</td>
<td><em>Allescheria boydii</em></td>
<td>Lesions of skin, bone, brain, joint</td>
<td>Smear and culture, serologic test, biopsy</td>
</tr>
<tr>
<td>Sporotrichosis</td>
<td><em>Sporothrix schenckii</em></td>
<td>Skin lesion, CSF, bone marrow, ear</td>
<td>Smear and culture, biopsy, serologic test</td>
</tr>
<tr>
<td>Tinea pedis (athlete’s foot)</td>
<td><em>Epidermophyton</em> spp. and <em>Candida albicans</em>, <em>Trichophyton mentagrophytes</em>, <em>Trichophyton rubrum</em></td>
<td>Skin</td>
<td>Hair, skin, nail scrapings for smear and culture</td>
</tr>
<tr>
<td>Tinea capitis (ringworm of scalp)</td>
<td><em>Microsporum</em> (any spp.) and <em>Trichophyton</em> (all except <em>T. concentricum</em>)</td>
<td>Skin, hair</td>
<td>Hair, skin scrapings for smear and culture</td>
</tr>
<tr>
<td>Tinea barbae (ringworm of beard, barber’s itch)</td>
<td><em>Trichophyton</em> and <em>Microsporum</em> spp.</td>
<td>Skin, hair</td>
<td>Hair, skin scrapings for smear and culture</td>
</tr>
<tr>
<td>Tinea cruris (jock itch)</td>
<td><em>Epidermophyton</em> spp. and <em>Candida albicans</em></td>
<td>Skin</td>
<td>Hair, skin scrapings for smear and culture</td>
</tr>
<tr>
<td>Tinea corporis (ringworm of the body)</td>
<td><em>Trichophyton rubrum</em>, <em>Trichophyton tonsurans</em></td>
<td>Skin</td>
<td>Skin scrapings for smear and culture</td>
</tr>
<tr>
<td>Tinea unguum (nail)</td>
<td><em>Trichophyton rubrum</em>, <em>Trichophyton tonsurans</em>, <em>Trichophyton verrucosum</em>, <em>Epidermophyton</em> spp.</td>
<td>Nail</td>
<td>Nail culture</td>
</tr>
</tbody>
</table>
4. Pluck hair stubs out with tweezers because the fungus is usually found at the base of the hair shaft. Use of a Wood's light in a darkened room helps identify the infected hairs.
5. Samples from infected nails should be procured from beneath the nail plate to obtain softened material from the nail bed. If this is not possible, collect shavings from the deeper portions of the nail and place them in a covered sterile container.

Common Diagnostic Methods for Fungal Diseases

1. A Wood's light is used to determine presence of a fungus directly on hair. A Wood's light is a lamp that uses ultraviolet rays of 3660A. In a darkened room, infected hairs fluoresce a bright yellow-green under the Wood's light.
2. Direct microscopic examination of tissue samples placed on a slide is performed to determine whether a fungus is actually present. The potassium hydroxide (KOH) test or Calcofluor white stain test is used to detect the presence of mycelial fragments, arthrospores, spherules, or budding yeast cells and involves mixing the specimen with the reagent on a glass slide. The slide is then microscopically examined for fungal elements.
3. A fluorescent brightener, Calcofluor white, fluoresces when exposed to ultraviolet light. This reagent stains the fungi, causing them to exhibit a fluorescence that can be detected microscopically. It can be used on tissue and has the same sensitivity as KOH. Moreover, it allows for easier and faster detection of fungal elements. Calcofluor white–stained specimens can also be examined under bright field or phase-contrast microscopy.
4. Cultures are done to identify the specific type of fungus. Fungi are slow growing and are subject to overgrowth by contaminating and more rapidly growing organisms. Fungemia (fungus in the blood) is an opportunistic infection, and often a blood culture reveals the earliest suggestion of the causative organism.
5. For fungal serology tests, single titers greater than 1:32 usually indicate the presence of disease. A fourfold or greater rise in titer of samples drawn 3 weeks apart is significant. However, serologic diagnosis of Candida and Aspergillus species can be disappointing. Complement fixation tests for histoplasmosis and coccidioidomycosis can aid in the diagnosis of these diseases. The immunodiffusion test is helpful for the diagnosis of blastomycosis.

Types of specimens

1. Skin
2. Nails
3. Hair
4. Ulcer scrapings
5. Pus
6. CSF
7. Urine
8. Blood
9. Bone marrow
10. Stool
11. Bronchial washings
12. Tissue biopsy specimens
13. Prostatic secretions
14. Sputum

Diagnosis of Spirochetal Disease

Spirochetes appear as spiral and curved bacteria. The four genera of spiral and curved bacteria—Borrelia, Treponema, Leptospira, and Spirillum (Table 7.7)—include several human pathogens. Most spirochetes multiply within a living host. Pathogenic Treponema organisms are transmitted from person to person through direct contact. Borrelia pass through an arthropod vector. Leptospira are usually contracted accidentally by humans through water contaminated with animal urine or a bite by an infected animal.

Table 7.7 Spirochetal Diseases and Their Laboratory Diagnosis

<table>
<thead>
<tr>
<th>Disease</th>
<th>Causative Organism</th>
<th>Source of Specimen</th>
<th>Diagnostic Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pinta</td>
<td>Treponema carateum</td>
<td>Skin</td>
<td>Skin smear, serologic test</td>
</tr>
<tr>
<td>Rat-bite fever</td>
<td>Spirillum minor, Streptobacillus moniliformis</td>
<td>Blood, joint fluid, abscess</td>
<td>Culture serology</td>
</tr>
<tr>
<td>Relapsing fever</td>
<td>Borrelia recurrentis</td>
<td>Blood</td>
<td>Blood smear</td>
</tr>
<tr>
<td>Syphilis</td>
<td>Borrelia pallidum</td>
<td>Skin lesion</td>
<td>Skin smear, nonspecific treponemal (VDRL, RPR) and specific treponemal (FTA-ABS) serologic tests</td>
</tr>
<tr>
<td>Weil's disease</td>
<td>Leptospira interrogans</td>
<td>Urine, blood, CSF</td>
<td>Culture serologic test</td>
</tr>
<tr>
<td>Yaws</td>
<td>Treponema pertenue</td>
<td>Skin</td>
<td>Culture, serologic test</td>
</tr>
<tr>
<td>Lyme disease</td>
<td>Borrelia burgdorferi</td>
<td>Skin lesion, blood, CSF</td>
<td>Serologic test</td>
</tr>
<tr>
<td>Nonvenereal syphilis</td>
<td>Treponema endemic</td>
<td>Skin, blood</td>
<td>Serologic test</td>
</tr>
</tbody>
</table>

Clinical Considerations

1. *Borrelia* appear in the blood at the onset of relapsing fever. Louse-borne relapsing fever is caused by *Borrelia*
recurrentis, tick-borne relapsing fever by several other *Borrelia* species, and Lyme disease by *Borrelia burgdorferi*.

2. *Treponema* (*Borrelia* vincentii) is the species responsible for ulcerative gingivitis (trench mouth).
   a. *Treponema pallidum* is the species responsible for venereal syphilis in humans.
   b. *Treponema pallidum* subsp. *pertenue* is the causative agent of yaws (an infectious nonvenereal disease).
   c. *Treponema carateum* causes pinta (carate).
   d. *Treponema pallidum* subsp. *endemicum* is the cause of endemic nonvenereal syphilis (bejel).

3. *Leptospira* is the genus of microorganism responsible for Weil's disease (infectious jaundice), swamp fever, swineherd's disease, and canicola fever.
   a. The organism is widely distributed in the infected person and appears in the blood early in the disease process.
   b. After 10 to 14 days, the organisms appear in considerable numbers in the urine.
   c. Patients with Weil's disease show striking antibody responses; serologic testing is useful for diagnosis of this disease.

4. *Streptobacillus moniliformis* and *Spirillum minor* are the species responsible for rat-bite fever. Although this condition occurs worldwide and is common in Japan and Asia, it is uncommon in North and South America and most European countries. Cases in the United States have been linked to bites by laboratory rats.

### Diagnosis of Viral and Mycoplasmal Disease

Viral diseases are the most common of all human infections. Once thought to be confined to the childhood years, viral infections in adults have increasingly been recognized as the cause of significant morbidity and mortality. They also affect immunosuppressed and elderly patients (*Chart 7.1*). Viruses are responsible for such infectious diseases as hepatitis, AIDS, and other sexually transmitted diseases (STDs).

#### Chart 7.1 Viral Infections in Infants, Children, and Adults

<table>
<thead>
<tr>
<th>Disease or Syndrome</th>
<th>Suspected Viral Agents</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>INFANTS AND CHILDREN</strong></td>
<td></td>
</tr>
<tr>
<td>Upper respiratory tract infection</td>
<td>Rhinovirus, coronavirus, parainfluenza, adenovirus, respiratory syncytial virus, influenza</td>
</tr>
<tr>
<td>Pharyngitis</td>
<td>Adenovirus, coxsackie A, herpes simplex, Epstein-Barr, rhinovirus, parainfluenza, influenza</td>
</tr>
<tr>
<td>Croup</td>
<td>Parainfluenza, respiratory syncytial</td>
</tr>
<tr>
<td>Bronchitis</td>
<td>Parainfluenza, respiratory syncytial</td>
</tr>
<tr>
<td>Bronchiolitis</td>
<td>Respiratory syncytial, parainfluenza</td>
</tr>
<tr>
<td>Pneumonia</td>
<td>Respiratory syncytial, adenovirus, influenza, parainfluenza</td>
</tr>
<tr>
<td>Gastroenteritis</td>
<td>Rotavirus, adenoviruses 40–41, calicivirus, astrovirus, Norwalk-like</td>
</tr>
<tr>
<td><strong>ADULTS</strong></td>
<td></td>
</tr>
<tr>
<td>Upper respiratory tract infection</td>
<td>Rhinovirus, coronavirus, adenovirus, influenza, parainfluenza</td>
</tr>
<tr>
<td>Pneumonia</td>
<td>Coxsackie B</td>
</tr>
<tr>
<td>Gastroenteritis</td>
<td>Norwalk-like virus</td>
</tr>
<tr>
<td><strong>ALL PERSONS</strong></td>
<td></td>
</tr>
<tr>
<td>Parotitis</td>
<td>Mumps, parainfluenza</td>
</tr>
<tr>
<td>Myocarditis/pericarditis</td>
<td>Coxsackie and echoviruses</td>
</tr>
<tr>
<td>Keratitis/conjunctivitis</td>
<td>Herpes simplex, varicella-zoster, adenovirus</td>
</tr>
<tr>
<td>Pleurodynia</td>
<td>Coxsackie B</td>
</tr>
<tr>
<td>Herpingania</td>
<td>Coxsackie A</td>
</tr>
<tr>
<td>Febrile illness with rash</td>
<td>Echo and coxsackie viruses</td>
</tr>
<tr>
<td>Infectious mononucleosis</td>
<td>Epstein-Barr, cytomegalovirus</td>
</tr>
<tr>
<td>Meningitis</td>
<td>Echo and coxsackie viruses, lymphocytic choriomeningitis, herpes simplex virus</td>
</tr>
<tr>
<td>Encephalitis</td>
<td>Herpes simplex, togaviruses, bunyaviruses, flaviviruses, rabies, enteroviruses, measles, human immunodeficiency virus (HIV), JC virus</td>
</tr>
<tr>
<td>Hepatitis</td>
<td>Hepatitis A, B, C, non-A, non-B; delta agent; E</td>
</tr>
<tr>
<td>Hemorrhagic cystitis</td>
<td>Adenovirus, BK virus</td>
</tr>
<tr>
<td>Cutaneous infection with rash</td>
<td>Herpes simplex, varicella-zoster, enteroviruses, Epstein-Barr, measles, rubella, parovirus, human herpes virus 6</td>
</tr>
<tr>
<td>Hemorrhagic fever</td>
<td>Ebola, Marburg, Lassa, hantavirus</td>
</tr>
<tr>
<td>Acute respiratory failure</td>
<td>Hantavirus</td>
</tr>
</tbody>
</table>

Viruses are submicroscopic, filterable, infectious organisms that exist as intracellular parasites. They are divided into two groups according to the type of nucleic acid they contain: RNA or DNA.

The *mycoplasmas* are scotobacteria without cell walls that are surrounded by a single triple-layered membrane; they are also known as *pleuropneumonia-like* organisms (PPLOs). Physiologically, mycoplasmal diseases are considered to be intermediate between those caused by bacteria and those caused by rickettsiae. One species, *Mycoplasma pneumoniae*, is recognized as the causative agent of primary atypical pneumonia and bronchitis. Other species are suspected as possible causal agents for urethritis, infertility, early-term spontaneous abortion, rheumatoid arthritis, myringitis, and
Viruses and mycoplasmas are infectious agents small enough to pass through bacteria-retaining filters. Although small size is the only property they have in common, viruses and mycoplasmas cause illnesses that are often indistinguishable from each other in terms of clinical signs and symptoms; in addition, both frequently occur together as dual infections. Therefore, the serologic (antigen-antibody) procedures commonly used for diagnosing viral disease are also used for diagnosing mycoplasmal infections (Table 7.8).

**Table 7.8 Viral Infections and Their Laboratory Diagnosis**

<table>
<thead>
<tr>
<th>Infection Type and Virus Information</th>
<th>Throat</th>
<th>Stool/Rectal</th>
<th>CSF Swab</th>
<th>Urine Vesicle Fluid/Swab</th>
<th>Conjunctival Swab/Scraping</th>
<th>Other</th>
<th>Blood Serology</th>
<th>Additional Information</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RESPIRATORY</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adenovirus</td>
<td>X</td>
<td></td>
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<td></td>
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<tr>
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<td>Herpes simplex virus (HSV)</td>
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<td>Nasopharyngeal Yes swab</td>
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<tr>
<td>Influenza virus</td>
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<td>Mumps virus</td>
<td>X</td>
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<td>X</td>
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<td>Yes</td>
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<td>Parainfluenza virus</td>
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<td></td>
<td>Nasopharyngeal Yes swab</td>
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<td>Respiratory syncytial virus (RSV)</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td>Nasopharyngeal Yes aspirate or swab</td>
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<td>Rhinovirus</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td>Nasal</td>
<td>Nasal specimen preferred</td>
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<td><strong>RASH Maculopapular</strong></td>
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<td>Enterovirus</td>
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<td>No *</td>
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<tr>
<td>Rubella virus</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>Viral culture rarely done</td>
<td>Yes</td>
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<td>Measles (rubeola)</td>
<td>X</td>
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<td>Viral culture rarely done</td>
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<td>Yes</td>
<td>Special culture required</td>
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<td><em>Vesicular</em></td>
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<td>Special culture required</td>
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<td>Coxsackie virus</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td>No *</td>
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<tr>
<td>HSV</td>
<td></td>
<td></td>
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<td></td>
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<td>Many strains of type A coxsackie virus do not grow in tissue culture</td>
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<td>Varicella-zoster virus (VZV)</td>
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<td></td>
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<td>Yes</td>
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<td>Vaccinia and other poxviruses</td>
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<td><strong>Central nervous system (aseptic meningitis, encephalitis)</strong></td>
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<tr>
<td>Arbovirus</td>
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<tr>
<td>Enterovirus</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td>Yes</td>
<td>Blood PCR is useful</td>
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<tr>
<td>HSV</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td>Brain biopsy Yes</td>
<td></td>
<td>Recovery of HSV type 1 from CSF is rare except in neonates</td>
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<tr>
<td>Mumps virus</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td>Yes</td>
<td>Blood</td>
</tr>
<tr>
<td>Rabies virus</td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>Yes</td>
<td>Skin biopsy of neck for fluorescent assay</td>
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<td><strong>CONGENITAL AND PERINATAL</strong></td>
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<td>Cytomegalovirus (CMV)</td>
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<td>Blood (leukocytes)</td>
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<td>Yes</td>
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<td>Virus</td>
<td>Site</td>
<td>Test Method</td>
<td>Presence</td>
<td></td>
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<tr>
<td>Enterovirus</td>
<td>Blood</td>
<td>(PCR)</td>
<td>No *</td>
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<tr>
<td>HSV</td>
<td>Blood</td>
<td>(PCR)</td>
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<tr>
<td>Adenovirus</td>
<td>GASTROINTESTINAL</td>
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<td>Parvovirus (Norwalk-like agents)</td>
<td>GASTROINTESTINAL</td>
<td>X</td>
<td>Research laboratories only</td>
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<tr>
<td>Adenovirus</td>
<td>EYE</td>
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<tr>
<td>Enterovirus</td>
<td>EYE</td>
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<td>No *</td>
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<tr>
<td>HSV</td>
<td>HEART</td>
<td>X</td>
<td>Yes</td>
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<tr>
<td>Coxsackie virus B</td>
<td>HEART</td>
<td>X</td>
<td>No *</td>
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<tr>
<td>CMV</td>
<td>HEART</td>
<td>X</td>
<td>Yes</td>
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<tr>
<td>Influenza A, V</td>
<td>HEART</td>
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<tr>
<td>Infectious mononucleosis</td>
<td>HEART</td>
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<tr>
<td>Epstein-Barr virus (EBV)</td>
<td>HEART</td>
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<td>Yes</td>
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<td>Immunodeficient patient CMV</td>
<td>HEART</td>
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<tr>
<td>CMV</td>
<td>HEPATITIS</td>
<td>X</td>
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<tr>
<td>EBV</td>
<td>HEPATITIS</td>
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<td>Yes</td>
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<tr>
<td>HEPATITIS A, B, AND C</td>
<td>HEPATITIS</td>
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<tr>
<td>HSV</td>
<td>URINARY</td>
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<td>Yes</td>
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</table>

*Enterovirus serology is not routinely available but can be performed with selected antigens under special circumstances.

**Approach to Diagnosis**

1. Isolation of the virus in tissue culture remains the gold standard for detection of many common viruses. Diagnostic modalities include the following:
   a. Tissue culture
   b. Direct detection in specimens
   c. Identification through specific cytopathic effect
   d. Use of immunofluorescence and immunoperoxidase, latex agglutination, or enzyme-linked immunosorbent
assay (ELISA) to identify
f. Direct nucleic acid probe and PCR technology

2. Serologic studies for antigen-antibody detection are valuable in regard to viral disease. Epstein-Barr virus (EBV) and human hepatitis viruses are routinely serodiagnosed. Classically, a fourfold rise in antibody titer is used to identify a particular infectious agent, provided that the pathogenesis of the agent agrees with the symptoms of the infected patient. An acute-phase serum is collected within the first several days after symptom onset. A convalescent-phase serum is collected 2 to 4 weeks later. A fourfold difference in antibody titer between the two sera is statistically significant. Alternatively, detection of specific immunoglobulin M (IgM) suggests acute infection. IgG antibody without IgM suggests infection sometime in the past.

3. Available cell cultures vary greatly in their sensitivity to different viruses. One cell type or species may be more sensitive than another for detecting the virus in low titers. For example, human embryonic kidney (HEK) or monkey kidney (MK) can be used for adenovirus, enterovirus, herpes simplex, measles, influenza, parainfluenza, and rubella; however, HEK cannot be used for cytomegalovirus (CMV) or influenza.

4. The critical first step in successful viral diagnosis is the timely and proper collection of specimens. The choice of which specimen to collect depends on typical signs and symptoms and the suspected virus. Improper specimen choice and collection is one of the biggest factors in diagnostic delays.

Specimen Collection

1. Collect specimens for viruses as early as possible during the course of the illness, preferably within the first 4 days after symptom onset. If specimen collection is delayed for 7 or more days after symptoms appear, diagnosis will be compromised. Virus titers are highest in the early part of the illness, when the host has not yet mounted a robust immune response. Little neutralizing antibody is present. Detection of a virus by culture, direct detection, or serology is greatly enhanced when the virus titers are high.

2. Sampling procedure
   a. For localized infection:
      1. Direct sampling of affected site (eg, throat swab, skin scraping)
      2. Indirect sampling. For example, if CSF is the target sample in a central nervous system infection, the indirect approach would involve obtaining throat or rectal swabs for culture.
   b. Sampling from more than one site, for example, in disseminated disease or with nonspecific clinical findings
   c. The type of applicator used to obtain specimens may affect accuracy of results. Do not use wooden applicators or cotton swabs because they are toxic to viruses. A self-contained transport system is recommended to ensure that the specimen remains moist.

3. When transporting specimens:
   a. Keep in mind that viral specimens are unstable and rapidly lose infectivity outside of living cells. Prompt delivery to the laboratory is essential. Samples must be refrigerated or placed on ice or cold packs while in transit.
   b. Freezing and thawing of specimens diminishes the quantity of available viable virus.

4. Accurate patient information must accompany the specimen to the laboratory. In addition to the required patient identification information, the requisition should include:
   a. Pertinent information that would influence processing of the specimen (eg, patient is immunocompromised owing to renal transplantation)
   b. Exact nature of the specimen
   c. Patient demographics
   d. Contact person or clinic so as to expedite the notification of positive results

5. Specimens of small volume (eg, vesicular fluid, fine-needle aspiration, biopsy samples) should be transported in a liquid medium. Suggested viral transport media are Hank’s balanced salt solution, 0.2 mol/L sucrose-phosphate, and bacteriologic broth (tryptic soy or veal infusion).

Often, a complete microbiologic workup of a specimen (tissue, bronchoscopy) is requested along with a viral workup. Because viral transport media contain antibiotics, sterile saline is recommended. Personnel in the laboratory can then divide the specimen for workup within the microbiology subsections. Specimens of a liquid nature (urine, CSF, sputum, body fluids) are collected in a sterile container. For patients with suspected viremia, a viral culture of theuffy coat of peripheral blood is submitted. Blood specimens are collected in evacuated tubes containing heparin or ethylene diamine tetraacetic acid (EDTA).

Clinical Considerations

1. Herpes simplex is the virus most frequently isolated and diagnosed virus in the laboratory.

2. Viral culture results are normally available within 3 to 5 days, although rapid test results (24 hours) are accurate and available for certain viruses, such as CMV.

3. Significance of viral cultures
   a. Positive viral culture results from the following sources are **diagnostically accurate**:
      1. Autopsy specimens
      2. Blood (leukocyte buffy coat)
      3. Biopsy
      4. CSF
      5. Other body fluids
      6. Cervix
      7. Eye
      8. Skin lesions
      9. Fine-needle aspirates
      10. Bronchial alveolar wash brushing
   b. **Probably diagnostically accurate are**:
      1. Throat
      2. Urine
      3. Sputum
4. Genital (cervical, penile)
5. Nasal aspirates or washes
6. Vesicular
7. Skin (mouth, lip)

c. Possibly diagnostically accurate is stool or rectal swab
d. Viruses do not compromise normal flora in the body. However, bacterial or fungal contamination of specimens can occur.

**Diagnosis of West Nile Virus, West Nile Fever, and West Nile Encephalitis**

The West Nile virus (WNV) is a single-stranded RNA flavivirus first isolated in Uganda about 1937. The virus first appeared in the United States in 1999 and, to date, has been documented in almost every state. The routes of transmission include the common household mosquito, which serves as the vector, and birds (eg, crows, sparrows, and jays), which are the reservoir hosts. The infected mosquito can then carry the virus particles in its salivary glands and infect susceptible bird species as well as humans. There is no evidence that the virus can be spread by human-to-human transmission or by handling infected birds. Symptoms of WNV include fever, headache, neck stiffness, and skin rash. West Nile fever is the mild form of the disease, characterized by flu-like symptoms lasting only a few days with no long-term effects. West Nile encephalitis is the more severe form of the disease, characterized by encephalitis and/or meningitis, which can lead to stupor, disorientation, coma, convulsions, and occasionally death.

This test is used to measure antibodies (IgM) produced early in the infected person.

**Reference Values**

**Normal** Negative for the West Nile virus IgM antibody

**Procedure** Collect either a blood or CSF sample. Not all laboratories are equipped to measure the antibody, and the sample may have to be forwarded to a commercial or public health laboratory.

**Clinical Alert**

The blood test may be negative early in the course of the infection; however, within 8 days of the onset of symptoms, 90% of infected people will become positive. The Centers for Disease Control and Prevention may perform a plague reduction neutralization test (PRNT) on a specimen for confirmation.

**Interfering Factors** Exposure to the St. Louis encephalitis virus may result in a false-positive test result for WNV.

**Clinical Alert**

Currently, there is no vaccination against WNV. Treatment is aimed at prevention of secondary infections (eg, pneumonia and urinary tract infection), airway management, and good nursing care.

**Interventions**

**Pretest Patient Care**

1. Explain purpose and procedure.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Care**

1. Interpret test results, monitor and counsel patient appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Diagnosis of Sexually Transmitted Disease**

STDs present a serious and increasing public health problem. They are caused by a variety of etiologic agents (Table 7.9). Some conditions, such as chlamydial and nongonococcal urethritis, have reached epidemic proportions. Although nongonococcal urethritis is a nonreportable disease in the United States, it is estimated that more than two million new cases occur each year. Manifestations of these infections range from the carrier state (asymptomatic) to diseases with obvious symptoms such as cervicitis, conjunctivitis, endometritis, epididymitis, infertility, pharyngitis, proctitis, lymphogranuloma venereum, salpingitis, trachoma, urethritis, and in the neonate, conjunctivitis and pneumonia.

**Table 7.9 Sexually Transmitted Diseases and Their Laboratory Diagnoses**

<table>
<thead>
<tr>
<th>Disease</th>
<th>Causative Agents</th>
<th>Diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chancroid</td>
<td>Haemophilus ducreyi</td>
<td>Culture of lesion or aspirate. Differential diagnosis should include syphilis, herpes, and lymphadenopathy associated virus (LAV) monoclonal antibody test</td>
</tr>
<tr>
<td>Gonorrhea</td>
<td>Neisseria gonorrhoeae</td>
<td>Gram stain of male urethra, culture or PCR of male urethra or female cervix, rectum, or pharynx. DNA probe, PCR of urine</td>
</tr>
<tr>
<td>Granuloma inguinale (donovanosis)</td>
<td>Calymmatobacterium granulomatis (formerly Donovania granulomatis)</td>
<td>Wright-Giemsa stain of lesion, tissue biopsy</td>
</tr>
<tr>
<td>Hepatitis B</td>
<td>Hepatitis B virus (HBV)</td>
<td>Serologic testing, hepatitis B antigen and antibody</td>
</tr>
<tr>
<td>Condition</td>
<td>Microbes/Agents</td>
<td>Diagnostic Methods</td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>---------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Genital herpes</td>
<td>Herpes simplex virus (HSV) types 1 and 2</td>
<td>Viral culture from unroofed blister, scrapings examined by fluorescent microscopy or cytologic stains</td>
</tr>
<tr>
<td>Lymphogranuloma venereum (LGV)</td>
<td>Chlamydia trachomatis serotypes L₁, L₂, and L₃</td>
<td>Culture of aspirate of bubo, serologic test</td>
</tr>
<tr>
<td>Molluscum contagiosum</td>
<td>Molluscum contagiosum virus</td>
<td>Clinical appearance of lesions (pearly white, painless, umbilicated papules), microscopic examination of scrapings</td>
</tr>
<tr>
<td>Chlamydia</td>
<td>Chlamydia trachomatis serotypes D–K</td>
<td>Cell culture, urogenital swabs for direct antigen test, PCR</td>
</tr>
<tr>
<td>Candidosis (monilia)</td>
<td>Candida albicans</td>
<td>Culture, potassium hydroxide (KOH) wet mount, Gram stain, DNA probe</td>
</tr>
<tr>
<td>Pelvic inflammatory disease (PID)</td>
<td>Neisseria gonorrhoeae, Chlamydia trachomatis</td>
<td>Clinical symptoms, cervical culture, DNA probe, PCR, laparoscopy or culdocentesis</td>
</tr>
<tr>
<td>Pediculosis pubis</td>
<td>Phthirus pubis (public or crab louse)</td>
<td>Adult lice or nits appear on body hairs</td>
</tr>
<tr>
<td>Scabies</td>
<td>Sarcoptes scabiei</td>
<td>Characteristic lesions, scrapings for microscopy</td>
</tr>
<tr>
<td>Syphilis</td>
<td>Treponema pallidum</td>
<td>Darkfield microscopy, serologic test</td>
</tr>
<tr>
<td>Trichomoniasis</td>
<td>Trichomonas vaginalis</td>
<td>Vaginal, urethral, prostatic secretion examined microscopically, culture; PCR</td>
</tr>
<tr>
<td>Nonspecific urethritis (NGU)</td>
<td>Chlamydia trachomatis (50% of cases), Ureaplasma urealyticum, Trichomonas vaginalis, Candida albicans, herpes simplex virus</td>
<td>Identification by smear, culture, or molecular tests of specific etiologic agent</td>
</tr>
<tr>
<td>Nonspecific vaginitis</td>
<td>Gardnerella vaginalis, Mobiluncus cortisii, Mobiluncus mulieris</td>
<td>Wet mount for “clue” cells or Pap smear; fishy smell is released when specimen fluid is mixed with 10% KOH. Culture; DNA probe. Culture or enzyme immunoassay to rule out gonorrhea</td>
</tr>
<tr>
<td>Condylomata acuminata (venereal warts)</td>
<td>Human papilloma DNA virus</td>
<td>Typical clinical lesion: cauliflower-like, soft, pink growth around vulva, anus, labia, vagina, glans penis, urethra, and perineum; rule out syphilis</td>
</tr>
<tr>
<td>Acquired immunodeficiency syndrome (AIDS)</td>
<td>Human immunodeficiency virus (HIV)</td>
<td>Serologic tests</td>
</tr>
<tr>
<td>Gastrointestinal (giardiasis, amebiasis, shigellosis campylobacteriosis, and anorectal infections)</td>
<td>Enteric infections: Giardia lamblia, Entamoeba histolytica, and Cryptosporidium spp.</td>
<td>Ova and parasite examination</td>
</tr>
<tr>
<td></td>
<td>Shigella spp., Campylobacter fetus, Strongyloides spp. (worms)</td>
<td>Stool culture</td>
</tr>
<tr>
<td></td>
<td>Anorectal: Neisseria gonorrhoeae, Chlamydia trachomatis, Treponema pallidum, Herpes simplex virus, Human papilloma virus</td>
<td>Stool culture, Anal swab specimen, culture, DNA probe, Anal swab culture, DNA probe, Tissue culture, Signs and symptoms, DNA probe</td>
</tr>
</tbody>
</table>

*The pathogens causing sexually transmitted diseases span the full range of medical microbiology; their only common characteristic is that they may cause genital disease or be transmitted by genital contact.

### Suggested Specimens

1. Urethral, vaginal, cervical swabs
2. Semen
3. Urine
4. Prostatic secretion
5. Tissue biopsy
6. Swabs of oral lesions
7. Blood for serologic tests

### Common Diagnostic Methods

1. Viral isolation in tissue cell cultures
2. Specific serologic antibody assays and syphilis detection tests
3. Cytologic techniques, such as Papanicolaou (Pap) and Tzanck smears to demonstrate giant cells associated with herpes virus infection
4. Gram stain and bacterial culture; saline wet prep
5. ELISA and immunoperoxidase assay to detect etiologic agent
6. Fluorescein or enzyme-tagged monoclonal antibodies to detect and identify etiologic agents
7. DNA probe
8. PCR

Clinical Considerations

1. Patients presenting with one STD are frequently infected with other types of sexually transmitted pathogens.
2. Asymptomatic carriers are common.
3. The disease may recur if the patient is reinfeected by the nontreated sexual partner.
4. Genital tract infections caused by sexually transmitted organisms in children are often the result of sexual abuse.
5. Cultures should always be obtained, especially for Chlamydia, when required as legal evidence.
6. For suspected herpetic lesions, the virus is best recovered from the base of an active lesion. The older the lesion, the less likely it is to yield viable virus. Open the vesicle with a small-gauge needle or Dacron swab. Rub the base of the lesion vigorously to recover infected cells onto the swab, and place the swab in a viral transport medium. If large vesicles are present, aspirate material directly by needle and syringe. A separate swab can be collected for a Tzanck preparation (histology stain).
7. For darkfield examination (eg, syphilis), cleanse the area around the lesion with sterile saline. Abrade the surface with sterile dry gauze until blood is expressed. Continue to blot until blood ceases; squeeze the area until serous fluid is expressed. Touch the material to a clean glass slide, add a cover slip, and examine the specimen immediately for motile spirochetes.
8. Complications of untreated STDs include ectopic (tubal) pregnancy, infertility, chronic pelvic pain, and poor pregnancy outcomes.

Diagnosis of Food Poisoning

Most cases of food poisoning in the United States are associated with Bacillus cereus, a species found in soil, water, airborne dust, vegetation, cereals, pasteurized food, and powdered milk. GI anthrax due to ingestion of contaminated meat has been reported in developing nations, not in the United States. (Spices in meat have been reported to be contaminated with the B. cereus spores.) B. cereus produces two toxins, an emetic toxin that causes vomiting and an enterotoxin that causes diarrhea. Unrefrigerated fried rice has been associated with the emetic toxin, whereas poultry, cooked meats, mashed potatoes, soups, and desserts have been associated with the enterotoxin.

This test is used to detect one of the two toxins produced by B. cereus.

Reference Values

Normal Negative for culture of B. cereus colonies by DNA probe or other microbiologic tests

Procedure Collect stool specimens (25–50 g) for culture. Call your laboratory about special stool culture collection.

Clinical Alert

● Suspected food specimens may also be tested.
● Refrigerate specimen in clean, sealed, leak-proof containers.
● If a delay of more than 2 hours is anticipated, the specimen should be placed in a Cary-Blair transport medium.

Clinical Implications Positive abnormal findings of the special characteristics of B. cereus are consistent with food poisoning.

Interventions


Posttest Patient Care Interpret test outcomes, monitor prescribed drug treatment (vancomycin, erythromycin).

BIOTERRORISM: INFECTIOUS AGENTS

Diagnosis of Botulism Infection

Human botulism is caused by the spore-forming, obligate anaerobe Clostridium botulinum. C. botulinum produces the botulinum toxin, which is the most poisonous substance known (1 g evenly dispersed would kill more than 1 million people). There are seven distinct antigenic types of the botulinum toxin designated types A through G. All forms of botulism are the result of absorption from a mucosal surface (eg, GI tract or lung) or a wound into the circulatory system. C. botulinum can be found in the soil and in undercooked food that is not kept hot. Cases of waterborne botulism have not been documented, although aerosolization of the toxin and subsequent inhalation has been done experimentally. Foil-wrapped baked potatoes held at room temperature after baking can cause botulism, as can contaminated condiments, such as sautéed onions or cheese sauce.

This test is used to confirm the presence of C. botulinum, which produces the botulinum toxin.

Reference Values

Normal Absence of botulinum toxin Absence of incremental response to repetitive nerve stimulation on an electromyogram

Procedure

1. Obtain specimens from blood, stool, gastric aspirates or vomitus, and, if available, suspected food.
2. Obtain at least 30 mL of venous blood in a red-topped Vacutainer.
3. Use an enema (with sterile water) to obtain an adequate fecal sample if the patient is constipated.
4. Refrigerate all samples.
5. Use the "mouse" bioassay to determine whether there is any botulinum toxin present.
Clinical Alert
1. In some cases, an electromyogram is performed to differentiate cause of acute flaccid paralysis.
2. Antibiotics can be used to treat secondary infections; however, they do not have a direct effect on the botulinum toxin.
3. Postexposure prophylaxis is limited owing to antitoxin scarcity and reactogenicity.
4. By law, in most areas of the country, suspected botulism must be reported to local public health authorities.

Clinical Implications
1. The identification of botulism neurotoxin is evidence of botulism poisoning.
2. Tell family that botulism paralyzes muscles and patients die quickly because they cannot breathe.

Interventions

Pretest Patient Care
1. Explain the purpose, procedure, and risks of obtaining a specimen.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Care
1. Do not isolate patients diagnosed with botulism because it is not contagious and cannot be transmitted from person to person.
2. Interpret test results, provide supportive care, and monitor appropriately.
3. Administer an antitoxin (ie, equine antitoxin) in a timely manner to minimize subsequent nerve damage.
4. The patient should be monitored for impending respiratory failure.
5. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

NOTE
In the United States, the botulinum antitoxin is available from the Centers for Disease Control and Prevention.

NOTE
The need for mechanical ventilation has varied from 20% in adults to 60% in children.

Clinical Alert
1. Laboratory personnel should observe universal standard precautions.
2. Sterile water, not saline, should be used for the enema solution because saline will interfere with the bioassay.
3. The use of anticholinesterases (eg, physostigmine salicylate or pralidoxime chloride) by the patient can interfere with the bioassay.
4. Decontaminate surfaces with 0.1% hypochlorite bleach solution.
5. Contaminated clothing should be washed with soap and water.

NOTE
About 20 state laboratories, including the Centers for Disease Control and Prevention, can test for botulism.

Diagnosis of Anthrax Infection

Anthrax is a communicable disease transmitted from animals to humans. Humans can contact Bacillus anthracis (spore-forming, gram-positive bacillus) from handling or consuming undercooked meat from infected animals. The organism can also be inhaled from animal products (eg, wool) or during intentional release of spores (ie, bioterrorism). There are three forms of anthrax: cutaneous, GI, and inhalational.

Clinical Alert
1. As soon as anthrax is suspected, notify the state public health laboratory and Centers for Disease Control and Prevention.
2. Take precautions to avoid production of aerosols of infected material.
3. If GI anthrax is suspected, collect samples of gastric aspirate, feces, or food along with three blood cultures.
4. Household bleach solutions (5.25% hypochlorite) diluted 1:10 can be used to decontaminate surfaces. Contaminated instruments should be autoclaved after immersion in decontamination solution.
5. Proper immunization is required for persons who work directly with contaminated animal hides or animal tissues or spores.
6. Skin infections constitute 95% of anthrax infections, with a 20% death rate in untreated skin (cutaneous) anthrax.
7. All cases of anthrax infection due to inhaled spores are fatal.
8. Abnormal chest x-ray findings show widening of the mediastinum due to hemorrhage.
9. Person-to-person transmission of anthrax infection has not been observed.

Reference Values
Normal  Negative for the *B. anthracis* organism (appears as two to four cells, encapsulated)

**Procedure**

1. Use sputum, blood, or stool specimens to isolate *B. anthracis*.
2. Perform procedure in a Biological Safety Level 2 (BSL 2) microbiologic laboratory.
3. Analyze samples in a certified Class II biologic safety cabinet (BSC).
4. Subculture a routine sputum, blood, or stool sample to sheep blood agar (SBA), MacConkey agar, or phenyl ethyl alcohol (PEA) plates.
5. Incubate cultures at 35°–37°C and examine within 18–24 hours of incubation.
6. Test isolates for motility, morphology, ß-hemolysis, and Gram stain to differentiate colonies of *B. anthracis* from other bacilli.
7. Remember that *B. anthracis* is an encapsulated gram-positive rod, with oval-shaped, nonswelling spores, and ground-glass appearance of colonies, and is nonmotile and nonhemolytic.
8. Soak two dry sterile swabs in vesicular fluid (previously unopened vesicle) for cutaneous anthrax.
9. Use a stool specimen for GI anthrax.
10. Use a sputum specimen for inhalation anthrax; in the later stages, use a blood sample.

**Procedural Alert**

*B. anthracis* grows well in SBA plates but does not grow on MacConkey agar.

**Clinical Implications** The isolation of *B. anthracis* rods confirms the diagnosis of anthrax.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose, procedure, and risks of obtaining a specimen.
2. Obtain current history of occupations, signs, and symptoms. Anthrax, usually fatal, is accompanied by fever, dyspnea, coughing, chest pain, heavy perspiration, and bluish skin due to lack of oxygen.
3. See guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Care**

1. Contact the FBI and state public health department if *B. anthracis* is identified.
2. Interpret test outcomes and monitor treatment, eg, penicillin or tetracycline. Report signs and symptoms.
3. See guidelines in Chapter 1 for safe, effective, informed posttest care.

**Diagnosis of Hemorrhagic Fever, Hantaan Virus, Ebola Virus, and Yellow Fever Infections**

This test is done to diagnose hemorrhagic fever (HF; with renal symptoms, HFRS), an endemic threat in the United States, and yellow fever associated with hepatitis. Signs and symptoms include fever, thrombocytopenia, renal failure, shock, multiorgan failure, and lung edema. Jaundice occurs in yellow fever. Heavy rains are associated with an increase in number of rodents (which are the vectors in HF) and mosquitoes in yellow fever. The hantavirus, called *sin nombre* (no name), is responsible for hantavirus pulmonary syndrome (HPS). See microbiology references for more information.

**Reference Values**

*Normal* No evidence of Hantaan virus, Ebola virus, or 17 other viruses that may cause HF in rodents or deer mice No evidence of yellow fever virus that may cause hepatitis. Negative for presence of hantaviral antigens in humans

**Procedure**

1. Obtain specimen of blood, sputum, tissue, and possibly urine using standard precautions.
2. Place specimens in biohazard bags. All specimens are considered infectious.

**Clinical Alert**

1. Follow airborne precautions with negative-pressure rooms if available.
2. Use personal protective equipment.

**Clinical Implications** Growth of Hantaan virus (or any of the other 17 causative viruses) in culture or presence of hantaviral antigens is evidence of disease. Thrombocytopenia is present in blood samples.

**Clinical Alert**

Evidence of HF is reported to local, state, and federal authorities.

**Interventions**

**Pretest Patient Care**

1. Explain necessity, purpose, and procedure of testing. Knowledge of signs and symptoms, eg, pneumonia, fever, muscular pain, sometimes jaundice, hemorrhage from nose or GI tract, facial swelling, is needed. Questions regarding occupation, living accommodations, and circumstances (eg, recent heavy rains, mosquitoes, tropical climate, port city) are important.
2. Be aware that no person-to-person transmission has been described, but close contact is usual.
3. See guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Care**
1. Counsel, monitor, and treat appropriately. Report signs and symptoms. Patients are usually very ill. If the HF viruses are isolated, or the Hantaan antigens (or other causative viruses) are identified, supportive therapy and antibiotics are used to treat secondary bacterial infections. Correct dehydration and treat acidosis and blood cell abnormalities.

2. Immunity to yellow fever occurs after recovery.

3. See guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

Monitor close and high-risk contact for 21 days.

Diagnosis of Plague, Bubonic Plague, and Primary Septicemic Plague Infection

This test is done to diagnose plague. Plague (zoonotic infection of rats, squirrels, prairie dogs, and other rodents) can be found on every continent except Australia. Plague is transmitted by the bite of an infected flea. Signs and symptoms include a sudden onset of fever, chills, generalized weakness, and bubo (swollen tender lymph nodes). Bubo typically develops in the axilla, cervical, or groin regions.

Reference Values

Normal: Negative for the presence of Yersinia pestis

Procedure

1. Obtain specimens of blood, sputum, or a lymph node aspirate following standard precautions.

2. Transport specimens per laboratory protocol.

Clinical Alert

Specimens should be processed in biosafety level 2 practices or biosafety level 3 if there is a high potential for aerosolization (eg, centrifugation procedures). There are no rapid available diagnostic tests for plague. Cultures take about 24 to 48 hours.

Clinical Implications

A positive test for Y. pestis is evidence of the disease, and treatment (eg, streptomycin, tetracycline, doxycycline) should begin immediately. Because person-to-person transmission has not been identified, isolation is not necessary.

Clinical Alert

1. Evidence of plague must be reported to the local, state, and federal health departments.

2. Persons having close contact (<2 meters) with an infected individual should receive postexposure antibiotic prophylaxis for 7 days.

Interventions

Pretest Patient Care

1. Explain necessity, purpose, and procedure of testing. Obtain history of pertinent signs and symptoms.

2. There is limited evidence of person-to-person spread, thus isolation is not necessary.

3. See guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Care

1. Counsel, monitor, and treat appropriately. Evidence does not support that residual Y. pestis poses an environmental threat, and the organism does not survive long outside the host.

2. See guidelines in Chapter 1 for safe, effective, informed posttest care.

Diagnosis of Smallpox Infection

Smallpox, a DNA virus, was once worldwide in scope; however, vaccination has all but eliminated the disease. Recently, concerns of its use as a “biological weapon” has prompted medical and public health care professionals to make recommendations for steps to be taken in case of exposure. There are two principal forms of smallpox: variola minor and variola major. Smallpox is spread from person to person by means of coughing, direct contact, or contaminated clothing or bedding. Only a few virions are required for transmission. Two other forms of smallpox, hemorrhagic and malignant, are difficult to recognize.

This test is used to determine the presence of the DNA virus responsible for smallpox.

Reference Values

Normal: No Guarnieri’s bodies isolated in scrapings of skin lesions. Absence of brick-shaped virions (ie, variola virus) by electron microscopy. Low levels of neutralizing, hemagglutinin-inhibiting or complement-fixing antibodies.

Procedure

1. Open skin lesions with a blunt instrument (eg, blunt edge of a scalpel) and collect the vesicular or pustular fluid on a cotton swab.

2. Remove scabs with a forceps; they can also be used.
3. Place specimens in a Vacutainer tube; re-stopper and seal it with adhesive tape.
4. Place the Vacutainer tube in a durable, watertight container for transport.
5. Ensure that the laboratory examining the specimens is Biosafety Level 4 (“safety equipment, and facility design and construction are applicable for work with dangerous and exotic agents that pose a high individual risk of life-threatening disease, which may be transmitted via the aerosol route”).
6. Confirm smallpox infection by the appearance of brick-shaped virions under the electron microscope.
7. Remember that definitive laboratory identification requires growth of the virus in cell culture.
8. Use a cotton swab to obtain specimens from the oral cavity or oropharynx if necessary.

Clinical Implications
1. Evidence of virions or Guarnieri's bodies indicate presence of smallpox infection.
2. High levels of antibodies indicate infection.

Interventions

Pretest Patient Care
1. Explain the purpose, procedure, and risks of obtaining a specimen. Obtain pertinent history of signs and symptoms (eg, chills, high fever, backache, pustules that leave a pockmark).
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Care
1. Isolate the individual immediately if the smallpox virus is identified.
2. Report signs and symptoms.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert
1. A confirmed case of smallpox should be brought to the attention of local, state, and federal health authorities.
2. Universal standard precautions should be observed (ie, gloves, gowns, and mask should be worn by laboratory personnel). All laundry and waste should be transported in biohazard bags and autoclaved before being laundered or incinerated.
3. Specimen collection and examination should be performed by laboratory personnel who have recently been vaccinated.
4. All surfaces should be cleaned with hypochlorite (bleach) or quaternary ammonia.
5. Because there are no antiviral drugs, supportive therapy and antibiotics for secondary bacterial infections should be offered to the patient.
6. Household members should be vaccinated and watched closely.

Diagnosis of Tularemia Infection
Tularemia is primarily a rural disease, although occasionally it occurs in urban and suburban areas. *Francisella tularensis* is an intracellular parasite that is spread to humans by infected animals (eg, mice, squirrels, rabbits) or contaminated water, soil, and vegetation. Animals become infected through tick, fly, and mosquito bites. Once infected, person-to-person transmission has not been documented. Two major subspecies of *F. tularensis* have been identified: type A (*F. tularensis* biovar. *tularensis*), which is highly virulent in humans, and type B (*F. tularensis* biovar. *palaearctica*), which is relatively avirulent.

Humans can contract *F. tularensis* through the skin, mucous membranes, lungs, and GI tract. Because *F. tularensis* is one of the most infectious pathogenic bacteria known, its use in biological terrorism cannot be overlooked. Some reports have indicated that it takes only 10 organisms to cause disease.

This test is used to determine the presence of the *F. tularensis* organism.

Reference Values
Normal Absence of the *F. tularensis* organism Negative serum antibody titers

Procedure
1. Obtain specimens of respiratory secretions (ie, sputum), blood, lymph node biopsy samples, or scrapings from infected ulcers.
2. Collect sputum samples after a forced deep cough and place in a sterile, screw-top container.
3. Obtain a 5- to 7-mL Vacutainer from a venipuncture for blood samples.
4. Obtain a skin scraping at the leading edge of a lesion from an infected ulcer and place in a clean, screw-top tube.
5. Perform presumptive identification of *F. tularensis* in a Biological Safety Level 2 clinical laboratory (“practices, equipment, facility design and construction are applicable to clinical, diagnostic, teaching, and other laboratories in which work is done with the broad spectrum of indigenous moderate-risk agents that are present in the community and associated with human disease”).
6. Ensure that confirmation of the organism is done in a Biological Safety Level 3 clinical laboratory (“work is done with indigenous or exotic agents with a potential for respiratory transmission, and which may cause serious and potentially lethal infection”).
Clinical Alert

1. Universal standard precautions should be observed by laboratory personnel (ie, face shields, gowns, gloves).
2. Contaminated clothing or linens should be disinfected per standard precautions protocols.
3. Decontaminate surfaces with a 10% bleach solution.
4. Laboratory personnel who may have had a potential infective exposure should be given prophylactic antibiotics if the risk for infection is high (eg, needle stick).

Clinical Implications Identification of *F. tularensis* and increased antibody titers indicate the presence of tularemia.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, and risks of obtaining a specimen. Obtain history of signs and symptoms, urban or rural living, and occupation (eg, handling infected animal carcasses).
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Care

1. If the *F. tularensis* organism is cultured from the patient, isolation is not recommended because human-to-human transmission has not been documented.
2. Postexposure treatment includes antibiotics such as streptomycin, gentamicin, or ciprofloxacin.
3. Interpret test results, counsel, and monitor appropriately.
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

1. Postexposure prophylactic antibiotic treatment of persons in close contact with the infected patient is not recommended because person-to-person transmission has not been documented.
2. Pregnant women should be treated with ciprofloxacin because rare cases of fetal nerve deafness and renal damage have been reported with some of the aminoglycosides.
3. Suspicion of inhalational tularemia (ie, signs and symptoms) must be reported to local or state public health authorities and the CDC.

DIAGNOSTIC PROCEDURES

Five different categories of laboratory tests are used for the diagnosis of infectious diseases: smears and stains, cultures, tissue biopsy, serologic testing, and skin testing. Cultures and skin testing are described in detail in this chapter; serologic testing is described in Chapter 8. A brief description of each of these procedures follows.

The Smear and Stain

A smear specimen for microscopic study is prepared by rolling a small quantity of the specimen material across a glass slide. If the material is also to be stained, it is generally fixed to the slide by rinsing in methanol. For direct examination of unstained material, phase-contrast microscopy can be used.

Smears are most often observed after they have been stained. Stains are salts composed of a positive and a negative ion, one of which is colored. Structures present in the specimen pick up the stain and make the organism visible under the light microscope. One staining procedure, called the negative stain, colors the background but leaves the organisms themselves uncolored. The gross structure of the organisms can then be studied.

Bacterial stains are of two major types: simple and differential. A simple stain consists of a coloring agent such as gentian violet, crystal violet, carbol-fuchsin, methylene blue, or safranine O. A thin smear of sampled organisms is stained and then observed under an oil-immersion lens. A differential stain is one in which two chemically different stains are applied to the same smear. Organisms that are physiologically different pick up different stains.

The Gram stain is the most important of all bacteriologic differential stains. It divides bacteria into two physiologic groups: gram-positive and gram-negative organisms. The staining procedure consists of four major steps: (1) staining the smear with gentian or crystal violet; (2) washing off the violet stain and flooding the smear with an iodine solution; (3) washing off the iodine solution and flooding the smear with 95% alcohol, or an acetone-alcohol mixture; and (4) counterstaining the smear with safranine O, a red dye. The Gram stain permits morphologic study of the sampled bacteria and divides all bacteria according to their ability or inability to pick up one or both of the stains. Gram-positive and gram-negative bacteria exhibit different properties, which helps to identify and differentiate them.

Stains other than the Gram stain are used for examining bacteriologic smears. Some, such as the acid-fast stain, can identify organisms of the genus *Mycobacterium*. Other stains differentiate certain structures, such as capsules, endospores, or flagella.

Cultures

Preparation of a culture involves growing microorganisms or living tissue cells on a special medium that supports the growth of a given material. Cultures may be maintained in test tubes or Petri dishes. The container holds the culture medium, which is either solid, semisolid, or liquid. Each organism has its own special requirements for growth (proper combination of nutritive ingredients, temperature, and presence or absence of oxygen). The culture is prepared in
accordance with the needs of the organism. Later, it is usually incubated to enhance growth.

**Tissue Biopsy**

At times, microorganisms are isolated from small quantities of body tissue that have been surgically removed. Such tissue is removed aseptically and transferred to a sterile container to be rapidly transported to the laboratory for analysis. Generally, the specimens are finely ground in a sterile homogenizer and then strained and cultured.

**Serologic Testing**

Infectious diseases can be diagnosed by detection of an immunologic response specific to an infecting agent in a patient's serum. Normal humans produce both IgM (first-response antibodies) and IgG (antibodies that may persist long after an infection) to most pathogens. For most pathogens, detection of IgM antibodies or a fourfold increase in the patient's antibody titer is considered to be diagnostic of current infection. If the infecting agent is rare or previous exposure is unlikely (eg, rabies virus, botulin), the presence of specific IgG antibody in a single serum specimen can be diagnostic. Methods for detecting the presence of antibodies include immunodiffusion assay, complement fixation, ELISA, indirect or direct fluorescent antibody, radioimmunoassay, and Western blot immunoassay (see Chap. 8).

**Skin Testing**

Skin testing determines hypersensitivity to the toxic products formed in the body by pathogens. In general, three types of skin tests are performed: scratch tests, patch tests, and intradermal tests.

**Blood Cultures**

Blood cultures are collected whenever there is reason to suspect bacteremia or septicemia. Although mild transitory bacteremia is a frequent finding in many infectious diseases, a persistent, continuous, or recurrent bacteremia indicates a more serious condition that may require immediate treatment. The expeditious detection and identification of pathogens (bacteria, fungi, viruses, and parasites) in the blood may aid in making a clinical and etiologic diagnosis.

**Indications for Blood Culture**

1. Bacteremia
2. Septicemia
3. Shock
4. Unexplained postoperative shock
5. Postoperative shock after genitourinary tract manipulation or surgery
6. Unexplained fever of several days' duration
7. Chills and fever in patients with:
   a. Infected burns
   b. Urinary tract infection
   c. Rapidly progressing tissue infection
   d. Postoperative wound sepsis
   e. Indwelling venous or arterial catheter
8. Debilitated patients receiving:
   a. Antibiotics
   b. Corticosteroids
   c. Immunosuppressives
   d. Antimetabolites
   e. Parenteral hyperalimentation
9. Following body piercing (nose, tongue, nipples, umbilicus) with signs of infection and bacteremia

**Note**

1. During an acute febrile illness, immediately draw two separate blood samples from opposite arms and promptly begin antibiotic therapy.
2. For fever of unknown origin, two blood cultures can be initially drawn 45 to 60 minutes apart. If necessary, two more sets of samples can be drawn 24 to 48 hours later.
3. In cases of acute endocarditis, draw blood cultures as above. If results are negative, two more sets of samples may be obtained on subsequent days.
4. Parasites in the blood (Plasmodium, Trypanosoma, and Babesia) are usually detected by direct microscopic observation.
5. For infants and small children, only 1 to 5 mL of blood can safely be drawn for culture. Quantities < 1 mL may be insufficient to detect bacterial organisms.

**Reference Values**

**Normal** Negative for pathogens

**Procedure of Obtaining Blood Culture** During venipuncture, because of the high potential for infecting the patient, aseptic technique must be used. Key points are listed as follows:

1. Observe standard precautions. The proposed puncture site should be scrubbed with an antiseptic agent such as povidone-iodine, 70% alcohol, or chlorhexidine. Allow to dry for 1 to 2 minutes.
2. Cleanse the rubber stoppers of culture bottles with iodine and allow to air dry. They should then be cleansed with 70% alcohol.
3. Perform venipuncture with a sterile syringe and needle; avoid contamination of the cleansed puncture site.
4. Withdraw about 10 to 30 mL of blood into a 20-mL syringe or directly into the culture tubes. Because of the danger of accidental needle sticks, the practice of changing needles to transfer the specimen into blood culture bottles has been replaced by direct injection with the original phlebotomy needle.
5. Inoculate the anaerobic bottle before the aerobic bottle if two culture bottles are to be inoculated (one anaerobic and one aerobic). Be certain to inoculate each bottle with the optimum blood volume.
6. Mix both bottles gently.
7. Properly label the specimen and immediately transfer them to the laboratory.
8. Cleanse the site with alcohol after the venipuncture because some patients are sensitive to iodine.

**Clinical Alert**

1. Handle all blood specimens according to universal precautions.
2. After disinfection, do not palpate the venipuncture site unless sterile gloves are worn. Palpation is the greatest potential cause of blood culture contamination.
3. The attending physician should be notified immediately about positive culture results so that appropriate treatment may be started.
4. Specimens can be drawn from two or three different sites to exclude a skin-contaminating organism.
5. Collection of more than three blood cultures in a 24-hour period does not improve the detection of bacteria.
6. It is recommended to draw blood below an intravenous line (if possible) to prevent dilution of the sample.

**Clinical Implications**

1. **Negative** cultures: If all cultures, subcultures (if performed), and Gram-stained smears are negative, the blood culture may be reported as no growth after 5 to 7 days of incubation.
2. **Positive** cultures: Pathogens most commonly found in blood cultures include:
   a. *Bacteroides* spp.
   b. *Brucella* spp.
   c. *Enterobacteriaceae*
   d. *Pseudomonas aeruginosa*
   e. *Haemophilus influenzae*
   f. *Listeria monocytogenes*
   g. *Streptococcus pneumoniae*
   h. *Enterococcus* spp.
   i. *S. aureus, Staphylococcus epidermidis*
   j. *Streptococcus* spp. including β-hemolytic streptococci
   k. *Candida albicans*
   l. *Clostridium perfringens*
3. Endocarditis following body piercing

**Interfering Factors** Blood cultures are subject to contamination, especially by skin bacteria. These skin organisms should be identified if possible.

**Interventions**

**Pretest Patient Care**

1. Explain culture purpose and procedure. Obtain pertinent history of signs and symptoms (chills, fever, shock)
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Care**

1. Interpret test results; monitor for bacteremia, septicemia, and other febrile illness; and counsel appropriately about treatment (triple antibiotic therapy).
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Urine Cultures**

Urine cultures are most commonly used to diagnose bacterial urinary tract infection (kidneys, ureter, bladder, and urethra). Urine is an excellent culture and growth medium for most organisms that infect the urinary tract. The combination of pyuria (pus in the urine) and significant bacteriuria strongly suggests the presence of a urinary tract infection.

**Collection of Specimens for Culture: General Principles**

1. Early-morning specimens should be obtained whenever possible because bacterial counts are highest at that time.
2. A clean-voided urine specimen of at least 3 to 5 mL should be collected into a sterile container. Catheterization and aspiration of a suprapubic or indwelling catheter are alternative methods for procuring urine specimens.
3. Urine specimens for culture must never be retrieved from a urine collection bag that is part of an indwelling catheter drainage system.
4. Ideally, urine should be taken to the laboratory and examined as soon as possible. When this is not possible, the urine can be refrigerated for up to 24 hours before being cultured.
5. Whenever possible, specimens should be obtained before antibiotic or antimicrobial therapy begins.
6. Professional health personnel should instruct the patient concerning proper specimen collection technique. Failure to isolate a causative organism is frequently the result of faulty cleansing or collection techniques that can come from lack of information about the proper collection procedure.
7. Provide proper supplies and privacy for cleansing and urine collection. Instruct patients in proper cleansing techniques. The patient who is unable to comply with instructions should be assisted by health care personnel.
8. The urine specimen should be properly labeled. Pertinent information includes:
   a. Patient's identification information
   b. Physician's name
   c. Suspected clinical diagnosis
   d. Method of collection
   e. Precise time obtained
   f. Specific chemotherapeutic agents being administered

   **Clinical Alert**

   Catheterization heightens the risk for introducing bacteria.

   **Clinical Alert**

   1. Urine is an excellent culture medium. At room temperature, it promotes the growth of many organisms. Specimen collection should be as aseptic as possible. Samples should be transported to the laboratory and examined as soon as possible. The specimen must be refrigerated if there is a delay in examination.
   2. In the case of suspected urinary TB, three consecutive early-morning specimens should be collected. Special care should be taken when cleaning the external genitalia to reduce the risk for contamination with commensal acid-fast Mycoplasma/Smegmatis.

**Reference Values**

**Normal**

**Negative**

**Procedure for Collection of Clean-Catch Urine Specimen or Midstream Specimen**

1. For women:
   a. Wash and dry hands thoroughly.
   b. Remove the cap from the sterile container and place it so that only the outer surface touches whatever it is placed on.
   c. Cleanse the area around the urinary meatus from front to back with an antiseptic sponge.
   d. Spread the labia apart with one hand. Hold the sterile container in the other hand, using care not to contaminate the inside surface.
   e. Void the first 25 mL into the toilet, then catch the rest of the urine directly into the sterile container without stopping the urine stream until sufficient quantity is collected. Hold the collection cup in such a way that it avoids contact with the legs, vulva, or clothing. Keep fingers away from the rim and inner surface of the container.
   f. Recap the specimen container, taking care not to contaminate the inside surface of the cap.
   g. Wash and dry hands thoroughly.
   h. Observe standard precautions when handling specimens.

2. For men:
   a. Wash and dry hands thoroughly.
   b. Remove the cap from the sterile container and place it so that only the outer surface touches whatever it is placed on.
   c. Retract the foreskin completely to expose the glans.
   d. Cleanse the area around the meatus with antiseptic sponges.
   e. Void the first 25 mL of urine directly into the toilet and then void a sufficient amount of urine into the sterile specimen container. Do not collect the last few drops of urine.
   f. Recap the specimen container, taking care not to contaminate the inside surface of the cap.
   g. Wash and dry hands thoroughly.
   h. Observe standard precautions when handling specimens.

3. For infants and young children:
   a. Use a suitable plastic collection apparatus to collect urine. Because the collection bag touches skin surfaces and picks up commensal organisms, the specimen must be analyzed as soon as possible.
   b. Cleanse and dry the urethral area thoroughly before applying the collection bag.
   c. Cover collection bag with a diaper or undergarment to prevent dislodging.
   d. Be aware that specimens collected by catheterization may be necessary to detect a urinary tract infection because of the contamination associated with collection bags.

**Clinical Implications**

1. A bacterial count > 100,000 colony forming units (CFU)/mL indicates infection. A mixed bacterial count < 10,000 CFU/mL does not necessarily indicate infection but rather indicates possible contamination. However, growth of a single potential pathogen > 10,000 CFU/mL may be clinically significant in a symptomatic patient.
2. The following organisms, when present in the urine in sufficient quantity, may be considered pathogenic:
   a. *Escherichia coli* and other Enterobacteriaceae
   b. *Enterococcus* spp.
   c. *N. gonorrhoeae*
   d. *M. tuberculosis* (requires special culture media)
   e. *P. aeruginosa*
   f. *S. aureus*
   g. *Staphylococcus saprophyticus*
   h. *Streptococci* (β-hemolytic, usually group B)
   i. *C. albicans* and other yeasts
3. Urine samples obtained by straight catheterization, suprapubic aspiration, or cystoscopy or during surgery represent bladder urine. Growth of any isolate is considered clinically significant.

**Interfering Factors**
1. Patients who are receiving forced fluids may have urine that is sufficiently dilute to reduce the bacterial count to < 100,000 CFU/mL.
2. Bacterial contamination comes from sources such as:
   a. Perineal hair
   b. Bacteria beneath the prepuce in male patients
   c. Bacteria from vaginal secretions, from the vulva, or from the distal urethra in female patients
   d. Bacteria from the hands, skin, or clothing

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test.
2. Ensure that the cleansing procedure is done correctly to remove contaminating organisms from the vulva, urethral meatus, and perineal area so that any bacteria found in the urine can be assumed to have come only from the bladder and urethra.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Care

1. Interpret test outcomes, monitor for urinary tract infection, and counsel appropriately about treatment and possible further testing.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

The urine culture sample should not be taken from a urinal or bedpan and should not be brought from home. The urine should be collected directly into the sterile container that will be used for culture.

Eye and Ear Cultures

Bacterial conjunctivitis, caused by S. pneumoniae, S. aureus, and H. influenzae, is the most common type of infectious conjunctivitis. Inflammation of the cornea usually follows some type of trauma to the ocular surface. Postsurgical and posttraumatic endophthalmitis is often associated with Bacillus spp. and Enterobacteriaceae in addition to the above organisms.

Acute otitis media occurs in the form of a pustule and is often caused by S. aureus. Swimmer's ear is related to maceration of the ear from swimming or hot, humid weather; it often is caused by P. aeruginosa. Otitis media often begins as a viral infection, with a bacterial infection occurring soon afterward. In children, the most common pathogens are S. pneumoniae, H. influenzae, and M. catarrhalis.

Reference Values

Normal Low counts of S. epidermidis, Lactobacillus spp., and Propionibacterium acnes may be found in eye cultures. The same is true for the flora of the external ear.

Procedure for Eye Cultures

1. Observe standard precautions. Purulent material from the lower conjunctival sac or inner canthus of the eye is collected on a sterile swab and placed in transport medium. Both eyes should be cultured separately.
2. Make scrapings of the cornea with a heat-sterilized platinum spatula directly onto the medium (blood or chocolate agar, brain-heart infusion medium for fungi, or thioglycolate broth) in cases of keratitis. For viral culture, the material is placed into viral transport broth.
3. Do not refrigerate specimens or transport on ice. Deliver to the laboratory as soon as possible after collection.

Procedure for Ear Cultures

1. Cleanse the ear with a mild germicide to exclude contaminating skin flora in cases of external otitis.
2. Use a sterile swab or syringe and needle to collect middle ear fluid. Cultures from the mastoid usually are taken during surgery.
3. Do not refrigerate specimens, and deliver to the laboratory as soon as possible after collection.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure for the culture. Record signs and symptoms of ear infection, pain, redness, and/or drainage.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Care

1. Interpret test outcomes, monitor site of infection, and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.
RESPIRATORY TRACT CULTURES

Explanation of Test

Four major types of culture may be used to diagnose infectious respiratory tract diseases: sputum, throat swabs, nasal swabs, and nasopharyngeal swabs. At times, the purposes for which certain tests are ordered overlap.

Reference Values The following organisms may be present in the nasopharynx of apparently healthy persons:

1. *C. albicans*
2. Diphtheroid bacilli
3. *Haemophilus hemolyticus*
4. Staphylococci (coagulase-negative)
5. Streptococci (a-hemolytic)
6. Streptococci (nonhemolytic)
7. Micrococci
8. *Lactobacillus* spp.

Clinical Alert

1. Twenty percent of normal adults carry *S. aureus*; 10% are carriers of group A hemolytic streptococci.
2. A rapid strep test gives results after 10 minutes instead of 24 to 48 hours. It has a false-negative rate of 5% to 10%, about the same as traditional methods. It permits rapid diagnosis and treatment.
3. Both throat and urine cultures are done to detect CMV.

Sputum Cultures

Sputum is not material from the postnasal region and is not spittle or saliva. A sputum specimen comes from deep within the bronchi. Effective coughing usually enables the patient to produce a satisfactory sputum specimen.

Indications for Collection Sputum cultures are important for diagnosis of the following conditions:

1. Bacterial pneumonia
2. Pulmonary TB
3. Chronic bronchitis
4. Bronchiectasis
5. Suspected pulmonary mycotic infections
6. Mycoplasmal pneumonia
7. Suspected viral pneumonia

Reference Values

**Normal** Negative: Normal oral flora

Procedure

1. Instruct patients to provide a deep coughed specimen into a sterile container. Often, an early-morning specimen is best. Expectorated material of 1 to 3 mL is sufficient for most examinations. Remember that good sputum samples depend on thorough health care worker education and patient understanding during the collection process.
2. Label specimens properly and note the suspected disease on the accompanying requisition.
3. Do not refrigerate specimens, and deliver to the laboratory as soon as possible.

Interventions

**Pretest Patient Care**

1. Record signs and symptoms (eg, coughing, productive sputum, blood in sputum)
2. Instruct the patient that this test requires tracheobronchial sputum from deep in the lungs. Instruct the patient to take two or three deep breaths, then to take another deep breath and forcefully cough with exhalation.
3. Ask respiratory therapy personnel to assist the patient in obtaining an “aerosol-induced” specimen if the cough is not productive. Patients breathe aerosolized droplets of a sodium chloride-glycerin solution until a strong cough reflex is initiated. The specimen often appears watery but is in fact material directly from alveolar spaces. It should be noted on the requisition as being aerosol induced.
4. Remember that when pleural empyema is present, thoracentesis fluid and blood culture are excellent diagnostic specimens. Bronchial washings, BAL, and bronchial brush cultures are excellent for detecting most major pathogens of the respiratory tract.
5. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Care**

1. Interpret test outcomes, counsel about treatment, and monitor for respiratory tract infections.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.
Throat Cultures (Swab or Washings)

1. Throat cultures are important for diagnosis of the following conditions:
   a. Streptococcal sore throat
   b. Diphtheria—obtain both throat and nasopharyngeal cultures
   c. Thrush (candidal infection)
   d. Viral infection
   e. Tonsillar infection
   f. Gonococcal pharyngitis
   g. *Bordetella pertussis*

2. Throat cultures can establish the focus of infection in:
   a. Scarlet fever
   b. Rheumatic fever
   c. Acute hemorrhagic glomerulonephritis

3. Throat cultures can be used to detect the carrier state of persons harboring such organisms as:
   a. ß-Hemolytic streptococcus
   b. *Neisseria meningitidis*
   c. *Corynebacterium diphtheriae*
   d. *S. aureus*

Reference Values

**Normal** Negative: Normal oral flora

Clinical Implications Positive findings are associated with infection in the presence of:

1. Group A hemolytic streptococci
2. *N. gonorrhoeae*
3. *C. diphtheriae*
4. *B. pertussis*
5. Adenovirus and herpesvirus
6. *Mycoplasma* and *Chlamydia*

Procedure

1. For adult patients:
   a. Place the patient’s mouth in good visual light.
   b. Use a sterile throat culture kit with a polyester-tipped applicator or swab and a sterile container or tube of culture medium.
   c. Tilt head back. Depress the patient’s tongue with a tongue blade and visualize the throat as well as possible. Rotate the swab firmly and gently over the back of the throat, around both tonsils or fossae, and on areas of inflammation, exudation, or ulceration.
      1. Avoid touching the tongue or lips with the swab.
   d. Place the swab into the designated receptacle so that it comes in contact with the culture medium. Immediately send the specimen to the laboratory.
   e. Refrigerate the throat culture if examination is delayed.

2. For pediatric patients:
   a. Seat the patient in the adult’s lap.
   b. Have the adult encircle the child's arms and chest to prevent the child from moving.
   c. Place one hand on the child's forehead to stabilize the head and to prevent movement.
   d. Proceed with the technique used for collection of the throat and nose culture as described for adults.

3. For throat washings:
   a. Have the patient gargle with 5 to 10 mL of sterile saline solution and then expectorate it into a sterile cup.
   b. Remember that this method provides more specimen than a throat swab and is more definitive for viral isolation.

Interventions

**Pretest Patient Care**

1. Explain purpose and procedure to patient or parents. Record signs and symptoms of sore throat, color of throat, etc.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Care**

1. Interpret test outcomes, monitor for throat infection, and counsel appropriately. Report signs and symptoms.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Nasopharyngeal Cultures (Swab)

Nasopharyngeal swabs are the optimal specimen for detection of *B. pertussis*. Nasopharyngeal swabs, aspirates, and washes are better suited for recovery of respiratory syncytial virus, parainfluenza virus, and viruses causing rhinitis.

**Indications for Collection**

1. Submitted primarily for viral cultures
2. B. pertussis
3. C. diphtheriae

Reference Values

Normal Negative: Normal oral flora

Procedure

1. Tip the patient’s head back to collect a nasopharyngeal specimen.
2. Insert a flexible, calcium alginate–tipped swab carefully through the nose into the posterior nasopharynx and rotate the swab.
3. Pass two swabs simultaneously through one nostril, leave in nasopharynx for 15–30 seconds. Repeat procedure on other nostril with same two swabs. Although the calcium alginate–tipped swabs are most commonly used, aspirated nasopharyngeal specimens, through a soft rubber bulb or plastic-tipped catheter, can be used.
4. Take specimens from both the nasopharyngeal area and the throat for C. diphtheriae confirmation.
5. Handle specimens as follows:
   a. Transport specimens for viral infection in appropriate transport media and refrigerate if not cultured within a few hours.
   b. Do not refrigerate samples unless for diphtheria or pertussis (whooping cough).

OTHER CULTURES AND SMEARS

Wound and Abscess Cultures

Wound infections and abscesses occur as complications of surgery, trauma, or disease that interrupts a skin surface. Material from infected wounds reveals a variety of aerobic and anaerobic microorganisms. Because anaerobic microorganisms are the preponderant microflora in humans and are consistently present in the upper respiratory, GI, and genitourinary tracts, they are also likely to invade other parts of the body to cause severe, and sometimes fatal, infections. Blood cultures should always be drawn from patients with bullous lesions, burn infections, or significant myonecrosis.

Reference Values

Normal Clinical specimens taken from wounds can harbor any of the following microorganisms. Pathogenicity depends on the quantity of organisms present. Quantitative or semi-quantitative reporting of culture results may provide information on the relative importance of the various organisms present in the lesion and also the response of the infection to antibiotic therapy.

1. Actinomyces spp.
2. Bacteroides and Fusobacterium spp.
3. C. perfringens and other species
4. E. coli
5. Other gram-negative enteric bacilli
6. Mycobacterium marinum
7. Nocardia spp.
8. Pseudomonas spp.
9. S. aureus
10. Corynebacterium jeikeium
11. Enterococcus sp.
12. Streptococci (ß-hemolytic)
13. Candida spp.

Procedure

1. Procedure for wound culture
   a. Observe standard precautions.
   b. Be aware that most wounds need some form of preparation to reduce the risk for introducing extraneous organisms into the collected specimen. In the presence of moderate to heavy pus or drainage, irrigate the wound with sterile saline until all visible debris has been washed away. When culturing chronically present wounds (pressure sores), débride the wound surface of any loose necrotic, sloughed material before culturing. Cultures of the surface alone may be misleading; biopsies of deeper tissue are recommended.
   c. Disinfect the surface of the wound with 70% alcohol or an iodine solution.
   d. Apply sterile gauze pads to absorb excess saline and to expose the culture site. Always culture highly vascular areas of granulation tissue. Wearing sterile gloves, separate margins of deep wounds with thumb and forefinger to permit insertion of the swab deep into the wound cavity. Press and rotate the swab several times over the clean wound surfaces to extract tissue fluid containing the potential pathogen. Avoid touching the swab to intact skin at the wound edges.
   e. Immediately place the swab into the appropriate transport container.
2. Procedure for anaerobic collection of aspirated material
   a. Decontaminate the culture site with surgical soap and 70% ethyl or isopropyl alcohol.
   b. Aspirate at least 1 mL of fluid using a sterile 3-mL syringe and a needle of appropriate gauge. Immediately transfer the aspirate to an anaerobic transport medium.
   c. Remember that aspiration cultures are commonly done for closed wounds, such as soft tissue abscesses, cellulitis, or infected skin flaps. Tissue biopsies are more often performed during surgery, when infected tissue is more easily accessible.
   d. Never submit a swab when a tissue sample can be obtained.
Properly label the specimen for the microbiology laboratory with the following:

1. Patient identification information
2. Physician’s name
3. Date and time the specimen was collected
4. Anatomic site or specific source of the specimen
5. Type of specimen (e.g., granulation tissue, abscess fluid, postsurgical wound)
6. Examination requested
7. Patient’s diagnosis
8. Current antibiotic therapy

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**Clinical Alert**

A microscopic examination of pus and wound exudates can be very helpful in diagnosing a pathogenic organism. Consider the following:

1. Pus from streptococcal infections is thin and serous.
2. Pus from staphylococcal infections is gelatinous.
3. Pus from *P. aeruginosa* infections is blue-green.
4. Actinomycosis infections show “sulfur” granules.
5. Bronze discoloration of the skin and fluid-filled blisters are present in gas gangrene.

**Clinical Implications** Clinically significant pathogens are likely to be present in the following specimens:

1. Pus from deep wounds or abscesses, especially if associated with a foul odor
2. Necrotic tissue or débrided material from suspected gas gangrene infection
3. Samples from infections bordering mucous membranes
4. Postoperative wound drainage
5. Lower-extremity ulcers from diabetic patients
6. Decubitus ulcers from elderly or bedridden patients

**Interventions**

**Pretest Patient Care**

1. Explain purpose and wound culture procedure. Record signs of wound infection, type of drainage, fever, etc.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Care**

1. Interpret test outcomes, monitor site of infection, and counsel appropriately about treatment.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Skin Cultures**

The most common bacteria implicated in skin infections are *S. aureus* and *S. pyogenes* (group A). The common abnormal skin conditions include:

1. Pyoderma
   a. Staphylococcal impetigo, characterized by bullous lesions with thin, amber, varnish-like crusts
   b. Streptococcal impetigo, characterized by thick crusts
2. Erysipelas
3. Folliculitis
4. Furuncles
5. Carbuncles
6. Secondary invasion of burns, scabies, and other skin lesions
7. Dermatophytes, especially athlete’s foot, scalp and body ringworm, and “jock itch”

**Reference Values**

*Normal* The following organisms may be present on the skin of a healthy person. When present in low numbers, some of these organisms may be considered normal commensals; at other times, when they multiply to excess, these same organisms may become pathogens.

1. *Clostridium* spp.
2. Coliform bacilli
3. Diphtheroids
4. Enterococci
5. Mycobacteria
7. Staphylococci
8. Streptococci
9. Yeasts and fungi

**Procedure for Obtaining Scrapings from Vesicular Lesions or Skin**
1. Observe standard precautions.
2. Clean the affected site with sterile saline, wipe gently with alcohol, and allow it to air dry.
3. Aspirate a fluid sample from fresh, intact vesicles with a 25-gauge needle attached to a tuberculin syringe, and transfer the specimen to the transport medium by ejecting it from the syringe.
4. If fluid cannot be aspirated, open the vesicles and use a cotton-, rayon-, or Dacron-tipped applicator to swab the base of the lesion to collect infected cells. Place the swab directly into transport medium (eg, self-contained foam pad with Stuart's media).
5. To make smears for stains, use a scalpel blade to scrape the base of the lesion, taking care not to macerate the cells. Spread scraped material in a thin layer on a slide.
6. Place the specimen in biohazard bag; do not refrigerate. Immediately transport the specimen to the laboratory for bacterial, fungal, or viral cultures.

Clinical Alert
The most useful and common specimens for detection of fungal infection are skin scrapings, nail scrapings, and hairs (see Diagnosis of Fungal Disease and Appendix K).

Clinical Implications When present on the skin in significant quantities, the following organisms may be considered pathogenic and indicative of an abnormal condition:

1. Enterobacteriaceae
2. Fungi (Sporotrichum, Actinomycoses, Nocardia, C. albicans, Trichophyton, Microsporum, Epidermophyton)
3. S. aureus
4. Streptococcus pyogenes
5. P. aeruginosa
6. Varicella-zoster virus
7. Herpes simplex virus

Interventions
Pretest Patient Care

1. Explain purpose and skin culture procedure.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Care

1. Interpret test outcomes, monitor site of infection, and counsel appropriately about treatment. Report rashes, fever, etc.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Stool and Anal Cultures and Smears

Stool cultures are commonly done to identify bacteria associated with enteric disease. Of all specimens collected, feces are likely to contain the greatest number and greatest variety of organisms. For a routine stool culture, the stool is examined to detect and to rule out Salmonella, Shigella, Campylobacter, Aeromonas, Plesiomonas, and predominating numbers of Staphylococcus organisms; cultures for yeast, Bevdomonas, Yersinia, Vibrio, and Shiga toxin–producing E. coli have to be specifically requested, depending on laboratory practice. Clostridium difficile causes antibiotic-associated colitis. It is diagnosed by detection of the toxins.

A single negative stool culture should not be considered the endpoint in testing. At least three stool cultures collected on separate days are recommended if the patient's clinical picture suggests bacterial involvement, despite previous negative cultures. Moreover, once a positive diagnosis has been made, the patient's personal contacts should also be tested to prevent a potential spread of infection.

Reference Values
Normal The following organisms may be present in the stool of apparently healthy people:

1. C. albicans
2. Enterococcus spp.
3. E. coli
4. Proteus spp.
5. P. aeruginosa
7. Staphylococcus spp.

Procedure

1. Procedure for stool specimen collection
   a. Observe standard precautions.
   b. Use a dry container or a clean, dry bedpan to collect feces. Do not contaminate stool specimen with urine, soap, or disinfectants.
   c. Remember that a freshly passed stool is best. Diarrheal stool usually gives acceptable results.
   d. Select portions containing pus, blood, or mucus. A 1- to 2-gram quantity is sufficient.
   e. Do not retrieve stool from the toilet for specimen use.
f. Do not place toilet tissue or diapers with the specimen. It may contain bismuth, which interferes with laboratory tests.
g. Transfer stool specimens from the bedpan to the container with tongue blades.
h. Properly label the sealed specimen container and immediately send it to the laboratory.
i. Place the specimen in a transport medium, such as Cary-Blair medium, if a delay of longer than 2 hours for stool culture is anticipated (from time of collection until receipt in the laboratory). Specimens processed within 2 hours of collection do not require added preservatives. Place the designated volume of stool into the transport container.

2. Procedure for obtaining a rectal swab
   a. Observe standard precautions.
   b. Insert the swab gently into the rectum (to a depth of at least 3 cm) and rotate it to retrieve a visible amount of fecal material (Fig. 7.1).

   FIGURE 7.1 Method for obtaining the rectal culture.

   c. Place the swab into the receptacle containing transport medium, such as Cary-Blair medium.
   d. Properly label the specimen and send it in a biohazard bag to the laboratory as soon as possible.

3. Procedure for performing cellophane tape test for pinworm (*Enterobius vermicularis*)
   a. Observe standard precautions. The tape test is indicated in cases of suspected enterobiasis (pinworms).
   b. Apply a strip of clear cellophane tape (not micropore or adhesive type tape) to the perineal region. Remove and spread the tape on a slide for microscopic examination.
   c. Remember that a paraffin-coated swab can be used in place of the cellophane tape test. If used, place the swab within a stoppered test tube.
   d. Be aware that it may be necessary to make four to six examinations on consecutive days before ruling out the presence of pinworms.
   e. Test for pinworm eggs in the morning, before the patient has defecated or bathed. For small children, it is best to collect the specimen just before the child awakens.

**Clinical Alert**

Fecal specimens are far superior to rectal swab specimens. Often, rectal swabs reach only the anal canal and provide material of limited diagnostic significance.

**Clinical Implications**

1. *C. albicans*, *S. aureus*, and *P. aeruginosa*, found in large numbers in the stool, are considered pathogenic in the setting of previous antibiotic therapy. Alterations of normal flora by antibiotics often change the environment so that normally harmless organisms become pathogens.
2. *Cryptosporidiosis* is a cause of severe, protracted diarrhea in immunosuppressed patients. *Cryptosporidium* organisms can be detected by ova and parasite examination.
3. *H. pylori* has been associated with gastritis and peptic ulcer disease. *H. pylori* is found only on the mucus-secreting epithelial cells of the stomach. Detection of *H. pylori* in gastric biopsy specimens necessitates collection of the specimens in sterile containers. Smears and cultures should be examined for the presence of this organism. Initial culture incubation requires 7 days. Therefore, results of gastric biopsy specimen cultures may take 8 to 10 days to obtain. A test for *H. pylori* antigen in the stool provides rapid detection of *H. pylori*.
4. *C. difficile*: Whenever normal flora are reduced by antibiotic therapy or other host factors, the syndrome known as *pseudomembranous colitis* occurs. This condition is caused by *C. difficile*. It may be present in small numbers in the normal person, or it may occur in the hospital environment. When normal flora are reduced, *C. difficile* can multiply and produce its toxins.

The definitive diagnosis of *C. difficile*-associated diarrhea is based on clinical criteria. Endoscopic visualization of a characteristic pseudomembrane or plaque, together with a history of antibiotic therapy, is diagnostic of *C. difficile*. Three laboratory tests are also available. These include stool culture for *C. difficile* (nonspecific; requires at least 48 hours); tissue culture for detection of cytoxin (48 hours); and rapid antigen tests for toxins that are sensitive and specific for *C. difficile*.

**Interfering Factors** Feces from patients receiving barium, bismuth, mineral oil, or antibiotics are not satisfactory specimens for identifying protozoa.

**Interventions**

**Pretest Patient Care for Stool Specimen Collection**

1. Explain purpose and procedure. Obtain history of diarrhea including type and length of time. Instruct the patient to defecate into a clean, dry bedpan or large-mouthed container.
2. Do not allow patient to defecate into the toilet bowl or urinate into the bedpan or collecting container because urine has an adverse effect on protozoa.
3. Do not place toilet paper into the bedpan or collection container; it may contain bismuth, which can interfere with testing.
4. Follow Chapter 1 guidelines for **safe, effective, informed pretest care**.

**Posttest Patient Care**

1. Interpret test outcomes, monitor for intestinal infection, and counsel appropriately about treatment and possible further testing. Report signs and symptoms.
2. Follow Chapter 1 guidelines for **safe, effective, informed posttest care**.

**Clinical Alert**

1. In the institutional setting, patients with diarrhea should remain in isolation until the cause for the diarrhea is determined.
2. When pathogens are found in the diarrheic stool, the patient usually remains isolated until the stool becomes formed and antibiotic therapy is completed.

**Cerebrospinal Fluid (CSF) Cultures and Smears**

Bacteriologic examination of CSF is an essential step in the diagnosis of any case of suspected meningitis. Acute bacterial meningitis is an infection of the meninges (the membrane covering the brain and spinal cord). It is a rapidly progressive, fatal disease if left untreated or if treated inadequately. Death can occur within hours of symptom onset. Prompt identification of the causative agent is necessary for appropriate antibiotic therapy and aggressive treatment. Meningitis is caused by a variety of gram-positive and gram-negative microorganisms. Bacterial meningitis also can be secondary to infections in other areas of the body.

A smear and culture should be performed on all CSF specimens obtained from persons with suspected meningitis, whether the CSF fluid appears clear (normal) or cloudy.

In bacterial meningitis (except TB meningitis), the CSF shows the following characteristics:

   1. Purulence (usually)
   2. Increased numbers of leukocytes
   3. Preponderance of polymorphonuclear cells
   4. Decreased CSF glucose concentration in relation to serum glucose
   5. Elevated CSF protein concentration

In meningitis caused by the tubercle bacillus, viruses, fungi, or protozoa, the CSF shows the following characteristics:

   1. Nonpurulent (usually)
   2. Decreased mononuclear white cell count; increased lymphocytes
   3. Normal or decreased CSF glucose concentration
   4. Elevated CSF protein concentration

In those persons with suspected meningitis, the CSF fluid is generally submitted for chemical and cytologic examinations as well as culture.

**Indications for Collection**

1. Viral meningitis
2. Pyogenic meningitis
3. TB meningitis
4. Chronic meningitis

**Reference Values**

**Normal**

1. Negative: No growth
2. Bacteria are not normally present in CSF. However, the specimen may be contaminated by normal skin flora during the process of CSF procurement.

**Procedure**

1. Collect the specimen under sterile conditions. Three or four tubes (1.0 mL per tube) of CSF should be collected. The third tube is used for cell count and differential; the others can be used for microbiologic and chemical studies.
2. Seal immediately to prevent leakage or contamination, and send it to the laboratory without delay.
3. Label the specimen properly. Alert laboratory staff so that the specimen can be examined immediately.
4. Notify the attending physician as soon as results are obtained so that appropriate treatment can be started in a timely fashion.
Clinical Alert

In cases of suspected meningitis, a culture should be done and a diagnosis made as quickly as possible. This is important because some causative organisms cannot tolerate temperature changes.

If a viral cause is suspected, a portion of the CSF fluid should be refrigerated (0°C to 4°C). Freezing is not recommended unless inoculation into tissue culture will take longer than 5 days. If PCR testing is to be performed, specimens may need to be frozen immediately.

Clinical Alert

Newborns have the highest prevalence of meningitis of any age group. Organisms causing disease in the newborn (usually acquired during the birth process) include group B streptococcus, *E. coli*, and *L. monocytogenes*.

Clinical Implications

1. Pathogens found in CSF include:
   a. *Cryptococcus* and other fungi
   b. *H. influenzae*
   c. *Naegleria* or *Acanthamoeba* spp.
   d. Viruses (usually enteroviruses)
   e. *L. monocytogenes*
   f. *M. tuberculosis*
   g. *N. meningitidis*
   h. *S. pneumoniae*
   i. *S. aureus*
   j. *S. epidermidis*
   k. *Streptococcus* (group B)
   l. *T. pallidum*
   m. *Toxoplasma gondii*

2. Positive CSF cultures occur in:
   a. Meningitis
   b. Trauma
   c. Abscess of brain or ependyma of spine
   d. Septic thrombophlebitis of venous sinuses

Maintenance of Culture

1. If the CSF specimen cannot be delivered to the laboratory immediately, the container should be stored at room temperature.
2. No more than 4 hours should elapse before laboratory analysis takes place because of the low survival rates of the organisms causing meningitis (especially *H. influenzae* and *N. meningitidis*).

Interventions

**Pretest Patient Care**

1. Explain purpose and lumbar puncture procedure (see Chap. 5). Record pertinent signs and symptoms.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Care**

1. Interpret test outcomes, monitor for meningitis, and counsel appropriately (see Chap. 4).
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Cervical, Urethral, Anal, and Oropharyngeal Cultures and Smears for Gonorrhea and Other Sexually Transmitted Diseases

These tests are done for patients with genital ulcers, vaginal lymphadenopathy, bacterial vaginosis, lesions affecting epithelial surfaces, signs and symptoms of bacterial STDs, pelvic inflammatory disease, urethritis, or abnormal discharge and itching.

**Reference Values**

*Normal* Negative: Normal flora; pathogens not detected

**Procedures for Specimens**

1. Cervical (female patients)
   a. Be aware that the cervix is the best site from which to obtain a culture specimen.
   b. Observe standard precautions.
   c. Moisten the vaginal speculum with warm water; do not use a lubricant. Remove cervical mucus, preferably with a cotton ball held in a ring forceps.
   d. Insert a sterile, cotton-tipped swab into the endocervical canal; move the swab from side to side; allow 30 seconds for absorption or organisms by the swab (Fig. 7.2).
2. Urethral (male patients)
   a. Use a sterile swab to obtain the specimen from the anterior urethra by gently scraping the urethral mucosa (Fig. 7.3).

   FIGURE 7.3 Method for obtaining the urethral specimen.

b. Rotate the swab 360 degrees to dislodge some of the epithelial cells for *Chlamydia. N. gonorrhoeae* organisms inhabit the exudate, whereas *C. trachomatis* organisms are intracellular (within the epithelial cells).

3. Anal canal
   a. Insert a sterile, cotton-tipped swab approximately 2.5 cm into the anal canal. (If the swab is inadvertently pushed into feces, use another swab to obtain the specimen.)
   b. Move the swab from side to side in the anal canal to sample the crypts; allow several seconds for absorption of organisms by the swab.
   c. Remember that this site is likely to be positive in a patient with STD, when a cervical specimen is negative.

Because *Trichomonas vaginalis* may be present in urethral or vaginal discharge, material for culture should be collected as described; however, an additional swab should be placed in a tube containing 0.5 mL of sterile saline and be delivered to the laboratory immediately. Swabs for culture should be transported to the laboratory in Stuart’s transport medium and should be held at room temperature until processed. If specimens are not processed within 12 hours, they should be refrigerated. Recovery of a pathologic organism may be more difficult because of delay in processing.

### Clinical Alert

1. If the male urethral culture is negative but gonorrhea is still suspected, prostatic massage may produce an increased number of organisms in the urethral discharge. The first morning specimen before urination may be the best.
2. In a female patient, the anal canal specimen can be obtained after the cervical specimen without changing the patient’s position and without using the anoscope. Observe standard precautions.

### Clinical Alert

The finding of repeated negative cultures for gonococci does not always exclude a diagnosis of gonorrhea.

### Interventions

**Pretest Patient Care**

1. Explain culture purpose and collection procedure. Obtain history of pertinent signs and symptoms (drainage, pain, itching).
2. Place the patient in the dorsal lithotomy position and appropriately drape for genital procedures. Provide as much privacy as possible.
3. Follow standard precautions.
4. Follow Chapter 1 guidelines for **safe, effective, informed pretest care**.

**Posttest Patient Care**

1. Interpret test outcomes and counsel appropriately.
2. Explain need for possible follow-up testing and treatment.
3. Follow Chapter 1 guidelines for **safe, effective, informed posttest care**.

### Tissue, Bone, and Body Fluid Cultures

Types of fluid collected for bacterial, viral, or fungal culture include pleural, ascitic, synovial, and pericardial fluid. Tissues may have to be minced or ground to release trapped bacteria before culturing.

### Reference Values

**Normal** Negative for pathogens

### Procedure for Collection of Specimens

1. Transport body fluids to the laboratory in a sterile tube or sterile capped syringe. Ten to 20 mL of fluid is adequate for culture examination.
2. Collect bone during surgery and send to the laboratory in a sterile container. Place fragments directly onto the agar surface or into enrichment broth.

3. Collect pieces of tissue during surgery or during needle biopsy procedures. They should be collected in a sterile specimen cup. Add a small amount of sterile, nonbacteriostatic saline to keep specimen moist.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure for the culture.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Care**

1. Interpret test outcomes; monitor site of collection, and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**SKIN TESTS**

Skin testing is done for three major reasons: (1) to detect sensitivity to allergens such as dust and pollen, (2) to determine sensitivity to microorganisms believed to cause disease, and (3) to determine whether cell-mediated immune functions are normal. The test that detects sensitivity to allergens is mentioned only briefly in this chapter below; most of this discussion focuses on skin tests used to determine sensitivity to pathogens.

**Reference Values**

**Normal** Positive reactions indicate lack of immunity to a specific disease (eg, TB-producing agent) or sensitivity to a specific allergen (eg, mold).

**Intradermal Tests**

The substance being tested is injected into the layers of skin with a tuberculin syringe fitted with a short-bevel, 26- or 27-gauge needle. A positive reaction produces a red, inflamed area at the site of the injection within a given time period (eg, 72 hours for the Mantoux test for TB).

Skin tests that indicate hypersensitivity to a toxin from a disease-producing agent may also signal immunity to the disease. Positive reactions may also indicate an active or inactive phase of the disease under study. Skin tests can be categorized according to their nature and purpose as follows:

1. Test to reveal a present or past exposure to the infectious agent; for example, tuberculin test (positive reaction indicates presence of active or inactive TB).
2. Tests to show sensitivity to materials toward which a person may react in an exaggerated manner; for example, allergenic extracts such as house dust and pollen (positive reaction indicates sensitivity to allergen extracts).
3. Tests to detect impaired cellular immunity. Intradermal skin testing with several common antigenic microbial substances (eg, purified protein derivative [PPD] tuberculin, mumps virus, C. albicans, streptokinase-streptodornase) can determine whether immune function is normal. This would be important in treating leukemia and cancer with chemotherapy. (Negative reaction to any intradermal antigen indicates impaired immunity.)

**Procedure for Skin Tests**

1. Follow the manufacturer’s instructions for the diagnostic skin tests. Most are prepackaged as sterile kits.
2. Inject 0.1 mL of the test material intradermally on the volar aspect of the forearm.
3. Remember that a positive reaction is manifested by redness or swelling > 1 cm in diameter at the injection site. A central area of necrosis is a highly significant finding.

**Clinical Alert**

Material for diagnostic skin tests may be inadvertently injected into subcutaneous tissue rather than intradermal tissue. A subcutaneous injection yields a false-negative result.

See individual skin tests for pretest and posttest interventions.

**Tuberculin Skin Test (TB Test); Two-Step TB Test**

The intradermal tuberculin skin test detects TB infection; it does not distinguish active TB from dormant TB. PPD tuberculin is a protein fraction of the tubercle bacilli; when it is introduced into the skin of a person with active or dormant TB infection, it causes a localized skin erythema and induration at the injection site because of accumulated small, sensitized lymphocytes.

The Mantoux test is the test of choice. The tuberculin is injected into the intradermal skin layer with a syringe and fine-gauge needle. The multiple puncture test (tine test) is used for screening purposes for asymptomatic persons, but the Mantoux test is far more accurate.
The two-step TB skin test is done to reduce the likelihood that a “boosted” reaction will be interpreted as a recent infection. The two-step skin test is not routine for contact case investigation.

Indications for Testing

1. Persons who exhibit signs (x-ray film abnormality) or symptoms (eg, cough, hemoptysis, weight loss) suggestive of TB
2. Recent close contacts with persons known to have or suspected of having TB
3. Persons who show abnormal chest radiographs compatible with past TB exposure
4. Members of groups at high risk for M. tuberculosis infection, such as immigrants from Asia, Africa, and Latin America; poverty-prone and “skid row” populations; personnel and long-term residents of health care facilities and institutions (eg, nursing homes, mental institutions, prisons)
5. The two-step test is indicated for noninfected new employees and new residents of institutions (eg, nursing homes, hospitals, homeless shelters, correctional institutions, alcohol and drug treatment centers), persons 55 years of age and older, and persons born in countries with high prevalence.

Reference Values

Normal Reaction negative or not significant

Procedure for Intradermal Skin Test (Mantoux)

1. Observe standard precautions. Draw up PPD tuberculin into a tuberculin syringe (follow manufacturer's directions carefully) with a 0.5-inch, 26- or 27-gauge needle. Use 0.1 mL (5 tuberculin units) for each test.
2. Cleanse the skin on the volar or dorsal aspect of the forearm with alcohol and allow it to dry.
3. Stretch the skin taut.
4. Hold the tuberculin syringe close to the skin so that the hub of the needle touches the skin as the needle is introduced under the skin. A discrete, pale elevation of the skin (wheal) 6 to 10 mm in diameter should be produced when the prescribed amount of PPD tuberculin is injected into the intradermal skin layer.
5. For the two-step test, administer the Mantoux intradermal skin test, as described, for all persons for whom testing is indicated. Strictly enforce reading of results in 48 to 72 hours. If the result is positive, do not administer a second PPD dose but refer the patient for follow-up. If induration is present but does not classify as positive, retest immediately on the patient's other arm and read the results in 48 to 72 hours. If the result of the first Mantoux test is negative, retest in 1 to 2 weeks, using the same PPD dose and the same arm as for the first test. Read the results in 48 to 72 hours. If the reaction at second test is negative (no duration), perform no further testing now. Make plans to administer the one-step Mantoux test yearly (or every 3 to 6 months if the patient is at high risk).

Clinical Implications

1. The test should be read at 48 to 72 hours after injection. The larger the area of the skin reaction, the more likely it is to represent TB infection. Positive tests show an indurated area of 5 to 15 mm. However, a significant reaction to the skin test does not necessarily signify the presence of TB.
2. A significant reaction does not distinguish between active and dormant TB infection; the stage of infection can be determined from the results of clinical bacteriologic sputum tests and chest roentgenograms.
3. A significant reaction in a clinically ill patient means that active TB should be considered as a cause for illness. With HIV infection, a reaction of 5 mm or more is considered positive.
4. A significant reaction in a healthy person usually signifies either healed TB or an infection caused by a different mycobacterium. Chest roentgenograms can confirm the absence of an active disease process.

Interfering Factors False-negative results may occur even in the presence of active TB or whenever sensitized T lymphocytes are temporarily depleted in the body.

Reading the Test Results

1. The test should be read 48 to 72 hours after injection.
2. Examine the injection site in good light.
3. The patient should flex the forearm at the elbow.
4. Inspect the skin for induration (hardening or thickening).
5. Rub finger lightly from the normal skin area to the indurated zone (if present).
6. Circle the zone of induration with a pencil and measure the diameter in millimeters perpendicularly to the long axis of the forearm. Disregard erythema; it is clinically insignificant.
7. Large reactions may still be evident 7 days after the test.
Clinical Alert

1. Tuberculin test material should never be transferred from one container to another.
2. Intradermal skin tests should be given immediately after the tuberculin is drawn up.
3. The greatest value of tuberculin skin testing is in the negative results; a negative test result in the presence of signs and symptoms of lung disease is strong evidence against active TB in most cases.
4. A presumptive diagnosis of TB must be bacteriologically confirmed.
5. In the United States, the incidence of TB is higher among older persons, men, nonwhites, and foreign-born persons.
6. Sixteen percent of TB cases are extrapulmonary.
7. TB is acquired through close, frequent, and prolonged exposure to infected persons.
8. A person diagnosed with TB has on average nine contacts, of whom 21% are infected.
9. Persons who have received Bacille Calmette-Guérin (BCG) vaccine prophylactically or for bladder cancer treatment test positive for TB. Reactions of 5 to 10 mm may be caused by BCG vaccination. However, unless the vaccination was very recent, tuberculin reactions greater than 10 mm should not be attributed to BCG.
10. Periodic chest x-ray films are valuable adjuncts for monitoring patients who test positive because there is no sure way of predicting who will develop active TB.
11. BCG is a freeze-dried preparation of a live, attenuated bovine strain of mycobacteria. It is used for TB immunization in children (eg, infant with a negative TB test who lives in a household with untreated or ineffectively treated cases of TB) in countries with a high incidence of TB.
12. Clinicians in contact with suspected or confirmed TB must wear a properly fitted, high-efficiency, dust- and mist-proof mask.

Interpreting the Test Results

1. The test interpretation is based on the presence or absence of induration.
2. Negative or insignificant reaction: zone of induration < 5 mm in diameter. Positive or significant reaction: zone of induration > 10 mm in diameter.
3. For persons in good health with no risk factors, an induration of 15 to 20 mm usually is considered positive. However, because those who are at increased risk for TB (in poor health) have decreased hypersensitivity, a 5-mm induration may be considered positive. Retest within 3 weeks. See Chart 7.2 for classification of test results.

Chart 7.2 Classification of the Tuberculin Skin Test Reaction

An induration of 5 or more millimeters is considered positive for:
- HIV-infected persons
- close contacts of a person with infectious TB
- persons who have abnormal chest radiographs
- persons who inject drugs and whose HIV status is unknown

An induration of 10 or more millimeters is considered positive for:
- foreign-born persons
- HIV-negative persons who inject drugs
- medically underserved, low-income populations
- residents of long-term care facilities
- persons with certain medical conditions *
- children < 4 years old without any other risk factors
- staff of long-term care facilities and health care facilities

An induration of 15 or more millimeters is considered positive for:
- persons who do not have any risk factors for TB

Footnote

*For example, diabetes mellitus, prolonged corticosteroid therapy, immunosuppressive therapy, gastrectomy, some hematologic and reticuloendothelial diseases, end-stage renal disease, silicosis, and body weight that is 10% or more below ideal. From Centers for Disease Control and Prevention (CDC), Tuberculosis Information, Diagnosis of TB Infection and TB Disease, August 25, 1997, Document 250102.

Potential Causes of False-Negative Results

Reactions can be categorized according to the following factors:

1. Factors related to person being tested
   a. Presence of infections
   b. Viral (measles, mumps, chickenpox)
   c. Live virus vaccinations (measles, mumps, polio)
   d. Nutritional factors (severe protein depletion)
   e. Diseases affecting lymphoid organs (Hodgkin's disease, lymphoma, chronic lymphocytic leukemia, sarcoidosis)
   f. Drugs (corticosteroids, other immunosuppressive agents)
   g. Age (newborns, elderly patients with "waned" sensitivity)
   h. Recent or overwhelming M. tuberculosis infection

2. Factors related to tuberculin injected
   a. Improper storage (exposure to light, heat)
   b. Improper dilution
c. Chemical denaturation

d. Contamination

e. Absorption (partially controlled by adding Tween-80)

f. Outdated material

3. Factors related to method of administration
   a. Injection of too little or too much antigen
   b. Delayed administration after drawing up dose
   c. Injection too deep or too shallow

4. Factors related to test interpretation and recording of results
   a. Test not read within prescribed time frame
   b. Inexperienced reader
   c. Conscious or unconscious bias
   d. Recording error
   e. Measurement error

Interventions

Pretest Patient Care

1. Explain TB skin test purpose and procedure and the necessity of returning for “reading” of the skin reaction. Obtain
   history of occupation, living condition, and reason for testing.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Care

1. Interpret test outcomes at the prescribed time; monitor and counsel appropriately about need for chest radiograph
   and sputum cultures for those with positive TB skin tests. Discuss initial and continued therapy and institute
   infection and case control as required. The possibility of TB disease must be ruled out before preventive therapy
   can start. TB is a reportable disease.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Mumps Test

Mumps, the common disease that produces swelling and tenderness of the parotid glands, is caused by a myxovirus.

An antigen made from infected monkeys or chickens is injected intradermally. A positive mumps skin test may indicate
either a previous infection or an existing infection; therefore, it is not very effective as a diagnostic tool. The test is used
primarily as part of a battery of skin tests to determine immunocompetence.

Reference Values

Normal Same as described subsequently under Clinical Implications

Procedure

1. Observe standard precautions. Before injecting antigen, assess for allergy to eggs. Persons who are allergic to
   eggs are at risk for an anaphylactic reaction to mumps antigen.
2. Inject mumps antigen intradermally.

Clinical Implications

1. A positive reaction indicates resistance to the mumps virus.
2. A negative reaction indicates susceptibility to mumps virus.

Interpretation of the Test Results

1. Read the test 48 hours after the time of injection.
2. Positive reaction: erythema and a lesion > 10 mm in diameter.
3. Negative reaction: no erythema and a lesion < 10 mm in diameter.

Interventions

Pretest Patient Care

1. Explain skin test purpose and procedure.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Care

1. Interpret test outcomes regarding immunocompetence.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Candida and Tetanus Toxoid Tests

Candida and tetanus toxoid are additional skin tests that can be done to detect delayed-type hypersensitivity. The
Candida antigen is a mixture of trichophytin and Oidium. Both antigens are administered in a manner similar to the
 tuberculin skin test.
To interpret these skin tests for anergy, the following Centers for Disease Control and Prevention guidelines are recommended:

1. For high-risk patients (HIV infection, intravenous drug abuse, immunocompromise), an induration area > 5 mm is considered positive.
2. For patients at moderate risk (institutionalized patients, health care workers), an indurated area > 10 mm is significant.
3. In patients with no significant risk factors, an indurated area of 15 mm or larger is considered positive.

These additional skin tests are helpful in evaluating a negative PPD test in an immunosuppressed person. No reaction to mumps, tetanus, or *Candida* testing may indicate a false-negative PPD test. However, an induration > 2 mm with the mumps, *Candida*, or tetanus antigen confirms the negative PPD result.

**BIBLIOGRAPHY**


Centers for Disease Control and Prevention. Basic laboratory protocols for the presumptive identification of *Bacillus anthracis*. Atlanta, Author, 2001

Centers for Disease Control and Prevention. 2002 Sexually transmitted diseases treatment Guidelines. MMWR 51(RR-06), 2002


Forbes BA, Sahm DF, Weissfeld AS: Bailey’s and Scott’s Diagnostic Microbiology, 11th ed. St Louis, Mosby, 2002


OVERVIEW OF IMMUNODIAGNOSTIC STUDIES

Types of Tests
Collection of Serum for Immunologic Tests
Interpreting Results of Immunologic Tests
Serologic Versus Microbiologic Methods

BACTERIAL TESTS
Syphilis Detection Tests
Lyme Disease Tests
Legionnaire's Disease Antibody Test
Chlamydia Antibody IgG Test
Streptococcal Antibody Tests: Anti-Streptolysin O Titer (ASO), Streptozyme, Anti-DNase B (ADB, Streptodornase)
Helicobacter pylori (HPI) IgG Antibody Serum, Stool, and Breath (PF) Test

VIRAL TESTS
Erythema-Majus Ulcer (EMU) Antibody Tests: Herpes Simplex Virus Type 1 (HSV-1) and Type 2 (HSV-2) Tests

VIRAL ANTIBODY TESTS TO ASSESS IMMUNE STATUS
Rubella Antibody Tests
Measles (Rubeola) Antibody Tests
Mumps Antibody Tests
Varicella-Zoster (Chickenpox) Antibody Test
Cytomegalovirus (CMV) Antibody Test

PARASITIC TESTS
Toxoplasmosis (TPM) Antibody Tests
Amebiasis (Entamoeba histolytica) Antibody Test
TORCH Test

IMMUNOLOGIC TESTS FOR IMMUNE DYSFUNCTION AND RELATED DISORDERS OF THE IMMUNE SYSTEM
Quantitative Immunoglobulins: IgA, IgG, IgM
Protein Electrophoresis (PEP), Serum and Urine
Immunofixation Electrophoresis (IFE), Serum and Urine
Cold Agglutinin
Cryoglobulin Test
Total Hemolytic Complement (CH50)
C3 Complement Component
C4 Complement Component

TESTS FOR AUTOIMMUNITY AND SYSTEMIC RHEUMATIC DISEASE (SRD)
Antinuclear Antibody (ANA) Test
Anticentromere Antibody Test
Anti-dsDNA Antibody Test, IgG
Rheumatoid Factor (RF) Antibody (RAF) Test
Antibodies to Extractable Nuclear Antigens (ENA): Anti-Ribonucleoprotein (RNP), Anti-Smith (Sm), Anti-Sjögren's Syndrome (SSA, SSB), Anti-Scleroderma (Scl-70)
Anti-Jo-1 (Jo-1)
Complement Antibodies, IgA, IgG, IgM
Autoimmune Thyroiditis, Thyroid Antibody Tests: Thyroglobulin Antibody, Thyroid Microsomal Antibody, Thyroxine Oxidase Antibody

AUTOIMMUNE LIVER DISEASE TESTS
Anti-SMOOTH MUSCLE ANTIBODY (ASMA) Test
Anti-MITOCHONDRIAL ANTIBODY (AMA) Test
Anti-Liver/Kidney Microsomal Type 1 Antibody (LKM) Test
Anti-Parietal Cell Antibody (APCA) Test
Antiglomerular Basement Membrane (AGBM) Antibody Test
Acetylcholine Receptor (ACHR) Binding Antibody Test
Anti-Insulin Antibody Test
Glutathione S-transferase (GST) Antibodies, IgG and IgM

ANTINEUTROPHIL CYTOPLASMIC ANTIBODIES (ANCA)
Anti-Sperm Antibodies
Allergy Testing
IgE Antibody, Single Allergen
Protein Chemistry Testing/serum proteins: acute-phase proteins and cytokines
Ceruloplasmin
Table 8.1 Some Tests That Determine Antigen-Antibody Reactions

<table>
<thead>
<tr>
<th>Name of Test</th>
<th>Observable Reaction</th>
<th>Visible Change</th>
<th>Tests for</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agglutination, hemagglutination (HA), immune hemagglutination assay (IHA)</td>
<td>Particulate antigen reacts with corresponding antibody; antigen may be in form of RBCs (hemagglutination, latex, or charcoal coated with antigen).</td>
<td>Clumping</td>
<td>Treponemal, heterophile, and cold agglutinin antibodies</td>
</tr>
<tr>
<td>Procedure</td>
<td>Description</td>
<td>Application</td>
<td></td>
</tr>
<tr>
<td>-------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Precipitation (eg, immunodiffusion [ID]; counter-immunoelectro-phoresis [CIE])</td>
<td>Soluble antigen reacts with corresponding antibody by ID or count.</td>
<td>Precipitates</td>
<td>Fungal antibodies, food poisoning</td>
</tr>
<tr>
<td>Complement fixation (CF)</td>
<td>Competition between two antigen-antibody systems (test and indicator systems)</td>
<td>Complement activation, hemolysis</td>
<td>Viral antibodies</td>
</tr>
<tr>
<td>Immunofluorescence (eg, indirect fluorescent antibody [IFA])</td>
<td>Fluorescent-tagged antibody reacts with antigen-antibody complex in the presence of ultraviolet light.</td>
<td>Visible microscopic fluorescence</td>
<td>Antinuclear antibodies (ANAs); antimitochondrial antibodies (AMAs)</td>
</tr>
<tr>
<td>Enzyme immunoassay (EIA)</td>
<td>Enzymes are used to label induced antigen-antibody reactions.</td>
<td>Chromogenic fluorescent Hepatitis and human or luminescent change in immunodeficiency virus (HIV) (screening)</td>
<td></td>
</tr>
<tr>
<td>Enzyme-linked immunosorbent assay (ELISA)</td>
<td>Indirect EIA for quantification of an antigen or antibody enzyme and substrate</td>
<td>Color change indicates enzyme substrate reaction.</td>
<td>Lyme disease, Epstein-Barr virus, extractable nuclear antibodies (connective tissue/systemic rheumatic disease)</td>
</tr>
<tr>
<td>Immunoblot (eg, Western blot [WB])</td>
<td>Electrophoresis separation of antigen subspecies</td>
<td>Detection of antibodies of specific mobility</td>
<td>Confirms HIV-1</td>
</tr>
<tr>
<td>Polymerase chain reaction (PCR)</td>
<td>Amplifies low levels of specific DNA sequences; each cycle doubles the amount of specific DNA sequence.</td>
<td>Exponential accumulation of DNA fragment being amplified; defects in DNA appear as mutations</td>
<td>Slightest trace of infection can be detected; more accurate than traditional tests for chlamydia; genetic disorders</td>
</tr>
<tr>
<td>Rate nephelometry</td>
<td>Measures either antigen or antibody in solution through the scattering of a light beam; antibody reagent used to detect antigen IgA, IgG, IgM; concurrent controls are run to establish amount of background scatter in reagents and test samples.</td>
<td>Light scatter proportionately increases as numbered size of immune complexes increases.</td>
<td>Quantitative immunoglobulins IgA, IgM, C-reactive protein, anti-streptolysin O recorded in mg/dL or IU/mL</td>
</tr>
<tr>
<td>Flow cytometry</td>
<td>Blood cell types are identified with monoclonal antibodies (mAbs) specific for cell markers by means of a flow cytometer with an argon laser beam; as the cells pass the beam, they scatter the light; light energy is converted into electrical energy cells and stained with green (fluorescence) or orange (phytoerythrin).</td>
<td>Light scatter identifies cell size and granularity of lymphocytes, monocytes, and granulocytes; color fluorochromes tagged to monoclonal antibodies bend to specific surface antigens for simultaneous detection of lymphocyte subsets.</td>
<td>Lymphocyte immunophenocytology differentiates B cells from T cells and T-helper cells from T-suppressor cells Epidemiology of nosocomial and community-acquired infections Infectious disease such as tuberculosis, hepatitis C virus, and HIV</td>
</tr>
<tr>
<td>Restriction fragment length polymorphism (RFLP)</td>
<td>DNA-based typing technique</td>
<td>Amplifies nucleic acid to identify presence of bacterial or viral load</td>
<td></td>
</tr>
<tr>
<td>cDNA probes</td>
<td>Uses cDNA probes directed against ribosomal RNA</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Collection of Serum for Immunologic Tests**

Specific antibodies can be detected in serum and other body fluids (eg, synovial fluid, cerebrospinal fluid [CSF]).

1. **Procure samples.** For diagnosis of infectious disease, a blood sample (serum preferred) using a 7-mL red-topped tube should be obtained at illness onset (acute phase), and the other sample should be drawn 3 to 4 weeks later (convalescent phase). In general, serologic test usefulness depends on a titer increase in the time interval between the acute and the convalescent phase. For some serologic tests, one serum sample may be adequate if the antibody presence indicates an abnormal condition or the antibody titer is unusually high. See [Appendix A](#) for standard precautions and [Appendix B](#) for latex precautions.

2. **Perform the serologic test before doing skin testing.** Skin testing often induces antibody production and could interfere with serologic test results.

3. **Label the sample properly and submit requested information.** Place specimen in biohazard bag. Send samples to the laboratory promptly. Hemolyzed samples cannot yield accurate results. Hemoglobin in the serum sample can interfere with complement-fixing antibody values.
Interpreting Results of Immunologic Tests

The following factors affect test results:

1. History of previous infection by the same organism
2. Previous vaccination (determine time frame)
3. Anamnestic reactions caused by heterologous antigens: an anamnestic reaction is the appearance of antibodies in the blood after administration of an antigen to which the patient has previously developed a primary immune response
4. Cross-reactivity: antibodies produced by one species of an organism can react with an entirely different species (e.g., Tularemia antibodies may agglutinate Brucella and vice versa, rickettsial infections may produce antibodies reactive with Proteus OX19)
5. Presence of other serious illness states (e.g., lack of immunologic response in agammaglobulinemia, cancer treatment with immunosuppressant drugs)
6. Seroconversion: the detection of specific antibody in the serum of an individual when this antibody was previously undetectable

Serologic Versus Microbiologic Methods

Serologic testing for microbial immunology evaluates the presence of antibodies produced by antigens of bacteria, viruses, fungi, and parasites. The best means of establishing infectious disease etiology is by isolation and confirmation of the involved pathogen. Serologic methods can assist or confirm microbiologic analysis when the patient is tested late in the disease course, antimicrobial therapy has suppressed organism growth, or culture methods cannot verify a causative agent.

BACTERIAL TESTS

Syphilis Detection Tests

Syphilis is a venereal disease caused by Treponema pallidum, a spirochete with closely wound coils approximately 8 to 15 µm long. Untreated, the disease progresses through three stages that can extend over many years.

Antibodies to syphilis begin to appear in the blood 4 to 6 weeks after infection (Table 8.2). Nontreponemal tests determine the presence of reagin, which is a nontreponemal autoantibody directed against cardiolipin antigens. These tests include rapid plasma reagin (RPR) and Venereal Disease Research Laboratory (VDRL). The U.S. Centers for Disease Control and Prevention (CDC) recommend these tests for syphilis screening; however, they may show negative results in some cases of late syphilis. Biologic false-positive results can also occur (Table 8.3).

Table 8.2 Sensitivity of Commonly Used Serologic Tests for Syphilis

<table>
<thead>
<tr>
<th>Test</th>
<th>Stage</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Primary (%)</td>
</tr>
<tr>
<td>Venereal Disease Research Laboratory test (VDRL)</td>
<td>70</td>
</tr>
<tr>
<td>Rapid plasma reagin card test (RPR); automated reagin test (ART)</td>
<td>80</td>
</tr>
<tr>
<td>SPECIFIC TREPONEMAL TESTS</td>
<td></td>
</tr>
<tr>
<td>Fluorescent treponemal antibody absorption test (FTA-ABS)</td>
<td>85</td>
</tr>
<tr>
<td>Treponema pallidum particle agglutination (TP-PA)</td>
<td>65</td>
</tr>
</tbody>
</table>

*Treated late syphilis.


Table 8.3 Nonsyphilitic Conditions Giving Biologic False-Positive Results (BFPs) Using VDRL and RPR Tests

<table>
<thead>
<tr>
<th>Disease</th>
<th>Approximate Percentage BFPs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Malaria</td>
<td>100</td>
</tr>
<tr>
<td>Leprosy</td>
<td>60</td>
</tr>
<tr>
<td>Relapsing fever</td>
<td>30</td>
</tr>
<tr>
<td>Active immunization in children</td>
<td>20</td>
</tr>
<tr>
<td>Infectious mononucleosis</td>
<td>20</td>
</tr>
<tr>
<td>Lupus erythematosus</td>
<td>20</td>
</tr>
<tr>
<td>Lymphogranuloma venereum</td>
<td>20</td>
</tr>
<tr>
<td>Pneumonia, atypical</td>
<td>20</td>
</tr>
<tr>
<td>Rat-bite fever</td>
<td>20</td>
</tr>
<tr>
<td>Condition</td>
<td>Score</td>
</tr>
<tr>
<td>-----------------------------------------------</td>
<td>-------</td>
</tr>
<tr>
<td>Typhus fever</td>
<td>20</td>
</tr>
<tr>
<td>Vaccinia</td>
<td>20</td>
</tr>
<tr>
<td>Infectious hepatitis</td>
<td>10</td>
</tr>
<tr>
<td>Leptospirosis (Weil's disease)</td>
<td>10</td>
</tr>
<tr>
<td>Periarteritis nodosa</td>
<td>10</td>
</tr>
<tr>
<td>Trypanosomiasis</td>
<td>10</td>
</tr>
<tr>
<td>Chancroid</td>
<td>5</td>
</tr>
<tr>
<td>Chickenpox</td>
<td>5</td>
</tr>
<tr>
<td>Measles</td>
<td>5</td>
</tr>
<tr>
<td>Rheumatoid arthritis</td>
<td>5–7</td>
</tr>
<tr>
<td>Rheumatic fever</td>
<td>5–6</td>
</tr>
<tr>
<td>Scarlet fever</td>
<td>5</td>
</tr>
<tr>
<td>Subacute bacterial endocarditis</td>
<td>5</td>
</tr>
<tr>
<td>Pneumonia, pneumococcal</td>
<td>3–5</td>
</tr>
<tr>
<td>Tuberculosis, advanced pulmonary</td>
<td>3–5</td>
</tr>
<tr>
<td>Blood loss, repeated</td>
<td>? (low)</td>
</tr>
<tr>
<td>Common cold</td>
<td>? (low)</td>
</tr>
<tr>
<td>Pregnancy</td>
<td>? (low)</td>
</tr>
</tbody>
</table>

Conversely, treponemal (ie, specific) tests detect antibodies to *T. pallidum*. These tests include the particle agglutination *T. pallidum* test (TP-PA) and the fluorescent treponemal antibody test (FTA-ABS). These tests confirm syphilis when a positive nontreponemal test result is obtained. Because these tests are more complex, they are not used for screening. Certain states require automatic confirmation for all reactive screening tests by using a treponemal test such as the TP-PA or FTA-ABS.

**Reference Values**

**Normal** Nonreactive negative for syphilis

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Fasting is usually not required.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**

1. Diagnosis of syphilis requires correlation of patient history, physical findings, and results of syphilis antibody tests. *T. pallidum* is diagnosed when both the screening and the confirmatory tests are reactive.
2. Treatment of syphilis may alter both the clinical course and the serologic pattern of the disease. Treatment related to tests that measure reagin (RPR and VDRL) includes the following measures:
   a. If the patient is treated at the seronegative primary stage (eg, after the appearance of the syphilitic chancre but before the appearance of reaction or reagin), the VDRL remains nonreactive.
   b. If the patient is treated in the seropositive primary stage (eg, after the appearance of a reaction), the VDRL usually becomes nonreactive within 6 months of treatment.
   c. If the patient is treated during the secondary stage, the VDRL usually becomes nonreactive within 12 to 18 months.
   d. If the patient is treated >10 years after the disease onset, the VDRL usually remains unchanged.
3. A negative serologic test may indicate one of the following circumstances:
   a. The patient does not have syphilis.
   b. The infection is too recent for antibodies to be produced. Repeat tests should be performed at 1-week, 1-month, and 3-month intervals to establish the presence or absence of disease.
   c. The syphilis is in a latent or inactive phase.
   d. The patient has a faulty immunodefense mechanism.
   e. Laboratory techniques were faulty.

**False-Positive and False-Negative Reactions** A positive reaction is not conclusive for syphilis. Several conditions produce biologic false-positive results for syphilis. Biologic false-positive reactions are by no means “false.” They may reveal the presence of other serious diseases. It is theorized that reagin (reaction) is an antibody against tissue lipids. Lipids are presumed to be liberated from body tissue in the normal course of activity. These liberated lipids may then induce antibody formation. Nontreponemal biologic false-positive reactions can occur in the presence of drug abuse, lupus erythematosus, mononucleosis, malaria, leprosy, viral pneumonia, recent immunization, or, on rare occasions, pregnancy. False-negative reactions may occur early in the disease course or during inactive or later stages of disease.

**Interfering Factors**

1. Excess chyle in the blood interferes with test results.
2. Alcohol decreases reaction intensity in tests that detect reagin; therefore, alcohol ingestion should be avoided for at least 24 hours before blood is drawn.

**Clinical Alert**

Avoid drawing the blood sample immediately after a meal.
Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure. Assess for interfering factors. Instruct the patient to abstain from alcohol for at least 24 hours before the blood sample is drawn.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and counsel appropriately. Explain biologic false-positive or false-negative reactions. Advise that repeat testing may be necessary.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

1. Sexual partners of patients with syphilis should be evaluated for the disease.
2. After treatment, patients with early-stage syphilis should be tested at 3-month intervals for 1 year to monitor for declining reactivity.

Lyme Disease Tests

Lyme disease is a multisystem disorder caused by the spirochete *Borrelia burgdorferi*. It is transmitted by the bite of tiny deer ticks, which reside on deer and other wild animals. Lyme disease is present worldwide, but certain geographic areas show higher incidences. Transmission to humans is highest during the spring, summer, and early fall months. The tick bite usually produces a characteristic rash, termed *erythema chronicum migrans*. If untreated, sequelae lead to serious joint, cardiac, and central nervous system (CNS) symptoms.

Serologic testing for antibodies to Lyme disease includes enzyme-linked immunosorbent assay (ELISA) and Western blot analysis. Antibody formation takes place in the following manner: immunoglobulin M (IgM) is detected 3 to 4 weeks after Lyme disease onset, peaks at 6 to 8 weeks after onset, and then gradually disappears. IgG is detected 2 to 3 months after infection and may remain elevated for years. Current CDC recommendations for the serologic diagnosis of Lyme disease are to screen with a polyvalent ELISA (IgG and IgM) and to perform supplemental testing (Western blot) on all equivocal and/or positive ELISA results.

Western blot assays for antibodies to *B. burgdorferi* are supplemental rather than confirmatory because their specificity is less than optimal, particularly for detecting IgM. Two-step positive results provide supportive evidence of exposure to *B. burgdorferi*, which could support a clinical diagnosis of Lyme disease but should not be used as a criterion for diagnosis.

Reference Values

Normal

Negative for both IgG and IgM Lyme antibodies by ELISA and Western blot

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Cerebrospinal fluid may also be used for the test.
2. Observe standard precautions.
3. Place specimen in a biohazard bag.

Clinical Implications

1. Serologic tests lack the degree of sensitivity, specificity, and standardization necessary for diagnosis in the absence of clinical history. The antigen detection assay for bacterial proteins is of limited value in early stages of disease.
2. In patients presenting with a clinical picture of Lyme disease, negative serologic tests are inconclusive during the first month of infection.
3. Repeat paired testing should be performed if borderline values are reported.
4. The CDC states that the best clinical marker for Lyme disease is the initial skin lesion erythema migrans (EM), which occurs in 60% to 80% of patients.
5. CDC laboratory criteria for the diagnosis of Lyme disease include the following factors:
   a. Isolation of *B. burgdorferi* from a clinical specimen
   b. IgM and IgG antibodies in blood or CSF
   c. Paired acute and convalescent blood samples showing significant antibody response to *B. burgdorferi*

Interfering Factors

1. False-positive results may occur with high levels of rheumatoid factors or in the presence of other spirochete infections, such as syphilis (cross-reactivity).
2. Asymptomatic individuals who spend time in endemic areas may have already produced antibodies to *B. burgdorferi*.

Interventions

Pretest Patient Preparation

1. Assess patient's clinical history, exposure risk, and knowledge regarding the test. Explain test purpose and procedure as well as possible follow-up testing.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes for a positive test. Advise that follow-up testing may be required to monitor response to antibiotic therapy.
2. Unlike other diseases, people do not develop resistance to Lyme disease after infection and may continue to be at high risk, especially if they live, work, or recreate in areas where Lyme disease is present.
3. If Lyme disease has been ruled out, further testing may include *Babesia microti*, a parasite transmitted to humans by a tick bite. Symptoms include loss of appetite, fever, sweats, muscle pain, nausea, vomiting, and headaches.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Legionnaire’s Disease Antibody Test**

Legionnaire’s disease is a respiratory condition caused by *Legionella pneumophila*. It is best diagnosed by organism culture; however, the organism is difficult to grow.

Detection of *L. pneumophila* in respiratory specimens by means of direct fluorescent antibody (DFA) technique is useful for rapid diagnosis but lacks sensitivity when only small numbers of organisms are available. Serologic tests should be used only if specimens for culture are not available or if culture and DFA produce negative results.

**Reference Values**

**Normal** Negative for Legionnaire’s disease by indirect fluorescent antibody (IFA) test or ELISA

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in a biohazard bag for transport to the laboratory.
2. Be aware that follow-up testing is usually requested 3 to 6 weeks after initial symptom appearance.
3. Alert patient that a urine specimen may be required if antigen testing is indicated.

**Clinical Implications**

1. A dramatic rise of titer to levels to more than 1:128 in the interval between acute- and convalescent-phase specimens occurs with recent infections.
2. Serologic tests, to be useful, must be performed on an acute (within 1 week of onset) and convalescent (3 to 6 weeks later) specimen.
3. Serologic testing is valuable because it provides a confirmatory diagnosis of *L. pneumophila* infection when other tests have failed. IFA is the serologic test of choice because it can detect all classes of antibodies.
4. Demonstration of *L. pneumophila* antigen in urine by ELISA is indicative of infection.

**Interventions**

**Pretest Patient Preparation**

1. Assess clinical history and knowledge about the test. Explain purpose and procedure of blood test.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and significance. Advise that negative results do not rule out *L. pneumophila*. Follow-up testing is usually needed.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Chlamydia Antibody IgG Test**

Chlamydia is caused by a genus of bacteria (*Chlamydia* spp.) that require living cells for growth and are classified as obligate cell parasites. Recognized species include *Chlamydia psittaci*, *Chlamydia pneumoniae*, and *Chlamydia trachomatis*. *C. psittaci* causes psittacosis in birds and humans. *C. pneumoniae* is responsible for approximately 10% of cases of community-acquired pneumonia. *C. trachomatis* is grouped into three serotypes. One group causes lymphogranuloma venereum (LGV), a venereal disease. Another group causes trachoma, an eye disease. The third group causes genital tract infections different from LGV. Culture of the organism is definitive for chlamydiae. *C. trachomatis* infection is the most common reportable sexually transmitted disease (STD) in the United States. The national infection rate for *C. trachomatis* is estimated to be 3 million cases annually.

Because *Chlamydia* organisms are difficult to culture and grow, antibody testing aids in diagnosis of chlamydial infection.

**Reference Values**

**Normal** Negative for chlamydia antibody by complement fixation (CF), IFA, and PCR tests

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**
1. Presence of antibody titer indicates past chlamydial infection. A fourfold or greater rise in antibody titer between acute and convalescent specimens indicates recent infection. Serologic tests cannot differentiate among the species of *Chlamydia*.

2. Infection with psittacosis is revealed in an elevated antibody titer. History will reveal contact with infected birds (pets or poultry).

3. LGV in males is characterized by swollen and tender inguinal lymph nodes. In females, swelling occurs in the intraabdominal, perirectal, and pelvic lymph nodes. *Chlamydia* causes urethritis in males. It can infect the female urethra and endocervix, and it is also a cause of pelvic inflammatory disease in females. Eye disease caused by *Chlamydia* is endemic in parts of Africa, the Middle East, and Southeast Asia, although its presence is established worldwide. Culture and stained smear identification of the organism is diagnostic.

**Interfering Factors** Depending on geographic location, nonspecific titers can be found in the general healthy population.

**Interventions**

**Pretest Patient Preparation**

1. Assess patient knowledge regarding the test and explain purpose and procedure. Elicit history regarding possible exposure to organism.

2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and significance of test results. Refer to page 532 for interpretation of immunodiagnostic test results.

2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Streptococcal Antibody Tests: Anti-Streptolysin O Titer (ASO), Streptozyme, Anti-DNase B (ADB, Streptodornase)**

Group A β-hemolytic streptococci are associated with streptococcal infections or illness.

These tests detect antibodies to enzymes produced by organisms. Group A β-hemolytic streptococci produce several enzymes, including streptolysin O, hyaluronidase, and DNase B. Serologic tests that detect these enzyme antibodies include antistreptolysin O titer (ASO), which detects streptolysin O; streptozyme, which detects antibodies to multiple enzymes; and anti-DNase B (ADB), which detects DNase B. Serologic detection of streptococcal antibodies helps to establish prior infection but is of no value for diagnosing acute streptococcal infections. Acute infections should be diagnosed by direct streptococcal cultures or the presence of streptococcal antigens.

The ASO test aids in the diagnosis of several conditions associated with streptococcal infections, such as rheumatic fever, glomerulonephritis, endocarditis, and scarlet fever. Serial rising titers over several weeks are more significant than a single result. ADB antibodies may appear earlier than ASO in streptococcal pharyngitis, and this test is more sensitive for streptococcal pyoderma.

**Reference Values**

**Normal**

ASO titer: <166 Todd units (or <200 IU) Anti-DNase B (ADB) Birth–4 years: <170 U 5–19 years: <480 U >20 years: <340 U Streptozyme: negative for streptococcal antibodies

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in a biohazard bag for transport to the laboratory.

2. Repeat testing 10 days after the first test is recommended.

**Clinical Implications**

1. In general, a titer >166 Todd units is considered a definite elevation for the ASO test.

2. The ASO or the ADB test alone is positive in 80% to 85% of group A streptococcal infections (eg, streptococcal pharyngitis, rheumatic fever, pyoderma, glomerulonephritis).

3. When ASO and ADB tests are run concurrently, 95% of streptococcal infections can be detected.

4. A repeatedly low titer is good evidence for the absence of active rheumatic fever. Conversely, a high titer does not necessarily mean rheumatic fever of glomerulonephritis is present; however, it does indicate the presence of a streptococcal infection.

5. ASO production is especially high in rheumatic fever and glomerulonephritis. These conditions show marked ASO titer increases during the symptomless period preceding an attack. Also, ADB titers are particularly high in pyoderma.

**Interfering Factors**

1. An increased titer can occur in healthy carriers.

2. Antibiotic therapy suppresses streptococcal antibody response.

3. Increased B-lipoprotein levels inhibit streptolysin O and produce falsely high ASO titers.
Clinical Alert

The ASO test is impractical in patients who have recently received antibiotics or who are scheduled for antibiotic therapy because the treatment suppresses the antibody response.

Interventions

Pretest Patient Preparation

1. Assess patient's clinical history and test knowledge. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes. Refer to page 532 for interpretation of immunologic test results. Explain test results. Inform patient that repeat testing is frequently required.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Helicobacter pylori (HPY) IgG Antibody Serum, Stool, and Breath (PY) Test

*H. pylori* (previously known as *campylobacter pylori*) is a bacterium associated with gastritis, duodenal and gastric ulcers, and possibly gastric carcinoma. The clinician orders this test when screening a patient for possible *H. pylori* infection. The organism is present in 95% to 98% of patients with duodenal ulcers and 60% to 90% of patients with gastric ulcers. A person with gastrointestinal symptoms with evidence of *H. pylori* colonization (eg, presence of specific antibodies, positive breath test, positive culture, positive biopsy) is considered to be infected with *H. pylori*. A person without gastrointestinal symptoms having evidence of the presence of *H. pylori* is said to be colonized rather than infected.

This test detects *H. pylori* infection of the stomach. Traditionally, the presence of *H. pylori* has been detected through biopsy specimens obtained by endoscopy. As with any invasive procedure, there is risk and discomfort to the patient. Noninvasive methods of detection include the following:

1. An *H. pylori* breath test, which uses a liquid scintillation counter to detect gastric urease and to measure $^{14}$CO$_2$ in breath specimens
2. Serology—blood serum
3. Stool: *H. pylori* stool antigen test (HpSa)

The presence of *H. pylori*–specific IgG antibodies has been shown to be an accurate indicator of *H. pylori* colonization. ELISA testing relies on the presence of *H. pylori* IgG-specific antibody to bind to antigen on the solid phase, forming an antigen-antibody complex that undergoes further reactions to produce a color indicative of the presence of antibody and is quantified using a spectrophotometer or ELISA microwell plate reader. The sensitivity is 94% and specificity 78%, compared with an invasive procedure, such as biopsy, for which the sensitivity is 93% and specificity 99%.

Reference Values

**Normal** Negative for *H. pylori* by ELISA indicates no detectable IgG antibody in serum or stool. A positive result indicates the presence of detectable IgG antibody in serum or stool. **Breath**: Negative: <50 DPM for *H. pylori* 50–199 DPM indeterminate for *H. pylori* >200 DPM positive for *H. pylori*

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in a biohazard bag for transport to the laboratory.
2. Be aware that a random stool specimen may be ordered to test for the presence of *H. pylori* antigen.
3. Remember that this is a complex procedure. Use a special kit for the breath test. Ensure that the collection balloon is fully inflated. Transfer the breath specimen to the laboratory. Keep at room temperature.

Clinical Implications

1. This assay is intended for use as an aid in the diagnosis of *H. pylori*, and additionally, false-negative results may occur. The clinical diagnosis should not be based on serology alone but rather on a combination of serology (and breath or stool tests), symptoms, and gastric biopsy–based tests as warranted.
2. The stool antigen test is used to monitor response during therapy and to test for cure after treatment.

Education Alert for Breath Test

1. The patient should have no antibiotics and bismuth for 1 month and no proton pump inhibitors and sucralfate for 2 weeks before test.
2. Instruct the patient not to chew the capsule.
3. The patient should be at rest during breath collection.

Interventions

**Pretest Patient Preparation**

1. Explain test purpose, procedure, and knowledge of signs and symptoms and risk factors for transmission: close living quarters, many persons in household, poor household sanitation and hygiene. The patient swallows a capsule before a breath specimen is obtained. The serum antibody test would be appropriate for a previously untreated patient with a documented history of gastroduodenal ulcer disease and unknown *H. pylori* infection.
status.

2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes in light of patient's history, including other clinical and laboratory findings. Explain treatment (4–6 weeks of antibiotics to eradicate *H. pylori* and medication to suppress acid production) and need for follow-up testing. Transmission is unknown but potential for transmission may occur during episodes of GI illness, particularly with vomiting. Many persons may be infected with *H. pylori* but are asymptomatic.

2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

VIRAL TESTS

**Epstein-Barr Virus (EBV) Antibody Tests: Infectious Mononucleosis (IM) Slide (Screening) Test, Heterophile Antibody Titer, Epstein-Barr Antibodies to Viral Capsid Antigen and Nuclear Antigen**

Epstein-Barr virus (EBV) is a herpesvirus found throughout the world. The most common symptomatic manifestation of EBV infection is a disease known as *infectious mononucleosis* (IM). This disease induces formation of increased numbers of abnormal lymphocytes in the lymph nodes and stimulates increased heterophile antibody formation. IM occurs most often in young adults who have not been previously infected, through contact with infectious oropharyngeal secretions. Symptoms include fever, pharyngitis, and lymphadenopathy. EBV is also thought to play a role in the etiology of Burkitt's lymphoma, nasopharyngeal carcinoma, and chronic fatigue syndrome.

The most common test for EBV is the rapid slide test (Monospot) for heterophile antibody agglutination. The heterophile antibody agglutination test is not specific for EBV and therefore is not useful for evaluating chronic disease. If the heterophile test is negative in the presence of acute IM symptoms, specific EBV antibodies should be determined. These include antibodies to viral capsid antigen (anti-VCA) and antibodies to EBV nuclear antigen (EBNA) using IFA and ELISA tests.

Diagnosis of IM is based on the following criteria: clinical features compatible with IM, hematologic picture of relative and absolute lymphocytosis, and presence of heterophile antibodies.

**Reference Values**

**Normal** Negative for IM and EBV antibodies by latex agglutination (IM) and IFA or ELISA (EBV)

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**

1. The presence of heterophile antibodies (Monospot), along with clinical signs and other hematologic findings, is diagnostic for IM.
2. Heterophile antibodies remain elevated for 8 to 12 weeks after symptoms appear.
3. Approximately 90% of adults have antibodies to the virus.
4. The Monospot test is negative more frequently in children and almost uniformly in infants with primary EBV infection.

**Interventions**

**Pretest Patient Preparation**

1. Assess patient's clinical history, symptoms, and test knowledge. Explain test purpose and procedure. If preliminary tests are negative, follow-up tests may be necessary.

2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes. Refer to page 532 for interpretation of immunologic test results. Explain treatment (eg, supportive therapy [IV fluids]). After primary exposure, a person is considered immune. Recurrence of IM is rare.
2. Remember that resolution of IM usually follows a predictable course: pharyngitis disappears within 14 days after onset; fever subsides within 21 days; and fatigue, lymphadenopathy, and liver and spleen enlargement regress by 21 to 28 days.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Hepatitis Tests: Hepatitis A (HAV), Hepatitis B (HBV), Hepatitis C (HCV), Hepatitis D (HDV), Hepatitis E (HEV), Hepatitis G (HGV)**

Hepatitis can be caused by viruses and several other agents, including drugs and toxins. Approximately 95% of hepatitis cases are due to five major virus types: hepatitis A, B, C, D, and E (Table 8.4). Diagnosing the specific virus is difficult because the symptoms (eg, chills, weight loss, fever, distaste for cigarettes and food, darker urine and lighter stool) presented by each viral type are similar. Additionally, some individuals may be asymptomatic or have very mild symptoms that are ascribed to the “flu.” Serologic tests for hepatitis virus markers have made it easier to define the specific type.
**Table 8.4 Hepatitis Test Findings in Various Stages**

<table>
<thead>
<tr>
<th>Disease Stages</th>
<th>Viral Specific and Serologic Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HAV</td>
</tr>
<tr>
<td>Acute</td>
<td>IgM anti-HAV</td>
</tr>
<tr>
<td>Chronic</td>
<td>Fecal HAV 1–2 wk before symptoms</td>
</tr>
<tr>
<td>Infection</td>
<td>None</td>
</tr>
<tr>
<td>Recovery</td>
<td>None</td>
</tr>
<tr>
<td>Viral load (viral genome)</td>
<td>None</td>
</tr>
<tr>
<td>Carrier state</td>
<td>None</td>
</tr>
<tr>
<td>Immunity</td>
<td>None</td>
</tr>
<tr>
<td>Acute viral panel</td>
<td>IgM anti-HEV</td>
</tr>
</tbody>
</table>

**Hepatitis A virus** (HAV), which is acquired through enteric transmission, infects the gastrointestinal tract and is eliminated through the feces. Serologically, the presence of the IgM antibody to HAV (IgM anti-HAV) and the total antibody to HAV (total anti-HAV) identifies the disease and determines previous exposure or recovery from HAV.

**Hepatitis B virus** (HBV) demonstrates a central core containing the core antigen and a surrounding envelope containing the surface antigen: less than 0.01 pg/mL for viral load. Detection of core antigen (HBcAg), envelope antigen (HBeAg), and surface antigen (HBsAg) or their corresponding antibodies constitutes hepatitis B serologic or plasma assessment. Viral transmission occurs through exposure to contaminated blood or blood products through an open wound (eg, needle sticks, lacerations). Hepatitis monitoring panel for serial testing includes four B markers: HBsAg, HBeAg, anti-HBe, and anti-HBs. Interpretation depends on clinical setting. Hepatitis B DNA Ultra Sensitive Quantitative PCR is the most sensitive test available for hepatitis B viral load.

**Hepatitis C virus** (HCV), formerly known as non-A, non-B hepatitis, is also transmitted parenterally. HCV infection is characterized by presence of antibodies to hepatitis C (anti-HCV) and levels of alanine aminotransferase (ALT), which fluctuate between normal and markedly elevated. Levels of anti-HCV remain positive for many years; therefore, a reactive test indicates infection with HCV or a carrier state but not infectivity or immunity. Polymerase chain reaction (PCR) or reverse transcriptase PCR (RT-PCR) (viral load), which detects HCV RNA, should be used to confirm infection when acute hepatitis C is suspected. A negative hepatitis C antibody (recombinant immunoblot assay [RIBA]) does not exclude the possibility of HCV infection because seroconversion may not occur for up to 6 months after exposure.

**Hepatitis D virus** (HDV) is encapsulated by the HBsAg. Without the HBsAg coating, HDV cannot survive. Because HDV can cause infection only in the presence of active HBV infection, it is usually found where a high incidence of HBV occurs. Transmission is parenteral. Serologic HDV determination is made by detection of the hepatitis D antigen (HDAg) early in the course of the infection and by detection of anti-HDV antibody (anti-HDV) in the later stages of the disease.

**Hepatitis E virus** (HEV) is transmitted enterically and is associated with poor hygienic practices and unsafe water supplies, especially in developing countries. It is quite rare in the United States. Specific serologic tests include detection of IgM and IgG antibodies to hepatitis E (anti-HEV).

**Hepatitis G virus** (HGV) is transmitted by contaminated blood supply and is seen when HCV and HBV are detected together. See Table 8.5 for a summary of the features of the different hepatitis agents.

**Table 8.5 Summary of Clinical and Epidemiologic Features of Viral Hepatitis Agents**

<table>
<thead>
<tr>
<th>Features</th>
<th>Hepatitis A</th>
<th>Hepatitis B</th>
<th>Hepatitis C</th>
<th>Hepatitis D</th>
<th>Hepatitis E</th>
<th>Hepatitis G scan with HBV + HCV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incubation period</td>
<td>45–50 d</td>
<td>30–150 d</td>
<td>15–110 d</td>
<td>30–150 d</td>
<td>230–240 d</td>
<td>Questionable</td>
</tr>
<tr>
<td>Onset Jaundice</td>
<td>Abrupt</td>
<td>Insidious</td>
<td>Insidious</td>
<td>Abrupt</td>
<td>Insidious</td>
<td>Unknown</td>
</tr>
<tr>
<td>Jaundice</td>
<td>Children: 10%; adults: 70%–80%</td>
<td>25%</td>
<td>25%</td>
<td>Varies</td>
<td>Unknown</td>
<td>No</td>
</tr>
<tr>
<td>Asymptomatic patients</td>
<td>Most children; adults: 50%</td>
<td>Most children; adults: 75%</td>
<td>Rare</td>
<td>Rare</td>
<td>Contaminated blood</td>
<td></td>
</tr>
<tr>
<td>Routes of transmission</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mode of Transmission</td>
<td>Yes</td>
<td>No</td>
<td>Possible</td>
<td>Possible</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>----------------------</td>
<td>-----</td>
<td>----</td>
<td>----------</td>
<td>----------</td>
<td>-----</td>
<td>----</td>
</tr>
<tr>
<td>Fecal/oral</td>
<td>Yes</td>
<td>No</td>
<td></td>
<td></td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Parenteral</td>
<td>Rare</td>
<td>Yes</td>
<td></td>
<td></td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Sexual</td>
<td>No</td>
<td>Yes</td>
<td></td>
<td></td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Perinatal</td>
<td>No</td>
<td>Yes</td>
<td></td>
<td></td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Water/food</td>
<td>Yes</td>
<td>No</td>
<td></td>
<td></td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Chronic state</td>
<td>No</td>
<td>No</td>
<td></td>
<td></td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Case fatality rate</td>
<td>0.6%</td>
<td>1.4%</td>
<td></td>
<td>1%–2%</td>
<td>30%</td>
<td></td>
</tr>
</tbody>
</table>

The following terms are used:

**ALT (alanine aminotransferase):** an enzyme normally produced by the liver; blood levels may increase in cases of liver damage.

**Anti-HBc:** antibody to hepatitis B core antigen

**Anti-HBe:** antibody to hepatitis B envelope antigen

**Anti-HBs:** antibody to hepatitis B surface antigen

**Antibody:** a Y-shaped protein molecule (immunoglobulin) in serum or body fluid that either neutralizes an antigen or tags it for attack by other cells or chemicals; acts by uniting with and firmly binding to an antigen. The prefix *anti-* followed by initials of a virus refers to specific antibody against the virus.

**Chronic hepatitis:** a condition in which symptoms and/or signs of hepatitis persist for >6 months

**Cirrhosis:** irreversible scarring of the liver that may occur after acute or chronic hepatitis

**Delta agent:** a unique RNA virus that causes acute or chronic hepatitis; requires HBV for replication and infects only patients who are HBsAg-positive; is composed of a delta antigen core and a HBsAg coat; also known as HDV

**Endemic:** present in a community at all times but occurring in a small number of cases

**Enteric route:** spread of organisms through the oral-intestinal-fecal cycle

**Flavivirus:** a family of small RNA viruses; HCV is similar to members of the Flavivirus family.

**Fulminant hepatitis:** the most severe form of hepatitis; may lead to acute liver failure and death

**HBcAg:** hepatitis B core antigen

**HBsAg:** hepatitis B surface antigen

**Hepatotropic:** having an affinity for or exerting a specific effect on the liver

**IgG:** a form of immunoglobulin that occurs late in an infectious process

**IgM:** a form of immunoglobulin that occurs early in an infectious process

**IgM anti-HAV:** M-class immunoglobulin antibody to HAV

**IgM anti-HBc:** M-class immunoglobulin antibody to HBcAg

**Immune globulin:** a sterile solution of water-soluble proteins that contains those antibodies normally present in adult human blood; used as a passive immunizing agent against various viruses such as HAV

**Negative-sense RNA virus:** a virus in which the viral proteins are encoded by messenger RNA molecules that are complementary to the viral genome

**Non-A, non-B hepatitis:** viral hepatitis caused by viruses other than A, B, or D (eg, C, E)

**Parenteral:** entering the body subcutaneously, intramuscularly, or intravenously, or other means whereby the organisms reach the bloodstream directly

**Positive-sense RNA virus:** a virus in which the parenteral (or genomic) RNA serves as the messenger RNA for protein synthesis

**Recombinant antigen:** an antigen that results from the recombination of genetic components, which then are artificially
introduced into a cell, leading to synthesis of a new protein

**Viral load**: the amount or concentration of virus in the circulation

**New viruses**—GBV-A, GBV-B, and GBV-C: may be causative agents in non-A through E hepatitis

These measurements are used for differential diagnosis of viral hepatitis, viral load. Serodiagnosis of previous exposure and recovery of viral hepatitis is complex because of the number of serum or plasma markers necessary to determine the stage of illness. Testing methods include ELISA, microparticle enzyme immunoassay (MEIA), PCR, and RT-PCR and tests for viral genome (viral load).

**Indications for Hepatitis B Vaccine**

1. Family members of adoptees from foreign countries who are HBsAg positive
2. Health care workers (dentist, DO, MD, RN, and trainees in health care fields)
3. Hemodialysis patients or patients with early renal failure
4. Household or sexual contacts of persons chronically infected with hepatitis B
5. Immigrants from Africa or Southeast Asia; recommended for children <11 years old and all susceptible household contacts of persons chronically infected with hepatitis B
6. Injection drug users
7. Inmates of long-term correctional facilities
8. Clients and staff of institutions for the developmentally disabled
9. International travelers to countries of high or intermediate HBV endemicity
10. Laboratory workers
11. Public safety workers (eg, police, fire fighters)
12. Recipients of clotting factors. Use a fine needle (<23 gauge) and firm pressure at injection site for >2 minutes.
13. Persons with sexually transmitted diseases or multiple sexual partners in previous 6 months, prostitutes, homosexual and bisexual men
14. Postvaccination blood testing is recommended for sexual contacts of HBsAg-positive persons; health care workers at high risk, recipients of clotting factors, those who are HBsAg-positive.
15. Persons in nonresidential day care programs should be vaccinated if an HBsAg-positive classmate behaves aggressively or has special medical problems that increase the risk for exposure to blood. Staff in nonresidential day care programs should be vaccinated if a client is HBsAg-positive.
   a. Observe enteric and standard precautions for 7 days after onset of symptoms and/or jaundice with hepatitis B. Hepatitis A is most contagious before symptoms and/or jaundice appear.
   b. Use standard blood and body fluid precautions for type B hepatitis and B antigen carriers. Precautions apply until the patient is HBsAg negative and the anti-HBs appears. Avoid "sharps" (eg, needles, scalpel blades) injuries. Should accidental injury occur, encourage some bleeding, and wash area well with a germicidal soap. Report injury to proper department, and follow up with necessary interventions. Put on gown when blood splattering is anticipated. A private hospital room and bathroom may be indicated.
16. Persons with a history of receiving blood transfusion should not donate blood for 6 months. Transfusion-acquired hepatitis may not show up for 6 months after transfusion. Persons who test positive for HBsAg should never donate blood or plasma.
17. Persons who have sexual contact with hepatitis B–infected individuals run a greater risk for acquiring that same infection. HBsAg appears in most body fluids, including saliva, semen, and cervical secretions.
18. Observe standard precautions in all cases of suspected hepatitis until the diagnosis and hepatitis type are confirmed.

**Reference Values**

**Normal**

1. Negative (nonreactive) for hepatitis A, B, C, D, or E by ELISA, MEIA, PCR or RIBA, or RT-PCR
2. Negative or undetected viral load (not used for primary infection, only to monitor). PCR requires a separate specimen collection.
3. Hepatitis B viral DNA (HBV-DNA) negative or nonreactive viral load (<0.01 pg/mL) in an infected individual before treatment

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube or two lavender-topped ethylenediaminetetraacetic acid (EDTA) tubes, 5 mL each, for plasma. Observe standard precautions. Centrifuge promptly and aseptically. Place specimen in a biohazard bag for transport to the laboratory. Send specimens frozen on dry ice. Check with your laboratory for protocols and whether plasma or serum is needed.
2. Be aware that some specimens need to be split into two plastic vials before freezing and sent frozen on dry ice. Check with your laboratory.

**Clinical Implications**

1. Individuals with hepatitis may have generalized symptoms resembling the flu and may dismiss their illness as such.
2. A specific type of hepatitis cannot be differentiated by clinical observations alone. Testing is the only sure method to define the category.
3. Rapid diagnosis of acute hepatitis is essential for the patient so that treatment can be instituted and for those who have close patient contact so that protective measures can be taken to prevent disease spread.
4. Persons at higher risk for acquiring hepatitis A include patients and staff in health care and custodial institutions, people in day care centers, intravenous drug abusers, and those who travel to undeveloped countries or regions
where food and water supplies may be contaminated.

5. Persons at higher risk for hepatitis B include those with a history of drug abuse, those who have sexual contact with infected persons, and those who have household contact with infected persons and especially those with skin and mucosal surface lesions (eg, impetigo, saliva from chronic HBV persons on toothbrush racks and coffee cups in their homes); additionally, infants born to infected mothers (during delivery), hemodialysis patients, and health care employees are at higher risk for infection. Of all persons with HBV infection, 38% to 40% contract HBV during early childhood.

6. Health care workers should be periodically tested for hepatitis exposure and should always observe standard precautions when caring for patients.

7. Persons at risk for hepatitis C include those who have received blood transfusions, engage in intravenous drug abuse, undergo hemodialysis, have had organ transplantation, or have sexual contact with an infected person; hepatitis C can also be transmitted during delivery from mother to neonate. Most people are asymptomatic at the time of diagnosis for hepatitis C. See Table 8.6 for hepatitis markers that appear after infection.

### Table 8.6 Hepatitis Markers That Appear After Infection

<table>
<thead>
<tr>
<th>Serologic Marker</th>
<th>Time Marker</th>
<th>Clinical Implications</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HEPATITIS A VIRUS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HAV-Ab/IgM</td>
<td>4–6 wk</td>
<td>Positive for acute stage of hepatitis A, develops early in disease course</td>
</tr>
<tr>
<td>HAV-Ab/IgG</td>
<td>8–12 wk</td>
<td>Indicates previous exposure and immunity to hepatitis A</td>
</tr>
<tr>
<td><strong>HEPATITIS B VIRUS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HBsAg-hepatitis B virus</td>
<td>12 wk</td>
<td>Positive in acute stage of hepatitis B; earliest indicator of acute antigen infection; also indicates chronic infection</td>
</tr>
<tr>
<td>HBeAg</td>
<td>4–12 wk</td>
<td>Positive in acute active stage with viral replication (infectivity factor); highly infective</td>
</tr>
<tr>
<td>HBcAb</td>
<td>6–14 wk</td>
<td>This marker may remain in serum for a longer time; together with HBsAB represents convalescent stage; indicates past infection</td>
</tr>
<tr>
<td>HBcAb IgM</td>
<td>6–14 wk</td>
<td>Indicates acute infection</td>
</tr>
<tr>
<td>HBeAb antibody</td>
<td>8–16 wk</td>
<td>Indicates acute infection resolution</td>
</tr>
<tr>
<td>HBsAb antibody</td>
<td>4–10 mo</td>
<td>Indicates previous exposure, clinical recovery, immunity to hepatitis B, not necessarily to other types of hepatitis; marker for permanent immunity to hepatitis B</td>
</tr>
</tbody>
</table>

Interventions

**Pretest Patient Preparation**

1. Assess patient's social and clinical history and knowledge of test. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Explain significance of test results and counsel appropriately regarding presence of infection, recovery, and immunity. Counsel health care workers and family regarding protective and preventive measures necessary to avoid transmission. Instruct patient to alert health care workers and others regarding their hepatitis history in situations in which exposure to body fluids and wastes may occur.
2. Be aware that pregnant women may need special counseling.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Clinical Alert**

1. Observe enteric and standard precautions for 7 days after onset of symptoms and/or jaundice in hepatitis A. Hepatitis A is most contagious before symptoms and/or jaundice appear.
2. Use standard blood and body fluid precautions with hepatitis B and hepatitis B antigen carriers. Precautions apply until the patient is HBsAg negative and anti-HBs appears. Avoid “sharps” (eg, needles, scalpel blades) injuries. Should accidental injury occur, encourage some bleeding, and wash area well with a germicidal soap. Report injury to proper department and follow up with necessary interventions. Put on gown when blood splattering is anticipated. A private hospital room and bathroom may be indicated.
3. If patient has had a blood transfusion, he or she should not donate blood for 6 months. Transfusion-acquired hepatitis may not show up for 6 months after transfusion. Persons who test positive for HBsAg should never donate blood or plasma.
4. Persons who have sexual contact with hepatitis B–infected individuals run a greater risk for acquiring the infection. HBsAg appears in most body fluids, including saliva, semen, and cervical secretions.
5. Standard precautions must be observed in all cases of suspected hepatitis until the diagnosis and hepatitis type are confirmed.
6. Immunization of persons exposed to the infection should be done as soon as possible. In the case of contact with hepatitis B, both hepatitis B immunoglobulin (HBIG) and HBV vaccine should be administered within 24 hours of skin-break contact and within 14 days of last sexual contact. For hepatitis A, IG should be given within 2 weeks of exposure. In day care centers, immune globulin (IG) should be given to all contacts (children and personnel).
Differential Diagnosis of Viral Hepatitis

<table>
<thead>
<tr>
<th>Virus</th>
<th>Transmission</th>
<th>Incubation Period</th>
<th>Test for Acute Infection</th>
<th>Social and Clinical History</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hepatitis A</td>
<td>Fecal-oral by person-to-person contact or ingestion of contaminated food</td>
<td>Average, 30 d (range, 15–50 d)</td>
<td>IgM antibody to hepatitis A capsid proteins</td>
<td>Household or sexual contact with an infected person, day care centers, and common source outbreaks from contaminated food</td>
</tr>
<tr>
<td>Hepatitis B</td>
<td>Sexual, blood and other body fluids</td>
<td>Average, 120 d (range, 45–160 d)</td>
<td>the best test for acute or recent infection is IgM antibody to HBcAg</td>
<td>Sexual promiscuity, male-to-female sexual practices, injection drug use, birth to an infected mother</td>
</tr>
<tr>
<td>Hepatitis C</td>
<td>Blood</td>
<td>Commonly 6–9 wk (range, 2 wk–6 mo)</td>
<td>ELISA is the initial test to show if ever infected; it should be confirmed by another test such as PCR</td>
<td>Injection drug use, occupational exposure to blood, hemodialysis transfusion, possibly sexual transmission</td>
</tr>
<tr>
<td>Hepatitis D</td>
<td>Sexual, blood and other body fluids</td>
<td>2–8 wk (from animal studies)</td>
<td>Total antibody to delta hepatitis shows if ever infected; IgM test is in research laboratories; ELISA</td>
<td>Requires active infection with HBV; injection drug users and persons receiving clotting factor concentrates are at highest risk for infection</td>
</tr>
<tr>
<td>Hepatitis E</td>
<td>Fecal-oral</td>
<td>Average, 26–42 d (range, 15–64 d)</td>
<td>Research laboratories</td>
<td>No known cases originated in the United States; international travelers are the only high-risk group to date</td>
</tr>
<tr>
<td>Hepatitis G</td>
<td>Blood</td>
<td>Unknown</td>
<td>Occurs with hepatitis B and hepatitis C</td>
<td>Recipient of contaminated blood</td>
</tr>
</tbody>
</table>

Viruses for Which Clinical Signs and Symptoms Mimic Hepatitis

<table>
<thead>
<tr>
<th>Virus</th>
<th>Transmission</th>
<th>Incubation Period</th>
<th>Test for Active Infection</th>
<th>Social and Clinical History</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epstein-Barr virus (EBV)</td>
<td>Oropharyngeal (saliva)</td>
<td>4–6 wk</td>
<td>IgM antibody to EBV viral capsid proteins</td>
<td>Serocconversion by age 5 yr in 50% of persons in the U.S.; children with an acutely infected sibling are at greater risk</td>
</tr>
<tr>
<td>Cytomegalovirus (CMV, human herpes-virus 5)</td>
<td>Intimate contact with infected fluids; sexual, perinatal, blood transfusion, and infected breast milk</td>
<td>About 3–8 wk for CMV</td>
<td>Culture, monoclonal antibody to early antigen</td>
<td>Household sexual contact with an infected person, male-to-male sexual practices, day care centers, perinatal transmission</td>
</tr>
</tbody>
</table>


These tests detect human immunodeficiency viruses types 1 and 2 (HIV-1/2), which cause AIDS. Infection with HIV-1 is most prevalent in the United States and Western Europe. Most cases associated with HIV-2 are reported in West Africa. Tests to detect the presence of HIV-1 antibody screen blood and blood products that will be used for transfusion and tissue and organs for transplantation. They are also used to test people at risk for developing AIDS, such as intravenous drug users, sexual partners of HIV-infected persons, and infants born to HIV-infected women. The diagnosis of AIDS must be clinically established. Tests used to determine the presence of antibodies to HIV-1 include ELISA, Western blot, and PCR. PCR has been evaluated as a means to detect viral load by viral nucleic acid test (NAT) after infection but before seroconversion.

A single reactive ELISA test by itself cannot be used to diagnose AIDS. The test should always be repeated in duplicate using the same blood sample. If repeatedly reactive, follow-up tests using Western blot should be done. A positive Western blot is considered confirmatory for HIV. The combination HIV-1/2 test has replaced the HIV-1 test for screening blood and blood products for transfusion. It is also used for testing potential organ transplant donors.

<table>
<thead>
<tr>
<th>Whom to Test</th>
<th>How to Test</th>
<th>When to Test</th>
</tr>
</thead>
</table>

Table 8.7 Diagnostic Testing for HIV
Men who have sex with men, IV drug users, recreational drug users, those engaging in unprotected sex, those attending STD clinics, pregnant women, those with signs and symptoms of unusual pneumonia, skin lesions, mononucleosis-like syndrome; persons known to be infected with HIV

Screening EIA and confirmatory tests. Western blot confirmatory test detects antibodies to HIV-1 core antigens: gp41, gp120, gp160, p18, p24, p31, p40, p65, p55/51. IFA confirmatory test detects potent antibodies by fluorescein-tagged secondary antibodies. Viral RNA and p24 antigen are used along with CD4 count to monitor treatment. Nucleic acid amplification testing (NAT) to monitor “viral load.” Rapid testing: single-use diagnostic system (SUDS) results in 1 hour.

As early a detection as possible so that proper treatment, decrease in transmission, and modified behaviors can occur.

Mother-to-child (vertical transmission) treated during pregnancy and delivery, and exposed infants within 48 hours of delivery; transmission of HIV can occur in utero, during birth, and by breast feeding.

Reference Values

Normal Negative for HIV antibodies against HIV antigens types 1 and 2 by ELISA, enzyme immunoassay (EIA) and Western blot HIV proviral RNA: not reactive or negative by PCR HIV proviral DNA: not reactive or negative HIV core P24 antigen: not reactive or negative NAT: viral load is low

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Plasma may also be used. Observe standard precautions.
2. Be aware that saliva specimens may be collected; usually indicated in clinic settings or outreach environments. See Appendix K for protocols.

Clinical Alert

Detectable HIV antibodies develop within 3 months after exposure to virus.

Clinical Implications

1. A positive test is associated with viral replication and appearance of HIV antibodies (IgM, IgG).
2. A positive ELISA that fails to be confirmed by Western blot or IFA should not be considered negative, especially in the presence of symptoms or signs of AIDS. Repeat testing in 3 to 6 months is suggested.
3. A positive result may occur in noninfected persons because of unknown factors.
4. Negative tests tend to rule out AIDS in high-risk patients who do not have the characteristic opportunistic infections or tumors.
5. An HIV infection is described as a continuum of stages that range from the acute, transient, mononucleosis-like syndrome associated with seroconversion to asymptomatic HIV infection to symptomatic HIV infection and, finally, to AIDS. AIDS is end-stage HIV infection.
6. Treatments are more effective and less toxic when begun early in the course of HIV infection.
7. HIV PCR method to determine viral load may be performed during HIV treatment to monitor patient prognosis and treatment.
8. Diagnosis of HIV in neonates is difficult because maternally acquired antibodies may be present until the child is 18 months of age. Additionally, PCR to detect antigen is usually not successful until the child is 6 months of age.

Interfering Factors

1. Nonreactive HIV test results occur during the acute stage of disease when the virus is present but antibodies are not sufficiently developed to be detected. It may take up to 6 months for the test result to become positive. During this stage, the test for the HIV antigen may confirm an HIV infection (Fig. 8.1).

![FIGURE 8.1 Time course for appearance of viral and serological markers during primary HIV infection. (Source: Grenert JP, Nasses PS, Heller BL: “Human immunodeficiency virus—diagnostic testing at the start of the 21st century.” Clinical Laboratory News 19–23, July 2002)](https://example.com/figure8.1)

2. Test kits for HIV are extremely sensitive. As a result, nonspecific reactions may occur if the tested person has been previously exposed to HIV human cells or the growth media.
Clinical Alert

1. Issues of confidentiality surround HIV testing. Access to test results should be given judiciously on a need-to-know basis unless the patient specifically expresses otherwise. Interventions to block general computer access to this information are necessary; each health care facility must determine how best to accomplish this.
2. Conversely, health care workers directly involved with the care of an HIV/AIDS patient have a right to know the diagnosis so that they may protect themselves from exposure.
3. All results, both positive and negative, must be somehow entered in the patient's health care records while maintaining confidentiality. People are more likely to test voluntarily when they trust that inappropriate disclosure of HIV testing information will not occur. Long-term implications include potential loss of jobs, housing, insurance coverage, and personal relationships.
4. The clinician must sign a legal form stating that the patient has been informed regarding test risks.
5. A person who exhibits HIV antibodies is presumed to be HIV infected; appropriate counseling, medical evaluation, and health care interventions should be discussed and instituted.
6. Positive test results must be reported to the state and federal health authorities according to prescribed state regulations and protocols.
7. Anonymous testing and reporting is available, such as commercial home tests.

Interventions

Pretest Patient Preparation

1. An informed, witnessed consent form must be properly signed by any person being tested for HIV/AIDS. This consent form must accompany the patient and the specimen (see Appendix H for sample form).
2. It is essential that counseling precedes and follows the HIV antibody test. This test should not be performed without the subject's informed consent, and persons who need to access results legitimately must be mentioned. Discussion of the clinical and behavioral implications derived from the test results should address the accuracy of the test and should encourage behavioral modifications (eg, sexual contact, shared needles, blood transfusions).
3. Assess frequency and intensity of symptoms: elevated temperature, anxiety, fear, diarrhea, neuropathy, nausea and vomiting, depression, and fatigue.
4. Infection control measures mandate use of standard precautions (see Appendix A).
5. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes. Explain significance of test results along with CD4+ cell counts. Advise patient that screening tests must be confirmed before the results are reported as HIV reactive. Provide options for immediate counseling if necessary. Explain treatment with potent antiviral drugs and protease inhibitors.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Additional Applications for Oral Specimen Testing

1. HIV-1 and HIV-2
2. Viral hepatitis A, B, and C
3. H. pylori
4. Measles
5. Mumps
6. Rubella
7. Syphilis
8. Cytomegalovirus (CMV)
9. Autoimmune diseases
10. Cancer (carcinoembryonic antigen [CEA], prostate-specific antigen [PSA], CA 125)
11. Diabetes types 1 and 2
12. Therapeutic drug and hormone management and detection of other drugs (see Appendix K for saliva, hair, nails specimen collection)

Procedure for Oral Testing

1. Use a special testing kit such as the commercial Orasure Testing System. The kit's components consist of a specially treated cotton pad on a nylon stick and a vial containing preservative solution. Salt solution in the pad facilitates absorption of the required fluid.
2. Use precise technique. Place pad between the lower cheek and gum, rub back and forth until moistened, and leave in place for 2 minutes. Remove specially treated pad and place it in the vial of special antimicrobial preservative solution. Place specimen container in a biohazard bag and transport to laboratory.
3. The Omni Sal device employs a different collection method in which a cotton pad is placed under the tongue. An indicator in the collecting device changes color when an adequate amount of oral fluid has been collected.
4. Recent food intake, smoking, oral hygiene, or treatment with anticholinergic drugs do not affect test results.
Rubella Antibody Tests

Rubella, a mild, contagious illness characterized by an erythematous maculopapular rash, is observed primarily in children 5 to 14 years of age and in young adults. The disease, commonly called German or 3-day measles, may be asymptomatic or may involve a 1- to 5-day prodromal period of malaise, headache, cold symptoms, low-grade fever, and suboccipital lymphadenopathy.

Although the illness is mild in children, it may cause the congenital rubella syndrome in the fetus of a mother infected early in pregnancy. As many as 85% of infants infected during the first eight weeks of gestation have detectable defects by 4 years of age. The classic abnormalities associated with the rubella syndrome include congenital heart disease, cataracts, and neurosensory deafness. After 20 to 24 weeks of gestation, congenital abnormalities are rare.

The quantitative measurement of IgG antibodies to rubella virus aids in the determination of immune status. Assay results of 10 International Units (IU) of antibody per milliliter (mL) are negative or not immune. Assay results >10 IU are considered positive or immune. A positive result of IgM antibody indicates a congenital or recent infection. The measurement of IgM class antibodies for determination of acute phase infection is recommended in all age groups. IgM rubella antibody determination is usually not recommended when the patient is >6 months of age. Unlike IgG class antibodies, IgM antibodies are larger molecules and cannot cross the placenta, thus determining that the infant has an active form of the disease.

Reference Values

Normal
Negative for rubella IgG and/or IgM antibodies by ELISA or chemilluminescence: not immune. Positive for rubella IgG antibody: immune; indicates a current or previous exposure or immunization to rubella. Positive for rubella IgM antibody (with or without positive IgG): indicates a current or recent infection with rubella virus.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in a biohazard bag for transport to the laboratory.
2. Follow-up testing may be required.

Clinical Implications

1. When testing for IgG antibody, seroconversion between acute and convalescent sera is considered strong evidence of a current or recent infection. The recommended interval between an acute and convalescent sample is 10 to 14 days.
2. A serum specimen taken very early during the acute stage of infection may contain levels of IgG antibody below 10 IU/mL.
3. While the presence of IgM antibody suggests current or recent infection, low levels of IgM may occasionally persist for more than 12 months after infection or immunization. Passively acquired rubella antibody levels (IgG) in the infant (which can cross the placenta because of their smaller molecular size) decrease markedly within 2 to 3 months postinfection.
4. IgM is detectable soon after clinical symptoms occur and reaches peak levels at 10 days.

Interventions

Pretest Patient Preparation

1. Assess patient's test knowledge. Explain test purpose and procedure. Advise pregnant women that rubella infection acquired in the first trimester of pregnancy is associated with an increased incidence of miscarriage, stillbirth, and congenital abnormalities.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel appropriately. Advise women of childbearing age who test negative to be immunized before becoming pregnant. Immunization is contraindicated during pregnancy. Patients who test positive are naturally immune to further rubella infections.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Measles (Rubeola) Antibody Tests

Classified as a paramyxovirus, measles produces a highly contagious respiratory infection. The disease is spread during the prodrome phase through direct contact with respiratory secretions in the form of droplets. Clinical infection with measles virus is characterized by high fever, cough, coryza, conjunctivitis, malaise, and Koplik's spots on the buccal mucosa. An erythematous rash then develops behind the ears and over the forehead, spreading to the trunk.

Serology has become increasingly important as a tool for determining the immune status of the young adult population entering college or the military. In addition, the linkage between measles infection and premature delivery or spontaneous abortion supports screening pregnant mothers for susceptibility.

These tests determine susceptibility and immunity to measles virus. Since intensive immunization began in the U.S. in the 1970s, the incidence of measles infection has been reduced from approximately one half million cases annually.
to fewer than 500 cases in recent years. Many individuals, however, may remain susceptible to measles virus because of vaccine failure or nonimmunization. A positive IgG coupled with a negative IgM result indicates previous exposure to measles virus and immunity to this viral infection. Positive IgM results, with or without positive IgG results, indicate a recent infection with measles virus.

Reference Values

Normal
Negative for measles IgG and/or IgM antibodies by ELISA: not immune. Positive for measles IgG antibody: immune; indicates a current or previous exposure or immunization to measles. Positive for measles IgM antibody (with or without positive IgG): indicates a recent infection with measles virus.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in a biohazard bag for transport to the laboratory.
2. Follow-up testing may be required.

Clinical Implications

1. When testing for IgG antibody, seroconversion between acute and convalescent sera is considered strong evidence of a current or recent infection. The recommended interval between an acute and convalescent sample is 10 to 14 days.
2. While the presence of IgM antibody suggests current or recent infection, low levels of IgM may occasionally persist for more than 12 months after infection or immunization.
3. IgM antibody response is detectable 2 to 3 weeks after appearance of the rash.

Interventions

Pretest Patient Preparation

1. Assess patient's test knowledge. Explain test purpose and procedure. Advise pregnant women that measles poses a high risk for serious complications and may be linked to premature delivery or spontaneous abortion.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel appropriately. Advise women of childbearing age who test negative to be immunized before becoming pregnant. Inform patients who test positive that they are naturally immune to further measles infection.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Mumps Antibody Tests

The mumps virus is a member of the paramyxovirus group and the etiological agent of mumps in humans. Mumps is a generalized illness, usually accompanied by parotid (salivary gland) swelling and mild symptoms. Parotitis as a presenting symptom in mumps is usually sufficient to preclude confirmation by serology. However, one third of mumps infections are subclinical and may require viral isolation to confirm mumps infection. Infection with mumps virus, whether symptomatic or subclinical, is generally thought to offer lifelong immunity.

ELISA testing can be both specific and sensitive for the detection and measurement of serum proteins. Current methods for serodiagnosis of mumps include in vitro serum neutralization, hemagglutination inhibition (HAI), indirect immunofluorescence (IFA), and complement fixation (CF). These test methods, however, lack specificity, which limits their usefulness in establishing immune status.

Reference Values

Normal
Negative for mumps IgG and/or IgM antibodies by ELISA: nonimmune. Positive for mumps IgG antibody: immune; indicates a current or previous exposure or immunization to mumps virus. Positive for mumps IgM antibody: indicates a current or recent infection.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in a biohazard bag for transport to the laboratory.
2. Follow-up testing may be required.

Clinical Implications

1. When testing for IgG antibody, seroconversion between acute and convalescent sera is considered strong evidence of a current or recent infection.
2. The recommended interval between an acute and convalescent sample is 10 to 14 days.

Varicella-Zoster (Chickenpox) Antibody Test

Varicella-zoster virus (VZV) is a herpesvirus and causes chickenpox with primary infection, a highly contagious disease characterized by widely spread vesicular eruptions and fever. The disease is endemic in the U.S. and most commonly affects children from 5 to 8 years of age, although adults and younger children, including infants, may develop chickenpox. VZV infection in a pregnant woman may spread through the placenta to the fetus, causing congenital
disease in the infant.

Though a primary infection results in immunity to subsequent chickenpox, the virus remains latent in the body. When it is reactivated, VZV causes shingles (herpes zoster). Fever and painful localized vesicular eruptions of the skin along the distribution of the involved nerves are the most common clinical symptoms.

The sensitivity, specificity, and reproducibility of ELISA immunoassays are comparable to other serological tests for antibody such as immunofluorescence, complement fixation, and hemagglutination. A positive IgG result coupled with a positive IgM result indicates a current infection with VZV.

Reference Values

Normal: Negative for varicella-zoster IgG and/or IgM antibodies by ELISA: non-immune. Positive for varicella-zoster IgG antibody: indicates a current or previous infection; in the absence of current clinical symptoms, may indicate immunity. Positive for varicella-zoster IgM antibody: indicates a current or recent infection.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in a biohazard bag for transport to the laboratory.
2. Follow-up testing may be required.

Clinical Implications

1. When testing for IgG antibody, seroconversion between acute and convalescent sera is considered strong evidence of a current or recent infection. The recommended interval between an acute and convalescent sample is 10 to 14 days.
2. Whereas the presence of IgM antibody suggests a current or recent infection, low levels of IgM may occasionally persist for more than 12 months after infection or immunization.
3. Immunosuppressed patients in hospitals may contract severe nosocomial infections from others infected with VZV. Therefore, serologic screening of direct health care providers (physicians, nurses, etc.) is necessary to avoid spread of infection.

Interventions

Pretest Patient Preparation

1. Assess patient's test knowledge. Explain test purpose and procedure. Advise pregnant women that VZV poses a high risk of congenital disease in the infant.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel appropriately. Inform patients who test positive for VZV IgG that they are naturally immune to chickenpox, but the virus can be reactivated and cause shingles at a later time.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Cytomegalovirus (CMV) Antibody Test

Cytomegalovirus (CMV) is a ubiquitous human viral pathogen that belongs to the herpesvirus family. Infection with CMV is usually asymptomatic and can persist in the host as a chronic or latent infection. Cytomegalovirus has been linked with sexually transmitted infections. Blood banks routinely screen for CMV antibodies and report these as CMV-negative or CMV-positive.

This test determines the presence of CMV antibodies and is routinely done in congenitally infected newborns, immunocompromised patients, and sexually active persons who present with mononucleosis-like symptoms. Antibody results must be evaluated in the context of the patient's current clinical symptoms and viral culture results. Tests to detect CMV antigen are available and aid in early detection. Viral culture confirms CMV infection.

Reference Values

Normal: Negative for CMV-specific IgG and IgM by ELISA.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in biohazard bag for transport to the laboratory.
2. It is recommended that posttransplant titers be monitored at weekly intervals, particularly following bone marrow transplant.

Clinical Implications

1. Infants who acquire CMV during primary infection of the mother are prone to develop severe cytomegalic inclusion disease (CID). CID may be fatal or may cause neurologic sequelae such as mental retardation, deafness, microcephaly, or motor dysfunction.
2. Transfusion of CMV-infected blood products or transplantation of CMV-infected donor organs may produce interstitial pneumonitis in an immunocompromised recipient.
3. When testing for IgG antibody, seroconversion or a significant rise in titer between acute and convalescent sera may indicate presence of a current or recent infection.
While the presence of IgM antibodies suggests current or recent infection, low levels of IgM antibodies may occasionally persist for more than 12 months postinfection.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results. Refer to page 532. Counsel appropriately.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Herpes Simplex Virus (HSV) Antibodies (HSV-1 and HSV-2 Tests)**

Two types of herpes simplex virus exist. Herpes simplex virus type 1 (HSV-1) causes orofacial herpes; type 2 (HSV-2) causes genital and neonatal herpes. Serologic differentiation is difficult; therefore, type-specific antibody tests are required.

These tests identify the herpes simplex infections. Human herpes simplex virus (HSV) is found worldwide and is transmitted by close personal contact. The clinical course is variable, and symptoms may be mild enough to go unrecognized. Major signs and symptoms include oral and skin eruptions, genital tract infections and lesions, and neonatal herpes. Herpes simplex is also common in individuals with immune system deficiencies (eg, cancers, HIV/AIDS, chemotherapy treatment). HSV antibody testing is also widely used for bone marrow recipients and donors.

**Reference Values**

**Normal** Negative for HSV-1 and HSV-2 by ELISA and IFA.

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in biohazard bag for transport to the laboratory.
2. Follow-up testing is usually required.

**Clinical Implications**

1. Most persons in the general population have been infected with HSV by 20 years of age. After the primary infection, antibody levels fall and stabilize until a subsequent infection occurs.
2. Diagnosis of current infection is related to determining a significant increase in antibody titers between acute-stage and convalescent-stage blood samples.
3. Serologic tests cannot indicate the presence of active genital tract infections. Instead, direct examination with procurement of lesion cultures should be done.
4. Newborn infections are acquired during delivery through the birth canal and may present as localized skin lesions or more generalized organ system involvement.

**Interventions**

**Pretest Patient Preparation**

1. Assess patient's knowledge regarding the test. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes. Refer to page 532 for interpretation of immunologic test results. Advise pregnant women that the newborn may be infected during birth when active genital-area infection is present. Explain need for repeat testing.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Human T-Cell Lymphotropic Virus (HTLV-I/II) Antibody Test**

This test detects antibodies to HTLV-I, a retrovirus associated with adult T-cell leukemia (ATL) and demyelinating neurologic disorders. The presence of HTLV-I antibodies in an asymptomatic person excludes that person from donating blood; however, this finding does not mean that leukemia or a neurologic disorder exists or will develop. Specimens with a positive test result by EIA are referred for Western blot. The results of Western blot are for investigational use only at the time of this printing.

**Reference Values**

**Normal** Negative for HTLV-I/II antibodies by EIA and Western blot

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**
1. Positive results (antibodies to HTLV-I) occur in the presence of HTLV-I infection. Infection transmitted to recipients of HTLV-I–infected blood is well documented.

2. The presence of antibodies to HTLV-I bears no relation to the presence of antibodies to HIV-1; its presence does not put a person at risk for HIV/AIDS, but they often occur concurrently because of similar risk factors.

3. HTLV-I is endemic to the Caribbean, Southeastern Japan, and some areas of Africa.

4. In the United States, HTLV-I has been detected in persons with ATL, intravenous drug users, and healthy persons as well as in donated blood products. Transmission can also take place through ingestion of breast milk, sexual contact, and sharing of contaminated intravenous drug paraphernalia.

**Interventions**

*Pretest Patient Preparation*

1. Assess patient's knowledge about test. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

*Posttest Patient Aftercare*

1. Interpret test results. Refer to page 532 for interpretation of immunologic test results. Counsel patient appropriately.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Parvovirus B-19 Antibody Test**

These tests detect parvovirus B-19, the only parvovirus known to cause human disease. The B-19 virus destroys red blood cell precursor cells and interferes with normal red blood cell production. In young children, it is associated with erythema infectiosum, a mild, self-limiting disease characterized by a low-grade fever and rash. Recently, it has been associated with aplastic crisis in patients with chronic hemolytic anemia and in immunodeficient patients who have bone marrow failure.

**Reference Values**

*Normal* Negative for parvovirus B-19–specific IgG and IgM antibodies by ELISA and indirect fluorescent antibody (IFA)

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place sample in a biohazard bag for transport to the laboratory.

**Clinical Implications**

1. Positive parvovirus B-19 infection has been implicated in aplastic anemia associated with organ transplantation. It is recommended, therefore, that this test be included in the serologic assessment of prospective organ donors.
2. Immunocompromised patients may have a delayed or absent antibody response. It is recommended that parvovirus DNA detection by PCR be considered.

**Interventions**

*Pretest Patient Preparation*

1. Assess patient's knowledge regarding test. Explain purpose and blood test procedure. Advise any prospective organ donor that this test is part of a panel of tests performed before organ donation to protect the organ recipient from potential infection.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

*Posttest Patient Aftercare*

1. Interpret test outcome. Refer to page 532 for interpretation of immunologic test results. Explain significance of test results.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Rabies Antibody Tests**

Serologic testing is diagnostic for the presence of rabies in animals. It also indicates the degree of antibody responses to rabies immunization (eg, for people who routinely work with animals).

**Reference Values**

*Normal* IFA <1:16 or DFA examination of the animal brain for presence of the virus.

**Procedure for Humans** Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in a biohazard bag.

**Procedure for Animals**

1. If the suspect animal exhibits abnormal behavior, standard procedure is to sacrifice it and examine its brain for Negri body inclusions in the neurons.
2. Rabies testing is usually performed in a public health laboratory.

**Clinical Implications** An elevated titer in humans indicates an adequate response after immunization. A rabies titer of 1:16 or greater is considered protective.
Interventions

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding *safe, effective, informed pretest care*.

**Posttest Patient Aftercare**

1. Interpret test results after immunization.
2. Follow guidelines in Chapter 1 regarding *safe, effective, informed posttest care*.

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**Clinical Alert**

1. Prevention: Pre-exposure vaccine (human diploid cell rabies vaccine [HDCV]) should be given to persons at high-risk such as veterinarians, wildlife personnel, zoo workers, quarantine kennel workers, and those employed in laboratories that use animals.
2. Postbite: Administer rabies immunoglobulin (RIG) as soon as possible after the bite, regardless of time interval, to neutralize the virus in the wound. HDCV in five 1-mL intramuscular doses should be given in the deltoid muscle. The first HDCV dose is given concurrently with the RIG, and subsequent doses are given 3, 7, 14, and 28 days after the first dose.
3. The animal brain should be tested as soon as possible. Holding the animal for observation is not recommended.

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**Fungal Tests**

**Fungal Antibody Tests: Histoplasmosis, Blastomycosis, Coccidioidomycosis**

Certain fungal species are associated with human respiratory diseases acquired by inhaling spores from sources such as dust, soil, and bird droppings. Serologic tests may be used for diagnosis. Fungal diseases are categorized as either superficial or deep. For the most part, superficial mycoses are limited to the skin, mucous membranes, nails, and hair. Deep mycoses involve the deeper tissues and internal organs. Histoplasmosis, coccidioidomycosis, and blastomycosis are caused by deep mycoses.

These tests detect serum precipitin antibodies and CF antibodies present in the fungal diseases of coccidioidomycosis, blastomycosis, and histoplasmosis. Coccidioidomycosis, also known as desert fever, San Joaquin fever, and valley fever, is contracted through inhalation of *Coccidioides immitis* spores found in dust or soil. Blastomycosis is caused by infection with organisms of the genus *Blastomyces*. Histoplasmosis is a granulomatous infection caused by *Histoplasma capsulatum*.

**Reference Values**

**Normal** Negative for fungal antibodies

**CF titer:** <1:8

**Immunodiffusion:** negative

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place in a biohazard bag for transport to the laboratory.

**Clinical Implications**

1. Antibodies to *Coccidioides, Blastomyces*, and *Histoplasma* appear early in the course of the disease (weeks 1–4) and then disappear.
2. Negative fungal serology does not rule out the possibility of a current infection.

**Interfering Factors**

1. Antibodies to fungi may be found in blood samples from apparently healthy people.
2. When testing for blastomycosis, cross-reactions with histoplasmosis may occur.
3. More than 50% of patients having active blastomycosis yield a negative result by CF.
4. Recent histoplasmosis skin tests must be avoided because they cause elevated CF test results, which may be due to the stimulation from the skin test and not the systemic infection.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding *safe, effective, informed pretest care*.
3. Remember that specimens for culture of the organism may also be required.

**Posttest Patient Aftercare**

1. Interpret test results. Refer to page 532 for interpretation of immunologic test results. Counsel appropriately.
2. Follow guidelines in Chapter 1 regarding *safe, effective, informed posttest care*.

**Candida Antibody Test**
Candidiasis is usually caused by *Candida albicans* and affects the mucous membranes, skin, and nails (see *Candida Skin Test*, page 527). Compromised individuals with depressed T-cell function are most likely to have invasive disease.

Identifying the *Candida* antibody can be helpful when the diagnosis of systemic candidiasis cannot be shown by culture or tissue sample. Clinical symptomatology must be present for the test to be meaningful. Tests used include immunodiffusion; counter-immunoelectrophoresis (CIE), which is particularly valuable on CSF and urine specimens; and latex agglutination for *Candida* antigen.

**Reference Values**

**Normal** Negative for Candida antibodies by immunodiffusion (ID)

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place in a biohazard bag for transfer to the laboratory.

**Clinical Implications**

1. A titer greater than 1:8 by latex agglutination for *Candida* antigen indicates systemic infection.
2. A fourfold rise in titers of paired blood samples 10 to 14 days apart indicates acute infection.
3. Patients on long-term intravenous therapy treated with broad-spectrum antibiotics and diabetic patients commonly have disseminated infections caused by *Candida albicans*. The disease also occurs in bottle-fed newborns and in the urinary bladder of catheterized patients.
4. Vulvovaginal candidiasis, common in late pregnancy, can transmit candidiasis to the infant through the birth canal.

**Interfering Factors**

1. Approximately 25% of the normal population tests positive for the presence of *Candida*.
2. Cross-reaction can occur with latex agglutination testing in persons who have cryptococcosis or tuberculosis.
3. Positive results can occur in the presence of mucocutaneous candidiasis or severe vaginitis.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.
3. Specimens for culture of the organism may also be required.

**Posttest Patient Aftercare**

1. Interpret test results. Refer to page 532 for interpretation of immunologic test results. Counsel patient appropriately. Repeat testing is usually indicated.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Aspergillus Antibody Test**

The aspergilli, especially *Aspergillus fumigatus*, *Aspergillus flavus*, and *Aspergillus niger*, are associated with pulmonary infections and invasive fatal disease sequelae in immunosuppressed patients. Manifestations of *Aspergillus* infections include allergic bronchopulmonary disease, lung mycetoma, endophthalmitis, and disseminated brain, kidney, heart, and bone disease.

This test detects antibodies present in aspergillosis, primarily allergic bronchopulmonary disease, or fungus ball.

**Reference Values**

**Normal** Negative for *Aspergillus* antibody by immunodiffusion <1:8 by CF

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. CSF can also be tested. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**

1. Positive test results are associated with pulmonary infections in compromised patients and *Aspergillus* infections of prosthetic heart valves.
2. If blood serum exhibits one to four bands using immunodiffusion, aspergillosis is strongly suspected. Weak bands suggest an early disease process or hypersensitivity pneumonitis.
3. Best use of the CF test is with paired sera taken 3 weeks apart to detect a rise in titer against a single antigen.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.
3. Specimens for culture of the organism may also be required.

**Posttest Patient Aftercare**

1. Interpret test outcome. Refer to page 532 for interpretation of immunologic test results. Counsel appropriately.
2. Follow guidelines in Chapter 1 regarding *safe, effective, informed posttest care*.

**Cryptococcus Antibody Test**

*Cryptococcus neoformans*, a yeast-like fungus, causes a lung infection thought to be acquired by inhalation. The organism has been isolated from several natural environments, especially where weathered pigeon droppings accumulate.

This test detects antibodies present in *Cryptococcus* infections. It appears that about 50% of patients who present with antibodies have a predisposing condition such as lymphoma or sarcoidosis or are being treated with steroid therapy. Infection with *C. neoformans* has long been associated with Hodgkin's disease and other malignant lymphomas. In fact, *C. neoformans*, in conjunction with malignancy, occurs to such a degree that some researchers have raised the question regarding the possible etiologic relation between the two diseases. Tests ordered for this disease include latex agglutination testing for antigens or antibodies.

**Reference Values**

*Normal* Negative for *Cryptococcus* antibody by indirect fluorescent antibody (IFA)

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. A 2-mL spinal fluid sample may also be used. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**

Positive *C. neoformans* tests are associated with infections of the lower respiratory tract through inhalation of aerosols containing *C. neoformans* cells disseminated by the fecal droppings of pigeons.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure. Obtain clinical history and assess for exposure.
2. Follow guidelines in Chapter 1 regarding *safe, effective, informed pretest care*.
3. Specimens for culture of the organism may also be required.

**Posttest Patient Aftercare**

1. Interpret test results. Refer to page 532 for interpretation of immunologic test results. Counsel patient appropriately.
2. Follow guidelines in Chapter 1 regarding *safe, effective, informed posttest care*.

**PARASITIC TESTS**

**Toxoplasmosis (TPM) Antibody Tests**

Toxoplasmosis is caused by the sporozoan parasite *Toxoplasma gondii* and is a severe, generalized, granulomatous CNS disease. It may be either congenital or acquired and is found in humans, domestic animals (eg, cats), and wild animals. Humans may acquire the infection through ingestion of inadequately cooked meat or other contaminated material. Congenital toxoplasmosis may cause fetal death. Symptoms of subacute infection may appear shortly after birth or much later. Complications of congenital toxoplasmosis include hydrocephaly, microcephaly, convulsions, and chronic retinitis. It is believed that one fourth to one half of the adult population is asymptotically infected with toxoplasmosis. The CDC recommends serologic testing during pregnancy.

The indirect fluorescent antibody (IFA) test helps to differentiate toxoplasmosis from IM. Toxoplasmosis antibodies appear within 1 to 2 weeks of infection and peak at 6 to 8 months. IFA is also a valuable screening test for latent toxoplasmosis.

**Reference Values**

*Normal* Titer <1:16; no previous infection (*except* for ocular infection) by IFA Negative by MEIA Negative: *T. gondii* DNA not detected by PCR

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**

The IFA test is considered positive under any of the following conditions:

1. Titer of 1:256 or higher indicates recent exposure or current infection; rising titer is of greatest significance.
2. Any titer value is significant in a newborn infant.
3. Titer of 1:1024 or greater is significant for active disease.
4. Titer of 1:16 or less occurs with ocular toxoplasmosis.

**Interventions**
Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results. Refer to page 532 for interpretation of immunologic test results. Counsel appropriately.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Amebiasis (Entamoeba histolytica) Antibody Test

Entamoeba histolytica, the causative agent of amebiasis, is a pathogenic intestinal parasite. The E. histolytica test determines the presence or absence of specific serum antibodies to this parasite. Stool examination is considered the definitive diagnostic tool; however, the absence of detectable stool organisms does not necessarily rule out the disease. Antibiotic therapy, oil enemas, and barium may interfere with the ability to isolate this organism in the stool.

Reference Values

Normal Negative for Entamoeba antibodies by indirect hemagglutination, latent agglutination, and CIE.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

Clinical Implications

1. Positive test (titer > 1:128) indicates active or recent infection.
2. Amebic liver abscess and amebic dysentery indicate the presence of amebiasis.
3. Titers range from 1:256 to 1:2048 in the presence of current active amebiasis.

NOTE

A positive test may only reflect past but not current infections.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results. Refer to page 532 for interpretation of immunologic test results. Counsel patient appropriately.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

TORCH Test

TORCH is an acronym that stands for Toxoplasma, rubella, CMV, and herpes simplex virus (HSV). These pathogens are frequently implicated in congenital or neonatal infections that are not clinically apparent but that may result in serious CNS impairment.

Both mothers and newborn infants are tested for exposure to these agents. The test differentiates acute, congenital, and intrapartum infections caused by T. gondii, rubella virus, CMV, and herpesvirus. The presence of IgM-associated antibodies in newborns reflects actual fetal antibody production. High levels of IgM at birth indicate fetal in utero response to an antigen. In this instance, an intrauterine infection should be considered. TORCH is more useful in excluding than in establishing etiology.

Reference Values

Normal Negative for Toxoplasma, rubella, CMV, and HSV antibodies

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

Clinical Implications

1. Persistent rubella antibodies in an infant >6 months of age highly suggests congenital infection. Congenital rubella is characterized by neurosensory deafness, heart anomalies, cataracts, growth retardation, and encephalitic symptoms.
2. A diagnosis of toxoplasmosis is established through sequential testing rather than by a single positive result. Sequential examination reveals rising antibody titers, changing titers, and the conversion of serologic tests from negative to positive. A titer of 1:256 suggests recent infection. About one third of infants who acquire infection in utero show signs of cerebral calcifications and chorioretinitis at birth; the rest are born without symptoms.
3. A marked and persistent rise in CF antibody titer over time is consistent with a diagnosis of rubella in infants <6 months of age.
4. Presence of herpes antibodies in CSF, together with signs of herpetic encephalitis and persistent HSV-1 or HSV-2 antibody levels in a newborn showing no obvious external lesions is consistent with a diagnosis of HSV.

Interventions

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results. Refer to page 532 for interpretation of immunologic test results. Monitor and counsel appropriately for intrauterine and congenital infections.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**IMMUNOLOGIC TESTS FOR IMMUNE DYSFUNCTION AND RELATED DISORDERS OF THE IMMUNE SYSTEM**

**Quantitative Immunoglobulins: IgA, IgG, IgM**

Five classes of immunoglobulins (antibodies)—IgA, IgG (with four subclasses, IgG 1, IgG 2, IgG 3, and IgG 4), IgM, IgD, and IgE—have been isolated. Immunoglobulins function to neutralize toxic substances, support phagocytosis, and destroy microorganism functions. For example, IgA takes two forms: serum and secretory. Serum IgA is present in blood serum; secretory IgA is found in saliva, tears, colostrum, and bronchial, gastrointestinal, and genitourinary secretions, where it can protect against microorganism invasion. IgE is involved in allergic reactions, whereas IgD is involved in humoral immunity.

IgG, the only immunoglobulin that can cross the placenta, is responsible for protection of the newborn during the first months of life. IgM possesses antibody activity against gram-negative organisms and rheumatoid factors and forms natural antibodies such as the ABO blood group. IgM does not cross the placenta and is therefore usually absent in the newborn. It is observed about 5 days after birth.

Quantitative immunoglobulin measurements can monitor the course of a disease and its treatment. If there is a monoclonal protein or M component present on serum protein electrophoresis (SPEP), a quantitative measurement of IgA, IgG, and IgM can identify the specific immunoglobulin. IgD and IgE are present in trace amounts.

**Reference Values** These values are derived from rate nephelometry.

1. **Adults**
   a. IgG: 700–1500 mg/dL
   b. IgA: 60–400 mg/dL
   c. IgM: 60–300 mg/dL
   d. IgE: 3–423 IU/mL (3–423 KIU/L)
   e. IgD: 565–1765 mg/dL

2. **Children**
   a. IgA (boys and girls)
      1. 0–4 months: 6–64 mg/dL
      2. 5–8 months: 10–87 mg/dL
      3. 9–14 months: 17–94 mg/dL
      4. 15–23 months: 17–94 mg/dL
      5. 2–3 years: 24–192 mg/dL
      6. 4–6 years: 26–232 mg/dL
      7. 7–9 years: 33–258 mg/dL
      8. 10–12 years: 45–285 mg/dL
      9. 13–15 years: 47–317 mg/dL
     10. 16–17 years: 55–377 mg/dL
   b. IgM (boys)
      1. 0–4 months: 14–142 mg/dL
      2. 5–8 months: 24–167 mg/dL
      3. 9–23 months: 35–200 mg/dL
      4. 2–3 years: 41–200 mg/dL
      5. 4–17 years: 47–200 mg/dL
   c. IgM (girls)
      1. 0–4 months: 14–142 mg/dL
      2. 5–8 months: 24–167 mg/dL
      3. 9–23 months: 35–242 mg/dL
      4. 2–3 years: 41–242 mg/dL
      5. 4–17 years: 56–242 mg/dL
   d. IgG (boys and girls)
      1. 0–4 months: 141–930 mg/dL
      2. 5–8 months: 250–1190 mg/dL
      3. 9–11 months: 320–1250 mg/dL
      4. 1–3 years: 400–1250 mg/dL
5. 4–6 years: 560–1307 mg/dL
6. 7–9 years: 598–1379 mg/dL
7. 10–12 years: 638–1453 mg/dL
8. 13–15 years: 680–1531 mg/dL
9. 16–17 years: 724–1611 mg/dL

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in biohazard bag for transport to the laboratory.

Clinical Implications

1. IgA accounts for 10% to 15% of total immunoglobulin. Increases occur in the following conditions:
   a. Chronic, nonalcoholic liver diseases, especially primary biliary cirrhosis (PBC)
   b. Obstructive jaundice
   c. Exercise
   d. Alcoholism
   e. Subacute and chronic infections
2. IgA decreases occur in the following conditions:
   a. Ataxia-telangiectasia
   b. Chronic sinopulmonary disease
   c. Congenital deficit
   d. Late pregnancy
   e. Prolonged exposure to benzene immunosuppressive therapy
   f. Abstinence from alcohol after a period of 1 year
   g. Drugs and dextrin
   h. Protein-losing gastroenteropathies
3. IgG constitutes 75% to 80% of total immunoglobulins. Increases occur in the following conditions:
   a. Chronic granulomatous infections
   b. Hyperimmunization
   c. Liver disease
   d. Malnutrition (severe)
   e. Dysproteinemia
   f. Disease associated with hypersensitivity granulomas, dermatologic disorders, and IgG myeloma
   g. Rheumatoid arthritis
4. IgG decreases occur in the following conditions:
   a. Agammaglobulinemia
   b. Lymphoid aplasia
   c. Selective IgG, IgA deficiency
   d. IgA myeloma
   e. Bence Jones proteinemia
   f. Chronic lymphoblastic leukemia
5. IgM constitutes 5% to 10% of total antibody. Increases in adults occur in the following conditions:
   a. Waldenström's macroglobulinemia
   b. Trypanosomiasis
   c. Malaria
   d. Infectious mononucleosis
   e. Lupus erythematosus
   f. Rheumatoid arthritis
   g. Dysgammaglobulinemia (certain cases)
6. IgM decreases occur in the following conditions:
   a. Agammaglobulinemia
   b. Lymphoproliferative disorders (certain cases)
   c. Lymphoid aplasia
   d. IgG and IgA myeloma
   e. Dysgammaglobulinemia
   f. Chronic lymphoblastic leukemia

Clinical Alert

Persons with IgA deficiency are predisposed to autoimmune disorders and can develop antibody to IgA, with possible anaphylaxis occurring if transfused with blood containing IgA.

Clinical Alert

In the newborn, a level of IgM >20 mg/dL indicates in utero stimulation of the immune system (eg, rubella virus, cytomegalovirus, syphilis, toxoplasmosis).

Interventions

Pretest Patient Preparation

1. Explain test purpose and specimen collection procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.
Posttest Patient Aftercare

1. See posttest care for protein electrophoresis.
2. Interpret test outcome. Follow-up immunoglobulin testing may be necessary, along with serum viscosity, to monitor a patient with monoclonal gammopathy.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Protein Electrophoresis (PEP), Serum and Urine

Serum proteins represent a diverse microenvironment. They are a source of nutrition and a buffer system. Immunoglobulins and related proteins function as immunologic agents. Carrier proteins (e.g., haptoglobin, prealbumin, transferrin) transport certain ions and molecules to their destinations. Antiproteases (e.g., alpha\textsubscript{1}-antitrypsin, alpha\textsubscript{2}-macroglobulin) regulate the activity of various proteolytic enzymes, and other classes of proteins regulate oncotic pressure, genetic component pressures (e.g., chromosomal), and metabolic substances (e.g., hormones). Blood serum and urine are commonly screened for the monoclonal immunoglobulin component by means of serum protein electrophoresis (SPEP). Immunoglobulins are the major component of the serum gamma-globulin fraction. In health, the immunoglobulins are polyclonal instead of monoclonal. When a monoclonal band is observed, it frequently signals a neoplastic process such as multiple myeloma or Waldenström’s macroglobulinemia. SPEP enhances follow-up procedures such as specific protein quantification of immunoglobulins (IgA, IgG, IgM) and immunofixation. It provides one of the best tools for general screening of the human health state.

These tests can diagnose some inflammatory and neoplastic states, nephrotic syndromes, liver disease, and immune dysfunctions and can evaluate nutritional states and osmotic pressures in edematous and malnourished patients. SPEP produces electrophoretic separation of the five major protein fractions (albumin, alpha\textsubscript{1}-globulin, alpha\textsubscript{2}-globulin, beta-globulin, and gamma-globulin) in serum and urine specimens so that a more definitive diagnosis can be made. Major components present in each protein fraction or zone exhibit characteristic, unique electrophoretic patterns and are defined as the albumin zone (albumin); the alpha\textsubscript{1} zone (alpha\textsubscript{1}-lipoproteins, high-density lipoprotein, alpha\textsubscript{2}-antritrypsin); the alpha\textsubscript{2} zone (alpha\textsubscript{2}-macroglobulin, haptoglobin, beta-lipoprotein); the beta zone (transferrin, C3 [complement]); and the gamma zone (fibrinogen, IgA, IgM, IgG).

Reference Values: Urine Protein Electrophoresis (UPE) A descriptive report is prepared by the pathologist.

Reference Values: Serum Protein Electrophoresis (SPE)

<table>
<thead>
<tr>
<th>Total Protein</th>
<th>Albumin</th>
<th>Reference Values: Serum Protein Electrophoresis (SPE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult: 6.0–8.0 g/dL</td>
<td>60–80 g/L</td>
<td>Albumin: 3.8–5.0 g/dL</td>
</tr>
<tr>
<td>&lt;5 d: 5.4–7.0 g/dL</td>
<td>54–70 g/L</td>
<td>Newborn: 2.6–3.6 g/dL</td>
</tr>
<tr>
<td>1–3 yr: 5.9–7.0 g/dL</td>
<td>59–70 g/L</td>
<td>1–3 yr: 3.4–4.2 g/dL</td>
</tr>
<tr>
<td>4–6 yr: 5.9–7.8 g/dL</td>
<td>59–78 g/L</td>
<td>4–6 yr: 3.5–5.2 g/dL</td>
</tr>
<tr>
<td>7–9 yr: 6.2–8.1 g/dL</td>
<td>62–81 g/L</td>
<td>7–9 yr: 3.7–5.6 g/dL</td>
</tr>
<tr>
<td>10–19 yr: 6.3–8.6 g/dL</td>
<td>63–86 g/L</td>
<td>10–19 yr: 3.7–5.6 g/dL</td>
</tr>
<tr>
<td>a\textsubscript{1}-Globulin: 0.1–0.3 g/dL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a\textsubscript{2}-Globulin: 0.6–1.0 g/dL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ß-Globulin: 0.7–1.4 g/dL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gamma-globulin: 0.7–1.6 g/dL</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. First voided morning urine specimen or 24-hour timed urine specimen is preferred. A 100-mL sample from a 24-hour urine collection is submitted for a urine protein electrophoresis.
3. If blood or urine sample demonstrates the presence of a paraprotein, a follow-up or confirmatory immunofixation electrophoresis (IFE; see page 578) can be performed on the same specimen submitted for the protein electrophoresis.
4. To quantify the amount of protein in each fraction, separate proteins are scanned and separated according to net molecular charge by means of a densitometer and are expressed in grams per deciliter (g/dL).

Clinical Implications The following are the most frequent protein abnormalities in protein quantification and SPEP:

1. Total serum protein (the sum of circulating serum proteins) increases (hyperproteinemia) in dehydration and hemoconcentration states because of fluid loss (e.g., vomiting, diarrhea, poor kidney function); increases are also found in the following conditions:
   a. Liver disease
   b. Multiple myeloma and other gammopathies
   c. Waldenström’s macroglobulinemia
   d. Tropical disease
   e. Sarcoidosis and other granulomatous diseases
   f. Collagen disorders such as systemic lupus erythematosus (SLE) and rheumatoid arthritis (RA)
   g. Chronic inflammatory states
   h. Chronic infections
2. Total serum protein decreases (hypoproteinemia) in the following conditions:
   a. Insufficient nutritional intake (starvation or malabsorption)
b. Severe liver disease or alcoholism
c. Renal disease, nephrotic syndrome
d. Diarrhea (Crohn's disease, ulcerative colitis)
e. Severe skin diseases or burns
f. Severe hemorrhage (when plasma volume is replaced more rapidly than protein)
g. Heart failure
h. Hypothyroidism
i. Prolonged immobilization (trauma, orthopedic surgery)

3. Serum albumin increases with intravenous infusions and dehydration (elevated hemoglobin and hematocrit indicate higher albumin levels).

4. Serum albumin decreases in the following conditions:
a. Decreased synthesis states such as liver disease, alcoholism, malabsorption syndromes, Crohn's disease, other protein-losing enteropathies, starvation states, and congenital analbuminemia
b. Increased albumin loss (eg, nephrotic syndrome, third-degree burns)
c. Poor nutrition states and inadequate iron intake
d. Low albumin-to-globulin (A/G) ratio (eg, collagen disease, chronic inflammation, liver diseases, macroglobulinemia, severe infections, cachexia, burns, ulcerative colitis)

5. Alpha-1-globulin increases with infections (acute and chronic) and febrile reactions.
6. Alpha-1-globulin decreases with nephrosis and alpha-antitrypsin difference.
7. Alpha-1-globulin increases in the following conditions:
   a. Biliary cirrhosis
   b. Obstructive jaundice
   c. Nephrosis
   d. Multiple myeloma (rare)
   e. Ulcerative colitis
8. Alpha-1-globulin decreases in acute hemolytic anemia.
9. Beta-globulin increases in biliary cirrhosis, obstructive jaundice, and multiple myeloma (occasional).
11. Gamma-globulin increases in the following conditions:
    a. Chronic infections
    b. Hepatic diseases
    c. Autoimmune diseases
    d. Collagen diseases
    e. Multiple myeloma
    f. Waldenström's macroglobulinemia
    g. Leukemia and other cancers
12. Gamma-globulin decreases in the following conditions:
    a. Agammaglobulinemia
    b. Hypogammaglobulinemia
    c. Nephrotic syndrome

Interfering Factors

1. Decreased albumin can be seen with rapid intravenous fluid infusions and hydration and during all trimesters of pregnancy.
2. Excessive hemolysis decreases albumin 0.5 g/mL when patients are in the supine position. Conversely, hemolysis and dehydration elevate the total serum protein.
3. Prolonged bed rest and the last trimester of pregnancy produce lower total protein levels.

Interventions

Pretest Patient Preparation

1. Explain test purpose and specimen collection procedure.
2. If a 24-hour urine specimen is to be collected, the patient will require specific instructions, an appropriate container, and a receptacle for catching the voided urine (see Chap. 3, Urine Studies).
3. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately. Very low levels of protein and albumin are associated with edema and hypocalcemia.
2. Assess the patient for signs and symptoms related to these conditions and report and document same. Rarely is any one type of electrophoretic analysis used to diagnose a gammopathy. Follow-up testing may include immunofixation (IFE), quantitative immunoglobulins, and bone marrow studies.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Clinical Alert

Normally, very little protein is excreted in the urine; however, relatively large amounts may be excreted in certain disease states. In the presence of lipoid nephrosis, selective proteinuria produces excess albumin excretion. With nonselective proteinuria (eg, glomerulonephritis), all types of serum proteins usually appear in the urine. Urine protein electrophoresis can identify Bence Jones proteins, which migrate in the ß-globulin and gamma-globulin regions. See Chap. 3 for a complete explanation of urine protein and albumin.
Immunofixation Electrophoresis (IFE), Serum and Urine

Monoclonal immunoglobulins consist of heavy and light chains. IFE identifies presence or absence of a monoclonal protein and determines its heavy-chain and light-chain types.

This test measures immune status and competence by identifying monoclonal and particle protein band immunoglobulins involved in the immune response. IFE is a follow-up test performed when monoclonal spike is observed on SPEP or when a monoclonal gammopathy is suspected on the basis of the patient's immunoglobulin concentrations.

Reference Values
Normal No abnormality present

Procedure
1. Collect a 7-mL blood serum sample in a red-topped tube and/or a 24-hour urine specimen. Observe standard precautions. Submit 25 mL from a 24-hour urine collection, if a urine IFE is to be run simultaneously.
2. If the IFE is a follow-up to a paraprotein being demonstrated by protein electrophoresis (see page 575), the same specimen (blood, urine, or both) used for the electrophoresis can be used for this procedure as well.
3. In IFE, high-resolution electrophoresis produces stained bands. By comparing the location of the stained immunofixed band with a band in the same location in the SPEP reference pattern, a particular protein band can be identified.

Clinical Implications
1. Monoclonal protein in the serum or urine suggests a neoplastic process; a polyclonal increase in immunoglobulins is seen in chronic liver disease, connective tissue disease, and infection.
2. In multiple myeloma, 99% of patients have a monoclonal protein in the serum or urine. Waldenström's macroglobulinemia is characterized by the presence of a serum monoclonal IgM protein in all cases.
3. A monoclonal light chain (K or Bence Jones protein) is found in the urine of about 75% of patients with multiple myeloma. Approximately 75% of patients with Waldenström's macroglobulinemia have a monoclonal light chain in the urine. Heavy-chain fragments as well as free light chains may be seen in the urine of patients with multiple myeloma or amyloidosis.

Interventions
Pretest Patient Preparation
1. Explain test purpose and specimen collection procedure.
2. Submit the same specimen for the serum protein electrophoresis, if a blood sample is needed. If the test is to be performed separately, another 7-mL blood sample collected in a red-topped tube is required. Note patient’s age; this procedure is seldom indicated in patients <30 years of age because monoclonal proteins are rarely identified in this age group.
3. Be aware that a 24-hour urine specimen is preferred. Provide instructions and a 24-hour collection container (see Chap. 3, Urine Studies, for protocols).
4. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcomes and monitor appropriately for neoplasms, infection, and liver and connective tissue disease.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Cold Agglutinin

This test most commonly diagnoses primary atypical viral pneumonia caused by Mycoplasma pneumoniae; it is used to diagnose certain hemolytic anemias (eg, cold agglutination disease) as well. The diagnosis depends on demonstrating a fourfold or higher increase in antibody titers between an early acute-phase blood serum sample and a blood serum sample taken in the convalescence phase, 7 to 10 days after the first sample. Positive reaction frequency and titer elevation both appear to be directly related to infection severity.

Patient's serum is serially diluted, human red cells are added, and the test is incubated at 4°C (refrigerator, 0°–10°C). The cold agglutinin antibodies react optimally at 4°C with the I antigen present on human red cells. The reaction is reversed by incubation of the agglutinated serum/cell mixture at 37°C.

Reference Values
Normal <1:16 by red cell agglutination at 4°C

Procedure
1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. Place specimen in a biohazard bag for transport to the laboratory.
2. Be aware that the sample should be prewarmed to 37°C for at least 15 minutes before the serum is separated from the cells. This allows the cold agglutinating antibodies to be eluted from the red cell membranes so that they can be detected in the agglutination procedure using O-negative indicator cells (pooled group O donors).

Clinical Implications
1. In viral pneumonia, the titer rises 8 to 10 days after onset, peaks in 12 to 25 days, and decreases 30 days after onset. Up to 90% of people with severe illness exhibit positive titers.

2. Chronic increased titer levels are associated with the following conditions:
   a. Cold antibody hemolytic anemia
   b. Chronic cold agglutinin disease
   c. Paroxysmal cold hemoglobinuria
   d. Severe Raynaud's phenomenon (may lead to gangrene)
   e. B-cell chronic lymphocytic leukemia

3. More important than any single high value is the rise in titer during the course of illness. The titer usually decreases by 4 to 6 weeks after the onset of illness.

4. Transient increases in titers are associated with primary atypical viral pneumonia, infectious mononucleosis, measles, mumps, CMV, congenital syphilis, hepatic cirrhosis, and trypanosomiasis.

Interfering Factors

1. A high cold agglutinin titer interferes with blood typing and crossmatching.
2. High titers are sometimes spontaneous in older persons and may persist for years.
3. Antibiotic therapy may interfere with cold agglutinin development.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results. Refer to page 532 for interpretation of immunologic test results. Counsel appropriately. Cold agglutinin titers rise during the second and third week of illness before rapidly returning to baseline levels. The test should be repeated at appropriate intervals.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Cryoglobulin Test

Cryoglobulins are proteins that reversibly precipitate or gel at 0° to 4°C. They are classified as follows:

1. Type I (monoclonal)
2. Type II (mixed cryoglobulins, in which a monoclonal is directed against a polyclonal immunoglobulin)
3. Type III (polyclonal, of which no monoclonal protein is found)

Types I and II are associated with monoclonal gammopathies, a group of diseases (see Immunofixation Electrophoresis, pages 578–579) in which a monoclonal protein is produced by neoplastic plasma cells or lymphocytes. Types II and III cryoglobulins are circulating immune complexes produced in response to a variety of antigens, including viral, bacterial, and autologous antigens.

The normal proteins of serum do not precipitate in the cold. Blood should be collected, allowed to clot, and centrifuged at 37°C. The serum should be separated at 37°C to ensure that the cryoglobulins will remain in the serum. The serum is then refrigerated and checked each day (up to 7 days) for the presence of a white precipitate or gel. Warming the serum to 37°C will reverse the precipitation.

The amount of cryoglobulin present can be quantified by filling a hematocrit tube with serum, incubating at 1°C, centrifuging at 1°C at 750 g for 30 minutes, and reading the cryocrit.

To characterize the cryoprotein, the precipitate is washed (cold saline) and redissolved (warm saline). IFE, as described on page 578, will identify the immunoglobulin classes present.

Reference Values

Normal Negative for cryoglobulin If positive after 3 to 7 days at 4°C, IFE on the cryoprecipitate is performed to identify the protein complex.

Procedure

1. Collect a 15-mL blood serum sample in a red-topped tube. Observe standard precautions. Keep the specimen at 37°C until the cells are separated.
2. Refrigerate the serum for a minimum of 72 hours, although 7 days is better to determine the presence of a cryoglobulin.

Clinical Implications The tendency of cryoglobulins to precipitate at low temperatures may occlude blood vessels; symptoms include Raynaud's phenomenon, vascular purpura, bleeding tendencies, cold-induced urticaria, pain, and cyanosis. Type I cryoglobulinemia is associated with monoclonal gammopathy of undetermined significance (MGUS), macroglobulinemia, or multiple myeloma. Type II cryoglobulinemia is associated with autoimmune disorders such as vasculitis, glomerulonephritis, systemic lupus erythematosus, rheumatoid arthritis, and Sjögren's syndrome. It may also be seen in diseases such as hepatitis, infectious mononucleosis, CMV, and toxoplasmosis. Type III cryoglobulinemia is usually associated with the same disease spectrum as type II and may take the full 7 days to appear. A cryoprecipitate in
plasma but not serum is caused by cryofibrinogen. Cryofibrinogens are rare and can be associated with vasculitis.

**Clinical Alert**

The presence of cryoglobulins may cause erroneous results with some automated hematology instruments.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results. Counsel and monitor appropriately for infections, collagen disorders, and malignant blood cell disease. Follow-up testing is usually needed.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Total Hemolytic Complement (CH50)**

Complement (C) is a complex sequential cascade system in which inactive proteins become active and interact much like the clotting system. The complement system is important as part of the body's defense mechanism against infection. Activation of complement results in cell lysis, release of histamine from mast cells and platelets, increased vascular permeability, contraction of smooth muscle, and chemotaxis of leukocytes and neutralization of certain viruses. These inactive proteins constitute about 10% of the globulins in normal blood serum. The complement system is also interrelated with the coagulation, fibrinolytic, and kinin systems. The action of complement, however, is not always beneficial. The potent reactions mediated by this complex system are not always contained. In the presence of gram-negative bacteremia, the complement can escape its built-in control mechanisms, causing severe damage to the body. It is not clear how this happens, but it is known that complement abnormalities develop before shock occurs.

This test screens for certain autoimmune diseases, estimates the extent of immune complex formation, and detects all inherited and most acquired immune deficiencies. Serial measurements monitor disease course and treatment in SLE, RA, and glomerulonephritis. It is a useful adjunct for rheumatoid factor and antinuclear antibody (ANA) testing when immune complexes appear to be the primary mediators of tissue injury.

**Reference Values**

*Normal* 60–144 complement activity enzyme (CAE) Units by ELISA

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. A joint fluid specimen of at least 1 mL can also be used and should be collected in a tube that does not contain additives.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Alert**

Complement deteriorates at room temperature in serum or fluid; samples should be brought to the laboratory as soon as possible. Separate serum from clot and freeze at -70°C until test is performed. Both blood and fluid must be processed and frozen within 2 hours after specimen collection. Failure to process the specimen in this manner may lead to falsely decreased functional activity levels.

**Clinical Implications**

1. *Increased total complement values* are associated with most inflammatory responses; these acquired elevations are usually transient, and concentrations return to normal when the situation is resolved.
2. *Decreased total complement values* are associated with hereditary defects of specific complement components. In C2 deficiency, autoimmune disorders occur as SLE, and C1q deficiency may cause hereditary angioedema.
   a. Complement consumption by activation of the alternative pathway, an amplification of the classic pathway not requiring an "immunologic" stimulus, can be seen in the following conditions:
      1. Gram-negative septicemia
      2. Subacute bacterial endocarditis
      3. Acute poststreptococcal glomerulonephritis
      4. Membranoproliferative glomerulonephritis
   b. Complement consumption due to activation of the classic pathway by immune complex formation occurs in the following conditions:
      1. SLE
      2. Serum sickness
      3. Acute vasculitis
      4. Severe RA
      5. Hepatitis
      6. Cryoglobulinemia

**Interventions** See ANAs, page 586.

**C3 Complement Component**

C3 constitutes 70% of the total protein in the complement system and is essential to the activation of both classic and
alternative pathways. Along with the other components of the complement system, C3 may be used up in reactions that occur in some antigen-antibody reactions. C3 is synthesized in the liver, macrophages, fibroblasts, lymphoid cells, and skin.

This test is done when it is suspected that individual complement component concentrations are abnormally reduced. This test and the C1q and C4 tests are the most frequently ordered complement measurements. There is a correlation between most forms of nephritis, the degree of nephritis severity, and C3 levels. C3 is useful for assessing disease activity in SLE.

**Reference Values**

**Normal** 75–175 mg/dL (0.75–1.75 g/L) by rate nephelometry

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. This amount is sufficient for both C3 and C4 testing.
2. Place specimen in biohazard bag for transport to laboratory.

**Clinical Implications**

1. **Decreased C3 levels** are associated with most active diseases with immune complex formation.
   a. Severe recurrent bacterial infections due to C3 homozygous deficiency
   b. Absence of C3b inactivator factor
   c. Acute poststreptococcal glomerulonephritis
   d. Immune complex disease
   e. Active SLE
   f. Membranoproliferative glomerulonephritis
   g. Nephritis
   h. End-stage liver disease
2. **Increased levels** are found in numerous inflammatory states.

**Interventions** See ANAs, page 586.

**Clinical Alert**

Patients with low C3 levels are in danger of shock leading to death.

**C4 Complement Component**

C4 is another of the components of the complement system and is synthesized in bone and lung tissue. C4 may be bypassed in the alternative complement pathway when immune complexes are not involved, or it may be used up in the very complicated series of reactions that follow many antigen-antibody reactions.

This is a follow-up test done when total complement levels are abnormally decreased. It can also be ordered to confirm hereditary angioedema if the C1 inhibitor result is decreased.

**Reference Values**

**Normal** 14–40 mg/dL (140–400 mg/L) by rate nephelometry

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. This amount is sufficient for both C3 and C4 testing.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**

1. **Decreased C4 levels** are associated with the following conditions:
   a. Acute SLE
   b. Early glomerulonephritis
   c. Immune complex disease
   d. Cryoglobulinemia
   e. Inborn C4 deficiency
   f. Hereditary angioneurotic edema
2. **Increased C4 levels** are associated with malignancies.

**Interventions** See ANAs, page 586.

**C’1 Esterase Inhibitor (C’1 INH)**

C’1 esterase inhibitor is a glycoprotein. It acts as a regulatory brake on the complement activation process. Decreased production of this glycoprotein results in hereditary angioedema (HAE).

This determination is an important tool for diagnosing HAE, a disorder caused by a low concentration of C’1 esterase inhibitor or by an abnormal structure of the protein. Affected persons are apparently heterozygous for the condition. It is
also used in the differential diagnosis of the more prevalent but less serious allergic and nonfamilial angioedema.

Reference Values

Normal 18–40 mg/dL (180–400 mg/L) by an immunoturbidimetric assay

Procedure

1. Collect a 7-mL serum specimen in a red-topped tube. Observe standard precautions. Place specimen in biohazard bag for transport to laboratory.
2. Spin down, separate from clot, and freeze 1.0 mL of serum at -70°C until testing is performed.

Clinical Implications

1. Decreased values are associated with HAE, a genetic disease characterized by acute edema of subcutaneous tissue, gastrointestinal tract, or upper airway tract.
2. During acute attacks of the disease, C4 and C2 components can be markedly reduced.

Interventions

See ANAs, page 586.

Clinical Alert

Prednisolone and transfusions of fresh-frozen plasma have been successfully used to treat HAE.

TESTS FOR AUTOIMMUNITY AND SYSTEMIC RHEUMATIC DISEASE (SRD)

Antinuclear Antibody (ANA) Test

Measurement of ANAs in serum is the most commonly performed screening test for autoantibodies in patients suspected of having systemic rheumatic disease (SRD). SRDs are also called connective tissue or collagen diseases. Examples of SRDs include SLE, mixed connective tissue disease, Sjögren's syndrome, scleroderma, CREST (calcinosis, Raynaud's phenomenon, esophageal dysfunction, sclerodactyly, and telangiectasia) syndrome, rheumatoid arthritis, and polymyositis dermatomyositis.

The diagnosis of SLE is difficult because clinical signs and symptoms are varied and mimic other SRDs. SLE is characterized by the production of autoantibodies to nuclear antigens, that is, anti-dsDNA. SLE is a multisystem disease that can affect every organ system in the body, especially the kidneys.

Results of tests for ANAs by ELISA show that ELISA and traditional indirect immunofluorescence methods for ANA are substantially equivalent. Many laboratories are using a combination of both methods. ANA samples are screened using an ELISA assay. All samples that screen positive or equivocal are titered using Hep-2 cells, and the titer and pattern are reported. In general, a titer =1:160 is considered a significant positive. Low-titer ANAs are common with advancing age. When cell culture substrates (Hep-2 cells) are used, the ANA incidence is 99% in SLE.

Reference Values

Normal Negative by ELISA and IFA methods. If positive by IFA, the specimen is titered and a pattern is reported.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in biohazard bag for transport to the laboratory.

Clinical Implications

1. A positive result does not confirm a disease; low titers of ANAs are present in elderly people and some apparently healthy normal persons.
2. The diagnosis of an SRD is based primarily on the presence of compatible clinical signs and symptoms. The results of tests for autoantibodies, including ANAs and specific autoantibodies (eg, RNP, Smith, SSA, SSB, Scl-70, Jo-1) are ancillary. Additional diagnostic criteria include consistent histopathology or specific radiographic findings.

Interfering Factors

1. Drugs, such as procainamide and hydralazine, may cause a positive ANA result.
2. Positive ANA levels may be found after viral illnesses and with some chronic infections.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure. A strong positive result, that is, >3 on ELISA or =1:160 on IFA, may require follow-up testing of specific autoantibodies.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results. Monitor and counsel patient about follow-up tests and treatment.
2. SRDs, such as SLE, must be dealt with on a continuing basis and may require certain lifestyle changes. Repeat testing evaluates the effectiveness of therapy. Minor symptoms, in the absence of major organ involvement, are
frequently treated with nonsteroidal anti-inflammatory drugs (NSAIDs) such as salicylates. Cutaneous manifestations respond to topical corticosteroid treatments. Short-acting corticosteroids, such as prednisone, are necessary if acute serologic changes and severe clinical manifestations appear.

3. Long-term moderate- to high-dose corticosteroids are central regimens prescribed for diffuse glomerulonephritis as well as rheumatoid arthritis.

4. Corticosteroid dosage may be reduced and renal disease favorably managed by adding immunosuppressive drugs to the therapy regimen. Infection, secondary to immunosuppressive treatment, is a leading cause of death in patients with SRDs. Patient education plays a major role in prevention of infection.

5. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Anticentromere Antibody Test

The variant of scleroderma, the CREST syndrome, is characterized by calcinosis, Raynaud's phenomenon, esophageal dysfunction, sclerodactyly, and telangiectasia. Characteristically, anticentromere antibodies appear in about 90% of these patients. This antibody is detected by using Hep-2 cells in various stages of cell division. The centromere region of the cell chromosomes will stain if an anticentromere antibody is present.

Reference Values

Normal Negative for anticentromere antibody by IFA or ELISA. If positive by IFA, the serum is titered.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in biohazard bag for transport to laboratory.

Clinical Implications Positive results are associated with the CREST syndrome in scleroderma.

Interventions See ANAs, page 586.

Anti-dsDNA Antibody Test, IgG

Although not completely understood, the primary mechanism of tissue injury in SLE and related autoimmune disease is the formation of antigen-antibody immune complexes. Not all ANAs are pathogenic. For the few that are harmful, pathogenicity depends on the specific immunoglobulin class, ability to activate complement, size of the immune complex, and site of tissue deposition. For example, studies of immune complex–mediated tissue injury in the kidney have shown a clear relation between deposition of immune complexes and glomerular disease.

The anti-dsDNA test is done specifically to identify or differentiate native (ie, double-stranded) DNA antibodies, found in 40% to 60% of patients with SLE during the active phase of their disease, from other nonnative DNA antibodies found in other rheumatic diseases. The presence of antibodies to dsDNA generally correlates with lupus nephritis. An anti-dsDNA test supports a diagnosis, allows monitoring of disease activity and response to therapy, and establishes a prognosis for SLE.

Reference Values


Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to laboratory.

Clinical Implications

1. Anti-dsDNA concentrations may decrease with successful therapy and may increase with an acute recurrence of SLE.
2. DNA–anti-dsDNA immune complexes play a role in SLE pathogenesis through the deposit of these complexes in the kidney and other tissues.

Interfering Factors

1. The Farr assay, an RIA method, detects both single-stranded (ss) and double-stranded (ds) DNA antibodies.
2. Antibodies to ssDNA are nonspecific but are associated with various other rheumatic diseases.

Interventions See ANAs, page 586.

Rheumatoid Factor (Rheumatoid Arthritis [RA] Factor) Test

The blood of many persons with RA contains a macroglobulin-type antibody called rheumatoid factor (RF). Evidence indicates that rheumatoid factors are anti–gamma-globulin antibodies; however, until a specific antigen that produces RF is discovered, the exact nature of RF can only be speculated. Even more uncertain is the role that RF plays in RA. Although RF may cause or perpetuate the destructive changes associated with RA, it may also be incidental to these changes or may even serve some beneficial purpose. RF is sometimes found in blood serum from patients with other diseases, even though RF incidence and values are higher in patients with RA.

This test is useful in the diagnosis of RA. It measures RFs (antibodies directed against the Fc fragment of IgG). These are usually IgM antibodies, but they may also be IgG or IgA. Four of the following clinical criteria must be present to
diagnose rheumatoid arthritis.

Revised American College of Rheumatology Criteria for Rheumatoid Arthritis

1. Morning stiffness for at least 6 weeks
2. Pain on motion or tenderness in at least one joint for at least 6 weeks
3. Swelling in at least one joint for at least 6 weeks
4. Swelling in at least one other joint for at least 6 weeks
5. Symmetrical joint swelling with simultaneous involvement of the same joint on both sides of the body
6. Subcutaneous nodules
7. X-ray changes, including bony decalcification

Reference Values
Normal 0–20 U/mL or 0–20 kU/L, based on rate nephelometry

Procedure
1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in biohazard bag for transport to laboratory.

Clinical Implications
1. When a patient who tests positive improves, subsequent tests also remain positive unless titers were initially low.
2. A positive RF test result often supports a tentative diagnosis of early-onset RA (eg, versus rheumatic fever).
3. RFs frequently occur in a variety of other diseases, such as SLE, endocarditis, tuberculosis, syphilis, sarcoidosis, cancer, viral infections, Sjögren's syndrome, and diseases affecting the liver, lung, or kidney as well as in patients who have received skin and renal allografts.
4. Absence of RF does not exclude the diagnosis or existence of RA.

Interfering Factors The result is normally higher in older patients and in those who have received multiple vaccinations and transfusions.

Interventions See ANAs, page 586.

Antibodies to Extractable Nuclear Antigens (ENAs): Anti-Ribonucleoprotein (RNP); Anti-Smith (Sm);
Anti-Sjögren's Syndrome (SSA, SSB); Anti-Scleroderma (Scl-70); Anti-Jo-1 (Jo-1)

The extractable nuclear antigens (ENAs), another group of nuclear antigens (nonhistone proteins) to which autoantibodies may develop, are so named because of their presence in saline solution extracts of certain nonhuman cells. The most common ENAs are ribonucleoprotein (RNP) and Smith (Sm).

Anti-RNP is elevated in 35% to 40% of SLE patients and in patients with other connective tissue diseases, notably mixed connective tissue disease (MCTD). MCTD is characterized by high levels of anti-RNP without autoantibodies to dsDNA or Sm. The disease resembles SLE but is not accompanied by renal involvement.

Anti-Sm is specific for SLE but occurs in only approximately 30% of the patients. The levels of anti-Sm may be related to disease activity in SLE.

Anti-SSA (Ro) has been detected in approximately 25% of patients with SLE and in 40% to 45% of patients with Sjögren's syndrome.

Anti-SSB (La) is found in approximately 10% to 15% of patients with SLE and up to 60% of patients with Sjögren's syndrome.

Anti-Scl-70 is considered specific for scleroderma (systemic sclerosis). These autoantibodies are found in up to 60% of scleroderma patients with extensive cutaneous disease and interstitial pulmonary fibrosis.

Anti-Jo-1 occurs in approximately 20% of patients with myositis, usually in patients with interstitial pulmonary fibrosis and symmetrical polymyositis.

The ELISA assay is a screen for several nuclear antibodies. If the ENA screen result is borderline or positive, the following tests (Table 8.8) will be set up to determine the particular systemic rheumatic disease SRD.

| Table 8.8 ELISA Screening for Specific Systemic Rheumatic Disease (SRD) |
|----------------|----------------------------|
| Test           | Specific SRD               |
| Anti-RNP       | Mixed connective tissue disease (MCTD) |
| Anti-Sm (with or without RNP) | SLE, MCTD, Sjögren's syndrome |
| Anti-SSA(Ro) and/or Anti-SSB(La) | Sjögren's syndrome |
| Anti-Scl-70    | Scleroderma                |
| Anti-Jo-1      | Polymyositis               |
Reference Values for ENA and Individual Autoantibody Tests

**Normal**
- Negative: <20 U by ELISA
- Borderline: 20–25 U
- Positive: >26 U

**Procedure**
1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to laboratory.

**Clinical Implications**
1. Results of serum tests for autoantibodies should not be relied on extensively to establish the diagnosis of a connective tissue disease. They must always be interpreted in conjunction with clinical findings.
2. Testing for autoantibodies to ENAs is not useful in patients without demonstrable antinuclear antibodies.

**Interventions** See ANAs, page 586.

**Cardiolipin Antibodies, IgA, IgG, IgM**

In patients with SLE, antibodies to cardiolipin (a negatively charged phospholipid) have been associated with both arterial and venous thrombosis, thrombocytopenia and recurrent fetal loss. Patients with the anticardiolipin syndrome have one of the above clinical features and have antibodies to cardiolipin and/or a positive lupus anticoagulant test.

The antibodies present to cardiolipin may be of the IgA, IgG, IgM isotypes. Testing for the various antibody isotypes to cardiolipin aid in the diagnosis of the antiphospholipid syndrome in patients with SLE or lupus-like disorders. They are also useful for the prognostic assessment of pregnant patients with a history of recurrent fetal loss.

**Reference Values**
- Normal: <12 APL (IgA phospholipid units): absent or none detected
- <15 GPL (IgG phospholipid units): absent or none detected
- <12 MPL (IgM phospholipid units): absent or none detected

**Procedure**
1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**
1. Most patients with antiphospholipid antibody syndrome have moderate or high levels of cardiolipin antibodies and are positive for IgG only or IgG and IgM.
2. Elevated values are seen in spontaneous thrombosis and in patients with connective tissue disease.
3. Patients with current or prior syphilis infection may have a false-positive result without the risk for thrombosis.

**Interventions**

*Pretest Patient Preparation*
1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

*Posttest Patient Aftercare*
1. Interpret test outcomes in light of the patient's history, physical findings, and other diagnostic procedures and results. Temporarily positive tests do occur for IgG and IgM antibodies, and it is recommended that positive results be confirmed by follow-up assay in 8 weeks.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Autoimmune Thyroiditis, Thyroid Antibody Tests: Thyroglobulin Antibody, Thyroid Microsomal Antibody, Thyroperoxidase Antibody**

There are several autoantibodies that are organ specific for the thyroid gland, but anti-thyroglobulin and anti-thyroperoxidase are ordered most frequently by clinicians when evaluating patients for hyperthyroidism, hypothyroidism, and thyroid cancer. In Graves' disease, which is autoimmune hyperthyroidism, and in Hashimoto's thyroiditis, which is autoimmune hypothyroidism, the presence of both antibodies can help confirm the diagnosis.

Thyroglobulin antibodies are directed against the glycoprotein thyroglobulin located in the thyroid follicles; thyroperoxidase, against the membrane-bound glycoprotein thyroperoxidase located in the cytoplasm of the epithelial cells surrounding the follicles.

Along with chemiluminescence technology has come the use of highly purified antigens to improve specificity. For the anti-thyroperoxidase test, instead of using the entire microsomal antigen, this assay uses just the thyroperoxidase (TPO) component. TPO is considered the primary autoantigenic component of the microsomal antigen. Test systems that use the purified TPO (in place of the microsomal antigen) have greater specificity for the clinically significant autoantibody. Assays using microsomal antigen are detecting TPO antibody but may also detect antibodies to other parts of the microsomal antigen that have little or no clinical significance.

**Reference Values**
- Normal: <1:100 for thyroglobulin and thyroid microsomal antibodies by hemagglutination
- Negative for thyroglobulin and
thyroid microsomal antibodies by ELISA. Negative for thyroglobulin and thyroperoxidase antibodies by chemiluminescence.

Procedure

1. Collect 7-mL blood serum in a red-topped tube. Observe standard precautions.
2. Place specimen in biohazard bag for transport to laboratory.

Clinical Implications

1. High titers of thyroglobulin and thyroid microsomal antibodies (>1:400) are found in Hashimoto's disease, but elevations can also be seen in other autoimmune diseases.
2. Increased thyroid antibodies also occur in the following conditions:
   a. Graves' disease
   b. Thyroid carcinoma
   c. Idiopathic myxedema
   d. Pernicious anemia
   e. SLE, RA, Sjögren's syndrome
   f. Subacute thyroiditis
   g. Nontoxic nodular goiter

Interfering Factors

1. About 10% of the normal population may have low levels of thyroid antibodies with no symptoms of the disease. Incidence of low titer is higher in women and increases with age.
2. Antibody production may be confined to lymphocytes within the thyroid, resulting in negative serum test results.

Interventions

Pretest Patient Preparation

1. Explain test purpose. Thyroid antibody testing is done to confirm diagnosis. It is not to be relied on, however, for management of the disease.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and determine the need for possible follow-up testing. The diagnosis of autoimmune thyroiditis is made on the basis of clinical observations, thyroid function tests (see Chap. 6), and the presence of circulating autoantibodies, such as thyroglobulin, microsomal (TPO).
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

AUTOIMMUNE LIVER DISEASE TESTS

Anti–Smooth Muscle Antibody (ASMA) Test

ASMA is associated with liver and bile duct autoimmune diseases. The immune response itself is believed to be responsible for the disease process.

Sera from patients with autoimmune chronic active hepatitis (CAH) contain antibodies to smooth muscle antigens that are detectable by IFA on tissues that contain smooth muscle, such as mouse stomach. The antibodies are predominantly IgG. This measurement differentiates CAH and primary biliary cirrhosis from other liver diseases in which ASMs are seldom present (e.g., SLE).

Reference Values

Normal Negative by IFA if positive, serum is titered.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. This amount is sufficient for both ASMA and antimitochondrial antibody (AMA) testing.
2. Place specimen in biohazard bag for transport to the laboratory.

Clinical Implications

1. ASMAs are found in chronic active hepatitis, a progressive disease of unknown etiology found predominantly in young women. It has factors characteristic of both acute and chronic hepatitis (80% of patients). If this disease is associated with a positive ANA test, the disease is often called lupoid hepatitis.
2. Antibody titers between 80 and 320 occur commonly in patients with CAH.

Prevalence of Autoantibodies in Liver Disease

<table>
<thead>
<tr>
<th>Disease</th>
<th>Anti–Smooth Muscle (%)</th>
<th>Antimitochondrial (%)</th>
<th>ANA (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chronic active hepatitis</td>
<td>70–90</td>
<td>30–60</td>
<td>60</td>
</tr>
<tr>
<td>Chronic persistent hepatitis</td>
<td>45</td>
<td>15–20</td>
<td>15–30</td>
</tr>
</tbody>
</table>
### Interventions

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and monitor appropriately. Detection of ASMA by immunofluorescence assists in determining the presence of chronic active hepatitis and need for therapy when used in conjunction with other laboratory tests such as those used to evaluate liver enzymes, ANAs, and IgG levels. All of these are elevated in the majority of patients with chronic active hepatitis.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

### Antimitochondrial Antibody (AMA) Test

AMA is non–organ and non–species specific and is directed against a lipoprotein in the inner mitochondrial membrane. AMA is a marker for primary biliary cirrhosis, a chronic inflammatory liver disease, characterized by the progressive destruction of interlobular bile ducts with development of cholestasis and eventually cirrhosis.

The mitochondrial antigens recognized by AMAs in patients’ sera have been classified as M1 through M9, with M2 recognized by AMAs in 99% of patients with primary biliary cirrhosis. The antibodies are predominantly IgG. This measurement aids in the diagnosis of primary biliary cirrhosis (PBC), a progressive disease most commonly seen in women in the second half of their reproductive years. There is also a genetic predisposition.

**Reference Values**

**Normal** Negative by indirect fluorescent antibody (IFA). If positive, serum is titered.

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions. This amount is sufficient for both AMA and ASMA testing.
2. Place specimen in biohazard bag for transport to laboratory.

**Clinical Implications**

1. Elevated concentrations of AMAs are present in >80% of patients with PBC.
2. High titers are also associated with longstanding hepatic obstruction, chronic hepatitis, and cryptogenic cirrhosis.
3. Elevated levels are occasionally present in the following conditions:
   a. SLE
   b. RA
   c. Thyroid disease
   d. Pernicious anemia
   e. Idiopathic Addison's disease

### Interventions

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and monitor appropriately. Immunofluorescence testing, along with quantitation of IgM and liver enzymes, both of which tend to be elevated in PBC, are reliable follow-up protocols.
2. Follow guidelines in Chapter 1 regarding safe, effective informed posttest care.

### Anti–Liver/Kidney Microsome Type 1 Antibody (LKM) Test

Antibodies to liver/kidney microsome antigens (anti–LKM-1) occur in a subset of patients with chronic autoimmune hepatitis (AIH). The clinical diagnosis of AIH is difficult because there are no particular signs, symptoms, or liver function test abnormalities that are specific enough to be considered diagnostic. Patients with this type of chronic AIH are predominantly children, but some patients are adults.

Different autoantibodies are found in the serum from patients with AIH. The discovery of the LKM-1 antibody led to the establishment of two subtypes of AIH. The percentage of type 2 AIH patients whose serum contains LKM-1 antibodies is...
90%.

Patient sera are incubated on slides with mouse kidney and stomach. Anti–LKM-1 antibody produces a characteristic pattern, which allows it to be differentiated from the patterns produced by smooth muscle and mitochondrial antibodies on mouse tissue.

Reference Values
Normal Negative for LKM-1 by indirect fluorescent antibody (IFA) If positive, serum is titered.

Procedure
1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

Interventions

Pretest Patient Preparation
1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcomes and monitor appropriately. The primary therapy for AIH is administration of corticosteroids. Steroid treatment should lead to rapid reduction in AST and ALT (liver enzyme) levels.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Antiparietal Cell Antibody (APCA) Test

The disruption of normal intrinsic factor production or function due to autoimmune processes can lead to pernicious anemia. Antibodies to two antigens of the gastric parietal cell—antiparietal cell antibodies (APCAs) and intrinsic factor antibodies—are found in pernicious anemia.

This measurement is helpful in diagnosing chronic gastric disease and differentiating autoimmune pernicious anemia from other megaloblastic anemias. Persons with other anemias do not have detectable APCAs.

Reference Values
Normal Negative for APCA by IFA If positive, serum is titered.

Procedure
1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in biohazard bag for transport to the laboratory.

Clinical Implications
1. APCAs occur in >80% of patients with autoimmune pernicious anemia; 50% have antibodies to intrinsic factor.
2. Occasionally, APCAs are present in the following conditions:
   a. Gastric ulcer
   b. Gastric cancer
   c. Atrophic gastritis
   d. Thyroid disease
   e. Diabetes mellitus
   f. Iron-deficiency anemia

Interfering Factors APCAs are present in many healthy adults >60 years of age.

Interventions

Pretest Patient Preparation
1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcomes and monitor appropriately. Detection of APCA may suggest need for more invasive testing, such as gastric biopsy to rule out gastrointestinal disease.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Antiglomerular Basement Membrane (AGBM) Antibody Test

Antibodies specific for renal structural components such as the glomerular basement membrane of the kidney can bind to respective tissue-fixed antigens to produce an immune response.

This test is primarily used in differentiating glomerular nephritis induced by antiglomerular basement membrane antibodies (AGBMs) from other types of glomerular nephritis. AGBMs cause about 5% of glomerular nephritis; about two thirds of these patients may also develop pulmonary hemorrhage (Goodpasture's syndrome).
Reference Values

**Normal** Negative: <5 EU/mL by ELISA Borderline: 5.1–20.0 EU/mL Positive: 20.1–400 EU/mL

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications** AGBM antibodies are detected in the following conditions:

1. AGBM glomerular nephritis
2. Tubulointerstitial nephritis
3. AGBM Goodpasture’s syndrome
4. Some patients with SLE

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and need for follow-up testing and treatments that involve immunosuppressants and plasmapheresis, which are effective if treatment is started before renal failure is well advanced.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Acetylcholine Receptor (AChR) Binding Antibody Test**

Acetylcholine receptor antibodies (AChRs) appear in myasthenia gravis (MG). It is believed that this disease involves destruction by the muscle cells of acetylcholine receptors bound by antibodies at the skeletal muscle motor endplate.

This measurement is considered to be the first-order test for MG in symptomatic patients. It also helps in managing response to immunosuppressive therapy. Second- and third-order tests for modulating and blocking antibodies, respectively, are ordered to confirm the diagnosis of acquired MG, distinguish acquired disease from congenital disease, and monitor the serologic process in the course of MG.

Reference Values

**Normal** Negative for AChR or <0.02 nmol/L by RIA

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**

1. AChR antibodies are found in about 90% of persons with generalized MG, 70% of persons with ocular MG, and 80% of persons in remission. These findings confirm the autoimmune nature of the disease.
2. Patients who have only eye symptoms tend to have lower titers than those with generalized myasthenia symptoms.

**Interfering Factors** Positive results can be found in patients with Lambert-Eaton myasthenic syndrome (LES) or autoimmune liver disease.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose. Assess for history of immunosuppressive drug treatment. Detection of acetylcholine receptor binding antibody is infrequent in such cases.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and possible need for other testing. Other tests now available to aid in the serologic diagnosis of MG include acetylcholine receptor blocking antibody, acetylcholine receptor modulating antibody, and striational antibodies. These are ordered according to presentation of neurologic symptoms. All of these antibodies are less frequently detected in the early stages of MG (within 1 year of onset) and in patients treated with immunosuppressive drugs. None are found in cases of congenital MG.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Anti-Insulin Antibody Test**

Persons with diabetes may form antibodies to the insulin they take and require larger doses because insulin is not available for glucose metabolism when it is partially complexed with these antibodies. Insulin antibodies are immunoglobulins called anti-insulin antibodies; they act as insulin-transporting proteins. The most common type of anti-insulin antibody is IgG, but it is found in all five classes of immunoglobulins in insulin-treated patients. These
immunoglobulins, especially IgE, may be responsible for allergic manifestations: IgM may cause insulin resistance.

This insulin antibody level provides information for determining the most appropriate treatment for certain diabetic patients. It may focus the reason for allergic manifestations. It can identify a state of insulin resistance in which the daily insulin requirement exceeds 200 U for more than 2 days, and may be associated with elevated anti-insulin antibody titers and insulin-binding capacity.

Reference Values

Normal Negative: <3% binding of the patient's serum with labeled beef, human, and pork insulin SI units when performed by RIA

Procedure

2. Place specimen in biohazard bag for transport to laboratory.

Clinical Implications Anti-insulin antibody elevations are associated with insulin resistance and allergies to insulin.

Interventions

Pretest Patient Preparation

1. Explain purpose of test. Fasting is required. Check with individual laboratory for time frames.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes. Based on antibody levels present and clinical findings, the dosage of insulin is changed to reduce or prevent further allergic manifestations and/or insulin resistance.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Gliadin Antibodies, IgA and IgG

Antibodies to gliadin (wheat protein) have been shown conclusively to be the toxic agent in celiac disease. Originally, a series of multiple intestinal biopsies were required to diagnose celiac and related intestinal diseases. More recently, serologic testing has been strongly suggested for screening patients with suspected gluten-sensitive enteropathy as well as for monitoring dietary compliance.

Celiac disease usually begins in infancy soon after introduction of cereals to the diet, but symptoms may disappear spontaneously in later childhood, despite continued signs of malabsorption. Strict avoidance of gluten in the diet is recommended to control the disease.

Both IgG and IgA gliadin antibodies are detected in sera of patients with gluten-sensitive enteropathy. IgG antigliadin antibodies seem more sensitive but are less specific than the IgA class antibodies. The best strategy for at-risk populations includes testing for both classes of gliadin antibodies.

Reference Values

Normal Values are given for >2 years of age. Negative: <25 U/mL by ELISA Weakly positive: 25–50 U/mL Positive: >50 U/mL

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to laboratory.

Clinical Implications

1. The gliadin antibody assay has a sensitivity of 95% for active, untreated celiac patients when both IgG and IgA are used. The test has an overall specificity of 90%.
2. A negative IgA result in an untreated patient does not rule out gluten-sensitive enteropathy, especially when associated with elevated levels of IgG gliadin antibodies.
3. Significant portions of celiac patients are IgA deficient, which can serve as an explanation for this occurrence.
4. In treated patients known to express IgA antibodies, the IgA gliadin antibody level represents a better indicator of dietary compliance than the IgG level.
5. False-positive results (high antibody levels without the corresponding histologic features) are possible; other gastrointestinal disorders, especially Crohn's disease, postinfection malabsorption, and food protein intolerance (e.g., cow's milk), are known to induce circulating antigliadin antibodies.
6. Results of this assay should be used in conjunction with clinical findings and other serologic tests.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcome in light of patient's dietary history, including related clinical, laboratory, and histologic data. Positive results are possible in patients with other gastrointestinal disorders.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Antineutrophil Cytoplasmic Antibodies (ANCA)

There are two types of antineutrophil cytoplasmic antibodies distinguished by different immunofluorescent staining patterns using human neutrophil substrates:

1. cANCA produces a diffuse cytoplasmic staining of neutrophils and monocytes and is specific for proteinase 3. cANCA is found in the sera of patients with Wegner's granulomatosis (WG).
2. pANCA produces a perinuclear staining of neutrophils and are specific for other neutrophil enzymes including myeloperoxidase (MPO), elastase, and lactoferrin. pANCA specific for MPO is found in the sera of patients with systemic vasculitis, most of whom have renal involvement characterized by pauci-immune necrotizing glomerulonephritis.

Tests for ANCA are performed by an indirect immunofluorescent technique. Slides prepared from neutrophils are used as a substrate to bind ANCA so that it can be detected microscopically. Depending on the pattern of staining, as mentioned previously, two types of ANCA exist: cANCA and pANCA.

Reference Values

Normal Negative for ANCAs by IFA. If positive for cANCA, results are titered. If positive for pANCA, MPO testing is performed by ELISA. Not all specimens positive for pANCA are MPO positive.

Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in biohazard bag for transport to the laboratory.

Clinical Implications

1. In patients with active generalized WG (pulmonary and/or renal involvement), the frequency of positive cANCA results approaches 85%. A negative test for cANCA does not rule out WG; however, false-positive results are rare.
2. In patients with known WG, rising titers of cANCA suggest relapse, and falling titers suggest successful treatment.
3. In patients with active renal disease, a positive pANCA suggests the presence of antibodies to MPO and pauci-immune necrotizing glomerulonephritis.
4. Results of tests for ANCA should be considered along with other clinical, laboratory, and histopathologic data in establishing the diagnosis of WG or systemic vasculitis.
5. Inflammatory bowel disease (IBD)-associated ANCAs are found in ulcerative colitis and Crohn's disease, specifically pANCA.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes in light of the patient's history, including other clinical, laboratory, and histopathologic data. Positive ANCA results (pANCA and, rarely, cANCA) may occur in patients with diseases other than WG or vasculitis, including Goodpasture's syndrome and SLE.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

SPERM ANTIBODIES

Antisperm Antibody Test

The majority of infertile males have blocking of the efferent testicular ducts. It is likely that, similar to vasectomy, reabsorption of sperm from blocked ducts results in the formation of autoantibodies to sperm.

Testing for sperm antibodies is not recommended for routine infertility testing. The following indications, however, might warrant ordering this evaluation:

1. Males
   a. Sperm agglutination in the ejaculate
   b. History of testicular trauma
   c. History of biopsy
   d. Vasectomy reversal
   e. Genital tract infection
   f. Obstructive lesions of the male ductal system

2. Females
   a. Abnormal postcoital test
   b. Unexplained infertility
Reference Values

Normal Normals display <20% total binding. Reported as percentage of sperm with positive binding by immunobead technique; >20% binding is usually required to lower patient's fertility. Significance of percentage of binding is inversely related to patient's sperm count.

Procedure A semen test sample is preferred for values. If semen procurement presents a problem for a male patient, a blood serum sample can be tested. For females, blood serum is preferred because of the difficulty of cervical mucus collection.

1. Blood
   a. Collect a 7-mL blood serum sample in a red-topped tube.
   b. Spin down and send 2.0 mL of serum to laboratory frozen in plastic vial on dry ice.

2. Semen
   a. Collect contents of semen ejaculate.
   b. Send specimen to laboratory frozen in plastic vial on dry ice.

3. Cervical mucus
   a. Collect 1.0 mL of cervical mucus.
   b. Send specimen to laboratory frozen in plastic vial on dry ice.

Clinical Implications Antisperm antibodies are associated with the following conditions:

1. Blocked testicular efferent ducts and the resultant resorption of sperm can produce antibodies.
2. After vasectomy, antibodies and probable cellular immunity to sperm develop in most males as a result of the interaction of sperm antigens with the immune system.
3. In some studies, about 75% of women with primary infertility had sperm agglutinins. However, 11% to 15% of pregnant women had the same sperm antibody titers.

Clinical Alert

The potential adverse consequences of an immune sperm response to sperm include possible systemic effects in other organ systems and possible infertility after vasectomy reversal.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure. See details under Procedure for “Specimen Required.” Patient should be advised of the need for repeat testing.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and counsel appropriately. It may be necessary to repeat this procedure on different sample types (eg, semen, blood) to establish a possible cause for infertility.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

ALLERGY TESTING

IgE Antibody, Single Allergen

A large number of substances have been found to have allergic potential. Measurements of IgE antibodies are useful to establish the presence of allergic diseases and to define the allergen specificity of immediate hypersensitivity reactions. Examples of allergic diseases include asthma, allergic rhinitis, dermatitis, anaphylaxis, and urticaria.

The fluorescent enzyme immunoassay (FEIA) tests measure the increase and quantity of allergen-specific IgE antibodies and diagnose an allergy to a specific allergen (eg, molds, weeds, foods, insects). These measurements are used in persons, especially children, with extrinsic asthma, hay fever, and atopic eczema and are an accurate and convenient alternative to skin testing. Although more expensive, they do not cause hypersensitivity reactions.

Additional antigens are continually being added; up-to-date information should be sought. Examples of categories that can be tested for include grasses, trees, molds, venoms, weeds, animal dander, foods, house dust, mites, antibiotics, and insects.

Reference Values

Normal Based on FEIA, the fluorescence is proportional to the amount of specific IgE present in the patient's sample.

Fluorescent Enzyme Immunoassay (FEIA)

Class Interpretation

<table>
<thead>
<tr>
<th>Class</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Negative</td>
</tr>
<tr>
<td>1</td>
<td>Equivocal</td>
</tr>
<tr>
<td>2</td>
<td>Positive</td>
</tr>
<tr>
<td>3</td>
<td>Positive</td>
</tr>
</tbody>
</table>
### Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in biohazard bag for transport to laboratory.

### Clinical Implications
Positive results greater than or equal to class 2 are strongly associated with allergic symptoms on exposure to allergen.

### Interventions

#### Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

#### Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately regarding results and need for other tests. Negative results effectively rule out allergy induced by that allergen.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

### Latex Allergy Testing (Latex-specific IgE)

Latex-containing medical devices include gloves, catheters, and bandages, among many others. Millions of people, especially those in the health care profession, are susceptible to allergic reactions ranging from mild to severe when exposed to such products. It is recommended that patients at risk for latex allergy be tested before undergoing medical procedures that would expose them to latex. High-risk groups include health care workers, workers with industrial exposure to latex, children with spina bifida or urologic abnormalities due to high exposure to latex, and people who have undergone multiple surgeries.

This test measures IgE-mediated latex sensitivity and not irritation or delayed (type IV) reaction to latex. The method for testing is EIA in which the color reaction measured is directly related to the amount of IgE specific for the test allergen in the sample.

### Reference Values

**Normal**
- Negative for latex allergen: <0.35 IU/mL by EIA
- Positive for latex allergen: >0.35 IU/mL

### Procedure

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to laboratory.

### Clinical Implications

1. Positive results are strongly associated with a latex allergy.
2. In studies comparing latex-specific IgE results with clinical history, symptoms, and other confirmatory tests, the sensitivity has been >80% and the specificity >90%.

### Interventions

#### Pretest Patient Preparation

1. Explain test purpose and procedure. Positive history for latex may include the following factors:
   a. Swelling or itching from latex exposure
   b. Hand eczema
   c. Previously unexplained anaphylaxis
   d. Oral itching from cross-reactive foods (eg, banana, kiwi, avocado, chestnuts)
   e. Multiple surgical procedures in infancy
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

#### Posttest Patient Aftercare

1. Interpret test outcomes based on patient's clinical history (ie, latex exposure and laboratory reference values). If negative by this test procedure, yet symptomatic, or if positive for this test, refer patient to an allergist.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

### PROTEIN CHEMISTRY TESTING/SERUM PROTEINS: ACUTE-PHASE PROTEINS AND CYTOKINES

#### Ceruloplasmin

Measurement of ceruloplasmin aids in the diagnosis of copper metabolism disorders, ie, Wilson's disease. Copper bound to ceruloplasmin constitutes the largest amount of Cu $^{2+}$ in circulation. In Wilson's disease, Cu $^{2+}$ mobilization from the
liver is drastically reduced because of the low production of ceruloplasmin.

The test gives a quantitative measurement of the amount of ceruloplasmin in the patient's serum. Values <14 mg/dL are expected in Wilson's disease. Values can vary considerably from patient to patient and may be 50% of normal (pointing to some other primary defect). Patients with Wilson's disease are not always extremely low in ceruloplasmin.

**Reference Values**

**Normal** 25–63 mg/dL (250–630 mg/L) by nephelometry

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to laboratory.

**Clinical Implications** Deficient ceruloplasmin is not the primary defect in Wilson's disease that remains unknown, and can, therefore, vary considerably in its deficiency from patient to patient.

**Interfering Factors**

1. Ceruloplasmin is affected by infections (a late acute-phase reactant) and liver function.
2. Birth control pills increase ceruloplasmin, as does pregnancy.

**Interventions**

**Pretest Patient Preparation**

2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and determine the need for follow-up testing. Values vary from patient to patient and may be 50% or more of normal, pointing to some other defect.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Alpha 1-Antitrypsin**

Alpha 1-antitrypsin (AAT) is the most abundant serum protease inhibitor and inhibits trypsin and elastin as well as several other proteases. The release of proteolytic enzymes from plasma onto surface organs and into tissue spaces results in tissue damage unless inhibitors are present.

Measurement of AAT aids in the diagnosis of juvenile and adult cirrhosis of the liver. AAT deficiency has been associated with neonatal respiratory distress syndrome, severe protein-losing disorders, and pulmonary emphysema. The test is useful for individuals suspected of familial chronic obstructive lung disease.

**Reference Values**

**Normal** 100–200 mg/dL (18.4–36.8 µmol/L) by nephelometry

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**

1. Patients with a serum AAT <70 mg/dL (<12.9 µmol/L) may have a homozygous deficiency and are at risk for early lung disease. AAT phenotyping should be done to confirm the presence of the homozygous deficiency.
2. If clinically indicated, patients with serum levels <125 mg/dL (<23 µmol/L) should be phenotyped to identify heterozygous individuals. The latter do not appear to be at increased risk for early emphysema.

**Interfering Factors** AAT is an acute-phase reactant, and any inflammatory process will elevate serum AAT levels.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose. Follow-up testing, that is, AAT phenotyping, may be necessary if decreased results are obtained.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results and determine the need for phenotyping, which confirms that the deficiency is homozygous (increased risk chronic lung disease) or heterozygous (little if no risk).
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**C-Reactive Protein (CRP) and High-Sensitivity C-Reactive Protein (hs-CRP)**
During any inflammatory process, a specific abnormal protein named C-reactive protein (CRP) appears in the blood. This protein is virtually absent from the blood serum of healthy persons. CRP is one of the most sensitive acute-phase reactants. Levels of CRP can increase dramatically (100-fold or more) after severe trauma, bacterial infection, inflammation, surgery, or neoplastic proliferation. Measurement of CRP has been used historically to assess activity of inflammatory disease, to detect infections after surgery, to detect transplant rejection, and to monitor these inflammatory processes.

There are two types of CRP assays. One measures a wide range of CRP levels to include those found in patients with acute infections. The reportable range is typically 0.3 to 20 mg/dL. The second is a high-sensitivity CRP (hs-CRP) assay. The latter can detect a lower level of CRP to include those that may be of value in measuring the risk for a cardiac event. The sensitivity is to 0.01 mg/dL. The hs-CRP is useful, therefore, for assessment of risk for developing myocardial infarction in patients presenting with acute coronary syndromes.

**Reference Values**

**Normal** <0.8 mg/dL (<8 mg/L) by rate nephelometry for CRP 0.020–0.800 mg/dL (0.2–8.0 mg/L) by immunoturbidimetric assay for hs-CRP

**Procedure**

1. Collect a 7-mL blood serum sample in a red-topped tube. Observe standard precautions.
2. Place the specimen in a biohazard bag for transport to the laboratory.

**Clinical Implications**

1. The traditional test for CRP has added significance over the elevated erythrocyte sedimentation rate (ESR), which may be influenced by altered physiologic states. CRP tends to increase before rises in antibody titers and ESR levels occur. CRP levels also tend to decrease sooner than ESR levels.
2. The traditional test for CRP is elevated in rheumatic fever, RA, myocardial infarction, malignancy, bacterial and viral infections, and postoperatively (declines after fourth postoperative day).
3. A single test for hs-CRP may not reflect an individual patient’s basal hs-CRP level; therefore, follow-up tests or serial measurements may be required in patients presenting with increased hs-CRP levels.

**Interventions**

**Pretest Patient Preparation**

1. Explain the test purpose and procedure. A fasting sample is preferred. Water may be taken.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results, counsel and monitor appropriately. Repeat testing is often necessary to establish an individual’s basal hs-CRP concentration. A positive test indicates active inflammation but not its cause. CRP is an excellent tool for monitoring disease activity. hs-CRP is a tool for assessing cardiovascular risk.
2. In RA, the traditional test for CRP becomes negative with successful treatment and indicates that the inflammation has subsided.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Prion Proteins**

Prions are proteins that occur in both heredity forms and infectious disease. Prions do not contain RNA or DNA. No immune response has been detected. This test is done to diagnose prion brain disease, such as Creutzfeldt-Jacob disease and spongiform encephalitis (“mad cow disease”).

**Reference Values**

**Normal** Structural form named PrPc is found in lymphocytes and in CNS neurons.

**Procedure** Brain tissue samples are examined for evidence of the infectious prion or mutated gene in chromosome 20.

**Clinical Implications**

1. Abnormal finding of PrPc protein (disease-causing form) is pathogenic, which affects the cerebral cortex and cerebellum.
2. Gerstmann-Sträussler-Scheinker syndrome (GSS), cause of hereditary dementia, occurs because of mutation in prion gene
3. Evidence of prion infectious disease may be transfusion-related

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure. Obtain from patient or family signs, symptoms, and history of encephalopathy or dementia (hereditary). Patients are usually very sick and infectious disease is usually fatal.
2. Behavioral changes include ataxia, peripheral sensory changes, dementia

**Posttest Patient Aftercare**

1. Interpret test results and explain possible treatment. Monitor for encephalitis and/or dementia.
2. Provide comfort and support and special counseling regarding progression of disease. Death occurs about 12 months after appearance of first signs.

Cytokines

Cytokines, a diverse group of proteins and peptides secreted by many cells (eg, lymphocytes, T cells, monocytes, B cells, eosinophils), respond to an immunologic challenge. They are involved in immunity, allergy, and long-term memory (ie, degenerative aspects of aging) and include interferons, interleukins, chemokines, inflammatory cytokines, and hematopoietic growth factors. Cytokines have been directly implicated in a number of diseases, such as asthma, interstitial cystitis, RA, septic shock, transplant rejection, cirrhosis, and multiple sclerosis. Most interleukins are produced by macrophages and lymphocytes. Interleukins need adequate amounts of fats and pyridoxine to be effective. Some, such as interleukin-3, are involved in fever, slow-wave sleep, bone resorption, and use of protein by muscles.

These tests are done to evaluate allergy, skin hypersensitivity, asthma, fever, inflammation, and healing. They are also used as tumor markers and to assess immune factors and rheumatic disorders.

Reference Values

**Normal**

1. **Interleukin-1**
   a. 3, 5, 7, 9, 11, 12, 13, 14, 15, 16, 17, and 18
   b. Normal: Physiologic levels are normally very low (few pg/mL or ng/L)

2. **Interleukin-1a**
   a. Plasma: 0.1 + 1.4 pg/mL or ng/L
   b. Urine median: 1–4 pg/μmol creatinine or μg/mol creatinine

3. **Interleukin-1b**
   a. Blood: 4.60 + 300 pg/mL or ng/L
   b. Serum: 0.07 + 0.02 ng/mL or μg/L

4. **Interleukin-2**
   a. Amniotic fluid (AMF): median, 1.35 ng/mL or μg/L
   b. Plasma: 0.3 + 0.47 pg/mL or 0.3 + 0.5 ng/L

5. **Interleukin-4**
   a. See eosinophil count; T cells stimulate eosinophil production.

6. **Interleukin-6**
   a. Urine: 237 + 92 ng/L or 237 + 92 μg/L
   b. Blood: 1609 + 710 pg/mL or 1.61 + 0.71 μg/L
   c. Plasma: 2.50 + 0.35 pg/mL or 2.50 + 0.55 ng/L
   d. Serum: 0.4–2.1 pg/mL or 0.4–2.1 ng/L
   e. CSF: 0.04–12.5 ng/mL or 0.0–12.5 μg/L

7. **Interleukin-8**
   a. Amniotic fluid 237 + 92 ng/L or same

8. **Chemokines**
   a. Feces: <22–4077 pg/g wet stool or 0.02–4.08 ng/g wet stool
   b. Plasma: 3.3 + 0.3 pg/mL or ng/L

9. **Tumor necrosis factors (TNF-a)**
   a. ACSF: 22.3 + 9.5 pg/mL or 1.31 + 0.56 pmol/L
   b. Feces: <1–231 pg/g wet stool or <1–231 ng/g wet stool
   c. Plasma: 6.4 + pg/mL or 6.4 + 4.6 ng/L
   d. Serum: 0.12 + 0.02 ng/mL or 7.0 + 1.2 nmol/L

10. **Interferon-?**
    a. Serum (S): 0.7 + 1.8 pg/mL or 0.7 + ng/L
    b. Plasma (P): 3 + 1 IU/mL or 3 + K IU/L

**Procedure**

1. Collect a stool, urine, or venous blood sample for serum analysis.
2. Be aware that cells from synovial fluid, bronchial secretions, and CSF may also be tested.

**Clinical Alert**

Examine specimens within 5 hours. Avoid a freezing/thawing cycle while stored.

**Clinical Implications**

1. Pathophysiologic blood levels may indicate inflammation or cancer. Increases are associated with severity of disease.
2. Elevated levels in synovial fluid, CSF, amniotic fluid, urine, feces, and bronchoalveolar fluid may indicate immune disorders, SLE, and other pathologic or degenerative conditions.

**Interfering Factors**
1. Cytokines can continue to be produced after sample collection by the various cells in the fluid, urine, or feces.
2. Collection tubes can become contaminated by microorganisms, a potent stimulus of cytokine production.
3. Cytokines can degrade in the collection container.
4. Cytokines can bind to cell receptors during storage.
5. Circadian rhythms may affect results.

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose, procedure, benefits, and risks of cytokine tests and the complexities involved. For specimens other than plasma or serum, refer to specific chapters regarding specimen collection and patient care (eg. urine in Chap. 3, spinal fluid in Chap. 5, CSF studies, amniotic fluid, stool [feces]).
2. Obtain properly signed informed consent when necessary (eg, spinal fluid sample collection). Explain that a local anesthetic will be injected into the skin. Assess for any previous reactions to any numbing or local anesthetic medicines.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Intratest Care** Provide psychological support during specimen collection that may require more invasive procedures.

**Posttest Patient Aftercare**

1. Interpret laboratory test outcomes and counsel appropriately about further testing and treatment. Explain the need for possible identification of chronic disease.
2. Provide the appropriate aftercare if more invasive specimen collection procedures were used (see Chap. 5 for spinal fluid collection aftercare).
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Tumor Markers**

The underlying cause of cancer can be divided into four major classifications: viral, chemical, physical, and genetic. Cancers caused by viruses can be either RNA viruses, eg., retrovirus human T-cell leukemia virus type I (HTLV-I), which causes adult T-cell leukemia, or DNA viruses, eg., hepatitis B virus, which causes hepatocellular carcinoma. Chemical carcinogens can be classified as either genotoxic (targeting the DNA), eg., nitroso compounds that when heated release toxic fumes or solvents such as trichloroethylene, or nongenotoxic (targeting cell death directly or hormonal effects), eg., synthetic pesticides or herbicides. Physical factors associated with causing cancer include ultraviolet light (sunlight), ionizing radiation (x-rays), and asbestos fibers. Hereditary or genetic cancer can account for up to 30% of some forms of childhood cancers and 5%-10% of adult cancers.

Cancerous cells differ from normal cells in many respects. Malignant cells grow more rapidly in an uncontrolled fashion, lack normal cell-to-cell interactions, and apoptotic (programmed cell death) mechanisms are disrupted when compared to normal cells. Normal cells, through a series of mutations (referred to as hits) and alterations of normal cell growth and cell interactions, can transition into cancerous cells. This unregulated and disorganized increase in cell growth is stimulated by the factors described above. Tumors, by definition, are spontaneous growth of abnormal cells leading to a swelling or enlargement of the underlying tissue. This abnormal cell growth, or cancer, can be detected by certain substances (tumor markers) found in the blood.

### Types of Cytokines, Cellular Origin, and Clinical Implications

<table>
<thead>
<tr>
<th>Cytokine (Synonym)</th>
<th>Cellular Origin</th>
<th>Target Cells</th>
<th>Specific Clinical Implications of Abnormal Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>INTERLEUKIN-1</td>
<td>Monocytes, macrophages, antigen-presenting cells (APCs), endothelial cells, T lymphocytes, Natural Killer (NK) cells</td>
<td>Monocytes, macrophages, hepatocytes, endothelial cells, epithelial cells, fibroblasts, keratinocytes, T lymphocytes, B lymphocytes, NK cells, osteoclasts</td>
<td>Increased in rheumatoid arthritis, septic shock, periodontitis, malignancy, asbestosis, tuberculosis, HIV infection, differential diagnosis, ie, nonallergic bronchial asthma, multisystem organ failure during acute pancreatitis, acute eosinophilic pneumonia, inflammatory intestinal disease. Therapeutically useful in protecting patients against lethal doses of radiation and in stimulation of hematopoiesis</td>
</tr>
<tr>
<td>Also known as</td>
<td>B-cell accelerating factor, catabolin, endogenous pyroge, epithelial cell-derived thymocyte-activity factor, fibroblast-activity factor, hemoposition-1, hepatogenic stimulatory factor (HSF), leukocyte endogenous mediator, lymphocyte-activating factor, mononuclear cell factor, osteoclast-activating factor, proteolysis-inducing factor, and serum amyloid A inducer</td>
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</tbody>
</table>

Minimum Error Rate: This document was processed with a minimum error rate of 0.015.
Also known as

T-cell growth factor (TCGF)

T lymphocytes (CD4, Th0 and Th1 [CD8], NK lymphocytes, B lymphocytes, mast cells

Pregnancy, increased in multiple sclerosis, sarcoidosis, multiple myeloma, acute rheumatic fever and chronic rheumatic heart disease. Decreased in advanced age, diabetes, Sjögren's syndrome and against (AIDS). Therapeutic use in clinical trials for treatment of cancer (kidney, melanoma, and acute myeloid leukemia). AIDS, leprosy, and other immunologic conditions resulting from administration of immunosuppressive agents

INTERLEUKIN-3

Colony-forming unit (CFU) stimulating activity, colony-stimulating factor (CSF), hematopoietic growth factor, mast cell growth factor, megakaryocytic growth factor (MCGF), histamine-producing cell-stimulating factor, multilineage hemopoietic growth factor, P-cell stimulating factor activity, persisting (P)-cell stimulating factor, synergistic activity (Thy-1 inducing factor, and WEHI-3 growth factor)

INTERLEUKIN-4

B-cell differentiating factor (BCDF), B-cell growth factor I (BSF-I), B-cell stimulating factor I (BSF-I), MCGF-II, T-cell growth factor II (TCGF-II)

Most interleukins are produced by macrophages or T lymphocytes (CD4, T, H1 and T H2), NK cells, mast cells, eosinophils.

Hematopoietic stem cells, progenitor cells, B lymphocytes, macrophages, polymorphonuclear leukocytes, mast cells, keratinocytes.

Increased production in monocytes and myeloid leukemia and most cell disorders. In blood only with severe immunization against stimuli such as graft-versus-host disease or parasitic infestations; plays a role in chronic allergic diseases. Therapeutic use in clinical trials to assess the benefit from temperature stimulation, shortening the duration of cytopenia associated with chemotherapy, irradiation, or following bone marrow transplantation

INTERLEUKIN-5

BCDF for IgM (BCDFu), BCGF-II, eosinophil colony-stimulating factor, eosinophil differentiation factor, IgA-enhancing factor, and T-cell replacing factor

T lymphocytes (CD4, T H2), basophils, eosinophils, mast cells

Eosinophils, B lymphocytes, mast cells

Increased in allergic diseases, asthma in late allergen manifestations; patients with acute eosinophilic pneumonia have high bronchoalveolar lavage fluid levels of IL-5, IL-IRA, and soluble type II IL-IR. Production high in severe atopic dermatitis. Therapeutic use for hypogammaglobulinemia

INTERLEUKIN-6


T lymphocytes, B lymphocytes, monocytes, macrophages, APCs, endothelial cells, epithelial cells, fibroblasts, mast cells

T lymphocytes, B lymphocytes, hepatoctyes, endothelial cells, keratinocytes, hematopoietic cells, malignant plasma cells

Increased in gram-negative bacterial infection and inflammatory reactions, bacterial sepsis, acute complications of lung transplantation. IL-6 is produced by macrophages in the periprosthetic membrane surrounding joint replacements, causing osteolysis and prosthetic loosening. Hyperproduction associated with a variety of malignancies, including plasmacytoma, multiple myeloma, uterine and cervical cancer, and Kaposi's sarcoma. Therapeutic use to support neutrophil and megakaryocyte growth in patients treated with high doses of chemotherapy and to treat patients infected with human T-cell leukemia virus type 1 (HTLV-1)

INTERLEUKIN-7

Lymphopoietin 1, B-cell growth factor, and pre-B-cell growth factor

Stromal cells (bone marrow, thymic), T lymphocytes, spleen cells, epithelial cells, fibroblasts

T lymphocyte progenitors, B lymphocytes, T lymphocytes (CD4, CD8)

Regulates HIV-1 replication in naturally infected peripheral blood mononuclear cells (ABMCs). Therapeutic use—may be useful in tumor therapy because of its ability to enhance the generation of T lymphocytes and lymphokine-activated killer (LAK) cells, even in the absence of IL-2; helpful for some immunodeficiencies or to accelerate recovery following bone marrow transplantation. May enhance immune system

Colony-forming unit (CFU)

Also known as

T-cell growth factor (TCGF)

T lymphocytes (CD4, Th0 and Th1 [CD8], NK lymphocytes, B lymphocytes, mast cells

Pregnancy, increased in multiple sclerosis, sarcoidosis, multiple myeloma, acute rheumatic fever and chronic rheumatic heart disease. Decreased in advanced age, diabetes, Sjögren's syndrome and against (AIDS). Therapeutic use in clinical trials for treatment of cancer (kidney, melanoma, and acute myeloid leukemia). AIDS, leprosy, and other immunologic conditions resulting from administration of immunosuppressive agents

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Most interleukins are produced by macrophages or T lymphocytes (CD4, T, H1 and T H2), NK cells, mast cells, eosinophils.

Hematopoietic stem cells, progenitor cells, B lymphocytes, macrophages, polymorphonuclear leukocytes, mast cells, keratinocytes.

Increased production in monocytes and myeloid leukemia and most cell disorders. In blood only with severe immunization against stimuli such as graft-versus-host disease or parasitic infestations; plays a role in chronic allergic diseases. Therapeutic use in clinical trials to assess the benefit from temperature stimulation, shortening the duration of cytopenia associated with chemotherapy, irradiation, or following bone marrow transplantation

INTERLEUKIN-5

BCDF for IgM (BCDFu), BCGF-II, eosinophil colony-stimulating factor, eosinophil differentiation factor, IgA-enhancing factor, and T-cell replacing factor

T lymphocytes (CD4, T H2), basophils, eosinophils, mast cells

Eosinophils, B lymphocytes, mast cells

Increased in allergic diseases, asthma in late allergen manifestations; patients with acute eosinophilic pneumonia have high bronchoalveolar lavage fluid levels of IL-5, IL-IRA, and soluble type II IL-IR. Production high in severe atopic dermatitis. Therapeutic use for hypogammaglobulinemia

INTERLEUKIN-6


T lymphocytes, B lymphocytes, monocytes, macrophages, APCs, endothelial cells, epithelial cells, fibroblasts, mast cells

T lymphocytes, B lymphocytes, hepatoctyes, endothelial cells, keratinocytes, hematopoietic cells, malignant plasma cells

Increased in gram-negative bacterial infection and inflammatory reactions, bacterial sepsis, acute complications of lung transplantation. IL-6 is produced by macrophages in the periprosthetic membrane surrounding joint replacements, causing osteolysis and prosthetic loosening. Hyperproduction associated with a variety of malignancies, including plasmacytoma, multiple myeloma, uterine and cervical cancer, and Kaposi's sarcoma. Therapeutic use to support neutrophil and megakaryocyte growth in patients treated with high doses of chemotherapy and to treat patients infected with human T-cell leukemia virus type 1 (HTLV-1)

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Regulates HIV-1 replication in naturally infected peripheral blood mononuclear cells (ABMCs). Therapeutic use—may be useful in tumor therapy because of its ability to enhance the generation of T lymphocytes and lymphokine-activated killer (LAK) cells, even in the absence of IL-2; helpful for some immunodeficiencies or to accelerate recovery following bone marrow transplantation. May enhance immune system
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<tbody>
<tr>
<td>INTERLEUKIN-8</td>
<td>(see Chemokines)</td>
<td>B-cell derived T-cell growth factor, cytokine synthesis inhibiting factor, MCGF, and thymocyte growth-promoting factor</td>
</tr>
<tr>
<td>INTERLEUKIN-9</td>
<td></td>
<td>T lymphocytes (CD4, TH2), lymphoma cells progenitor cells</td>
</tr>
<tr>
<td>INTERLEUKIN-10</td>
<td></td>
<td>Increased: T-cell tumors, especially of TH2 lymphocytes. In Hodgkin's lymphoma acts as autoimmune signal for cellular proliferation. No therapeutic applications</td>
</tr>
<tr>
<td>INTERLEUKIN-11</td>
<td>Adipogenesis inhibitor factor</td>
<td>Stromal cells (bone marrow), trophoblasts, glial cells, fibroblasts</td>
</tr>
<tr>
<td>INTERLEUKIN-12</td>
<td>Cytotoxic lymphocyte maturation factor, K-cell stimulating factor, and T-cell stimulating factor</td>
<td>B lymphocytes, monocytes, macrophages, APCs</td>
</tr>
<tr>
<td>INTERLEUKIN-13</td>
<td>NC30 (human CDNA) and P6001 mouse CDNA clone</td>
<td>T lymphocytes (CD4, TH1, CD8), mast cells</td>
</tr>
<tr>
<td>INTERLEUKIN-14</td>
<td>High-molecular-weight BCGF</td>
<td>T lymphocytes, B lymphocytes, dendritic cells, malignant cells</td>
</tr>
<tr>
<td>INTERLEUKIN-15</td>
<td></td>
<td>UVB-induced down-regulation of keratinocytes. IL-15 production may contribute to the relative state of immunosuppression induced by sun exposure. IL-15 triggers the growth of leukemia B cells through IL-2R system subunits, relating to the role of cytokine in regulating neophasic proliferation in chronic lymphoproliferative disorders. Synovial fluid T lymphocytes proliferate in response to IL-15, thereby contributing to rheumatoid arthritis pathogenesis</td>
</tr>
<tr>
<td>INTERLEUKIN-16</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Lymphocytic chemoattractant factor</strong></td>
<td><strong>T lymphocytes</strong> (CD8), eosinophils, epithelial cells</td>
<td><strong>T lymphocytes</strong> (CD4), eosinophils (CD4), monocytes (CD4)</td>
</tr>
<tr>
<td><strong>INTERLEUKIN-17</strong></td>
<td><strong>T lymphocytes</strong> (CD4-activated memory CD4)</td>
<td><strong>T lymphocytes</strong>, fibroblasts, epithelial cells, endothelial cells</td>
</tr>
<tr>
<td><strong>INTERLEUKIN-18</strong></td>
<td><strong>Liver cells</strong></td>
<td><strong>NK cells, T lymphocytes</strong></td>
</tr>
<tr>
<td><strong>INTERFERONS: IFN-a and IFN-ß</strong></td>
<td><strong>IFN-a: Monocytes, macrophages, lymphocytes, IFN-ß: Fibroblasts, epithelial cells. Interferons are produced by virus-infected cells and are the body’s first line of defense against many viruses.</strong></td>
<td><strong>T lymphocytes, NK cells, macrophages</strong></td>
</tr>
<tr>
<td><strong>IFN-?</strong></td>
<td><strong>T lymphocytes (CD4, T lymphocytes, NK T $T_{H0}$, $T_{H1}$, CD8), NK cells, macrophages, endothelial cells, APCs, B lymphocytes</strong></td>
<td>Increased IFN-? production is associated with food-sensitive atopic dermatitis, Crohn’s disease (local mucosal synthesis), rheumatoid arthritis synovial fluid (possible role in the ongoing immunologic reaction of the inflamed joint), and poor antibody-mediated immunity in hypovitaminosis A. Also, IFN-? is associated with immunodeficiency after allogeneic bone marrow transplantation, the pathophysiology of aplastic anemia, and atherogenesis. Therapeutic use as an antiviral; antiproliferative and immunomodulatory effects. Its instability limits its clinical application.</td>
</tr>
</tbody>
</table>
TFN-a: cachectin, cytotoxin, cytotoxic factor, differentiation-inducing factor, hemorrhagic factor, macrophage cytotoxic factor, and necrosis. TNF-ß: Cytotoxin, differentiation-inducing factor, and lymphotoxin (LT)

Monocytes, macrophages, T lymphocytes, B lymphocytes, NK cells, mast cells, endothelial cells, APCs, fibroblasts

T lymphocytes, polymorphonuclear leukocytes (PMNs), macrophages, osteoclasts, fibroblasts, hepatocytes, tumor cells

Monocytes, macrophages, PMNs, T lymphocytes, B lymphocytes, NK cells, mast cells, endothelial cells, epithelial cells, APCs, stromal cells, fibroblasts, platelets

Monocytes, macrophages, PMNs, T lymphocytes, B lymphocytes, NK cells, mast cells, endothelial cells, epithelial cells, APCs, stromal cells, fibroblasts, megakaryocytes

TNF-a is associated with septic shock and diverse infectious pathology (protective immune response in pulmonary tuberculosis). TNF-a is important in the pathogenesis of allergic respiratory reactive and acute stages of rheumatoid arthritis. Injured patients reveal significantly increased plasma levels of both TNFs compared with healthy persons; severe injury reveals higher levels on day of admission than in patients with minor injury. TNF-a levels higher in peritoneal fluid of women with endometriosis than in healthy women. Increased TNF-a, IL-6, IL-1ß levels following transient cerebral ischemia suggest cytokine may be involved in pathophysiologic changes in hippocampus and striatum. TNF-a may play a role in acute pancreatitis and mediate the systemic sequelae of the disease. Increased serum tissue TNF levels observed in alcoholic liver disease; increased plasma levels in nephropathia epidemica. TNF-a involved in pathogenesis of both acute and chronic transplant rejection; those with higher levels of this cytokine are at risk for a number of T-lymphocyte–mediated autoimmune diseases such as type 1 diabetes. TNF-a and IFN-ß are involved in the progression of neurologic disorders such as multiple sclerosis and AIDS complex, anti–TNF-a antibodies, lupus nephritis, inflammatory arthritis, cerebral edema, and neurologic dysfunction.

CHEMOKINES (CKS)
(Formerly known as intercrines, the scy [small cytokine] family, and small, inducible, secreted cytokines.)
A condensation of the term chemoattractant cytokines.
Now defined as a superfAMILY of low-molecular-weight proteins (8–10 kd).

Regulate the motility and orientation of leukocytes (leukocyte activation/inflammation), lymphocyte activation and diversification, platelet activation, biologic modifier of erythrocyte function, angiogenic activity factors and cell adhesion. Play an important role as pathogenic mediators in several pathologies, including asthma, allergic diseases, autoimmune diseases, inflammatory/infectious processes (inflammatory bowel disease), rheumatoid arthritis, sarcoidosis, pulmonary fibrosis, AIDS/HIV infections, transplant rejection, kidney pathology and proteinuria, hemolytic decreases/transfusion and atherosclerosis.

There are a number of factors that have either a protective effect or promote cancer growth.

<table>
<thead>
<tr>
<th>Tumor Protectors</th>
<th>Tumor Promoters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genetic resistance</td>
<td>Genetic susceptibility</td>
</tr>
<tr>
<td>Tumor suppressor genes</td>
<td>Age</td>
</tr>
<tr>
<td>Immune system</td>
<td>Smoking</td>
</tr>
<tr>
<td>Programmed cell death</td>
<td>Asbestos exposure</td>
</tr>
<tr>
<td>DNA repair</td>
<td>Resistance to cytotoxicity</td>
</tr>
</tbody>
</table>

Tumor markers include genetic markers (abnormal chromosomes or oncogenes), DNA analysis, oncofetal antigens, enzymes, hormones, placental proteins, steroid receptors, glucoproteins, tumor-associated antigens, tumor-specific antigens, and circulating immune complexes.

Tumor cells differ from normal cells in many ways. Physical examination and standard x-ray techniques can usually detect tumors as small as 1 cm in volume. A tumor mass of this size has completed 30 doublings (two thirds of its growth) and contains 1 billion \((10^{9})\) cells. Certain tumor antigens, hormones, oncofetal proteins, and enzymes are secreted into the bloodstream by these tumors.
Malignant tumor cells are produced when DNA is damaged by some form of carcinogen, virus, radiation, or chemical causing the process of mitosis to go out of control. These growing, changed (mutant) cells express oncogenes. These oncogenes are capable of inducing or transforming cells into cancer cells or tumors. Tumor cells capable of forming metastases are likely to invade blood vessel walls; be released into the bloodstream, regional lymphatics, or interstitial stoma; and eventually spread to other organs. Tumor testing has focused on identifying certain tumor-related substances that might allow early detection of malignancy, determination of prognosis, and evaluation of tumor burden (ie, size, location, and encroachment on other tissues or organs).

Tumor markers are used and developed to obtain greater sensitivity and specificity in determining the presence of cancer and tumor activity. These substances are found in body cells, fluids, and tissue. In general, these markers lack specificity for cancer; none is pathognomonic for any one type of neoplasm. Diagnosis still derives from a biopsy and tissue examination, comprehensive patient history, physical examination, and other diagnostic procedures.

**Education Alert**

Tumor marker studies do not replace biopsy and pathologic tissue examination and are not ideal for screening for specific cancers, making a diagnosis, or predicting programs for symptomatic patients, but they are effective for tumor staging, monitoring response to therapy, detecting disease recurrence, and early detection of cancer recurrence.

The following are diagnostic prognostic and predictive markers:

1. **Oncofetal antigens** (Oncofetal antigens, normally produced in the fetus, are reactivated with cancer cell transformation.)
   a. Carcinoembryonic antigen (CEA)
   b. Alpha-fetoprotein (AFP)
   c. Proteins
   d. CA 125
   e. CA 19-9
   f. CA 15-3
   g. CA 549
   h. Tissue polypeptide antigen (TPA)
   i. Prostate-specific antigen (PSA)
   j. hK2 and hK3 of gene family = kallikreins
   k. Human glandular kallikrein for prostate cancer

2. **Placental proteins**
   a. Human chorionic gonadotropin (hCG and ß-hCG)
   b. Human placental lactogen (HPL)
   c. Placental alkaline phosphatase (PALP)

3. **Enzymes and isoenzymes**
   a. Prostatic acid phosphatase (PAP)
   b. Creatine kinase (CK)-BB isoenzyme
   c. Alkaline phosphatase (ALP)
   d. Neuron-specific enolase (NSE)
   e. Lactate dehydrogenase isoenzyme (LDI)
   f. Lysozyme (muramidase)

4. **Hormones**
   a. Hormones, both normally produced by the tissue and ectopic
   b. GGT
   c. F LNT
   d. Amylase
   e. TDD (terminal deoxynucleotydyl transferase)
   f. hCG—trophoblastic tumors
   g. Nonseminomatous testicular tumors
   h. HIAA
   i. Epinephrine and norepinephrine—pheochromocytoma and related malignancies
   j. DMA
   k. 17KS
   l. Gastrin-Zollinger-Ellison syndrome (gastrinoma)
   m. Renin-producing by kidney
   n. Calcitonin—medullary carcinoma of the thyroid (not normally produced by the tissue)
   o. Adrenocorticotrophic hormone (ACTH)—small cell carcinoma of the lung
   p. Antidiuretic hormone (ADH)
   q. Parathyroid-related peptide
   r. Erythropoietin
   s. Gastrin
   t. HIAA

5. **(Serotonin) immunoglobulins**
   a. IgG
   b. IgA
   c. IgM
   d. IgD
   e. IgE
   f. Kappa and lambda light chains

6. **Steroid receptors**
a. Estrogen and progesterone receptors (ER and PR)
b. Epidermal growth factor (EDFR)
c. HER-2 (human epidermal growth factor receptors—determine suitability for HER epton therapy)
d. Androgen receptors
e. Corticosteroid receptors

7. Immunocomplex typing
   a. Lymphoid cells
   b. Myeloid cells
c. Cytokines (see page 606 for more information)

8. DNA analysis
   a. Ploidy and S-phase fraction
   b. See Chapter 11 for more information.

9. Molecular diagnosis
   a. Oncogene and suppressor genes
   b. Genetic changes
c. See Chapter 11 for more information.

Reference Values

Normal: See Clinical Implications for value for each specific tumor marker.

Procedure

1. Be aware that most tumor marker tests involve obtaining either venous plasma or serum; urine or bladder washings or CSF; some may require fasting.
2. Follow the specific directions from the laboratory or cancer center involved in the testing procedure. Be sure to note factors that interfere with test results.

Clinical Implications

1. Tumor markers, substances produced and secreted by tumor cells and found in serum, urine, or tissue of persons with cancer, are indicative of tumor activity.
2. Table 8.9 includes tumor-specific or tumor-associated antigens (proteins and oncofetal antigens), enzymes, hormones, and cytokines.

Table 8.9 Tumor-specific and Tumor-associated Agents

<table>
<thead>
<tr>
<th>Name of Test</th>
<th>Clinical Marker in Current Use and Selected Normal Values</th>
<th>Type of Cancer in Which Tumor Marker May Be Found</th>
<th>Conditions Other Than Cancer Associated With Abnormal Values</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ENZYMES</strong></td>
<td>1. Prostatic acid phosphatase (PAP). Major pretreatment tumor marker and used to predict recurrence. Increased values due to increased metabolism and catabolism of cancer cells—levels increase with stage of cancer and age of individual. Prostate-specific antigen (PSA) done to monitor prostate cancer preferred screening marker (&lt;2.7 ng/mL or &lt;2.7 µg/L). Monitor therapy with antineoplastic drugs.</td>
<td>1a. Carcinoma of prostate with the following elevation: carcinoma with no metastasis 10%–20%; metastasis with one 20%–40%; metastases with bone involvement 70%–90% (usually osteoblastic). In three fourths of patients, arises in posterior lobe of prostate. Leukemia (hair-eye-cell) increased. Cancer metastatic to bone (increased osteoblastic lesions)</td>
<td>1. Increased in noncancer prostatic condition, prostate palpation, hyperplasia, infection of prostate following cystostomy, prostate surgery, and chronic prostatitis. Other increases: Gaucher's disease (lipid storage disease), Nieman-Pick disease, Paget's disease, osteoporosis, renal osteopathy, hepatic cirrhosis, pulmonary embolism, and hyperparathyroidism</td>
</tr>
<tr>
<td></td>
<td>2. Lactate dehydrogenase (LDH); increased isoenzymes I and II. Total LDH: 166–280 U/L. Detect and monitor testicular cancer.</td>
<td>1b. Is specific for prostate cancer</td>
<td>2. Increased in cellular injury/hemolysis, early in myocardial infarction, hepatic diseases (see Cardiac Marker Tests in Chap. 6).</td>
</tr>
<tr>
<td></td>
<td>3. Neuron-specific enolase (NSE) &lt;12.5 µg/mL; normal staining. Produced by neurons and neuroendocrine cells of the central and peripheral nervous system. Used to monitor disease progression, small cell lung cancer and pheochromocytoma neuroblastoma, medullary thyroid cancer.</td>
<td>2. Increased neuroblastic carcinoma of tests. Elevated in 60% of those with stage III testicular cancer—serial LDH may help to detect recurrence of cancer. Ewing's sarcoma, acute lymphocytic leukemia, non-Hodgkin's lymphoma, LD-1 increased in germ cell tumors; LD-3 increased in leukemia; LD-5 increased in breast, lung, stomach, and colon; elevated in metastatic carcinoma</td>
<td>3. Occasionally with benign liver diseases</td>
</tr>
<tr>
<td></td>
<td>4. Cytokines</td>
<td>3. NSE increased in neuroblastosomas, APUD system tumor—small cell lung cancers, pancreatic islet cell, medullary thyroid carcinoma, seminoma (20%) in prostate, breast, and gastrointestinal (GI) tract, also Wilm's tumor and pheochromocytoma</td>
<td></td>
</tr>
</tbody>
</table>

5. Other enzymes: gamma-glutamyl transpeptidase (GGT); muramidase, creatinine, phosphokinase isozyme BB, beta-glucuronidase, terminal deoxynucleotidyl transferase, ribonuclease, histaminase (medullary cancer of thyroid), amylase, and cystine aminopeptidase

6. Squamous cell cancer antigen (SCCA). Used to monitor and detect recurrence of squamous cell cancer of uterus, cervix, head and neck, esophagus, lung, skin, and sinus; also advanced cancer

HORMONES

1. Human chorionic gonadotropin (hCG) produced by placental syncytiotrophoblast; not usually found in sera of healthy, nonpregnant persons. <2 ng/mL. Useful to monitor testicular tumors and tumors of ovary and to monitor changes.

2. Calcitonin (CT): malignant C-cell thyroid tumor produces increased CT in ascites. Calcitonin (see Chap. 6) is a hormone produced by perifollicular C cells of thyroid gland. Ranges vary with method. Serum: Adult: <150 pg/mL. Plasma: male, <19 pg/mL; female, <14 pg/mL.

3. Other hormones: adrenocorticotropic hormone (ACTH) (lung—oat cell), parathyroid hormone (PTH) (lung—epidermoid), insulin (lung), glucagon (pancreas), gastrin (stomach and other carcinomas), prostaglandins and erythropoietin (kidney).

4. Serotonin (5-hydroxyindole acetic acid [5-HIAA]). Hydroxyacetic acid: used to detect and monitor carcinoid tumors.

5. Creatine phosphokinase (CPK)-BB increased in prostate, lung (small cell), bladder, breast and GI tract cancer; amylase increased in lung and ovarian cancer

6. Increased in uterine cancer (89% of stage IV disease). Alert: occurs in saliva, sweat, and respiratory secretions.

ONCOFETAL ANTIGENS

1. Alpha-fetoprotein (AFP) is a glycoprotein produced by fetal liver, yolk sac, and intestinal epithelium. Disappears from blood soon after birth and is not present in healthy individuals <40 ng/mL. Diagnose and monitor AFP tumors. Follow-up for therapy of testicular, ovarian, and primary liver tumors; used with hCG.

2. Calcitonin (CT): malignant C-cell thyroid tumor produces increased CT in ascites. Calcitonin (see Chap. 6) is a hormone produced by perifollicular C cells of thyroid gland. Ranges vary with method. Serum: Adult: <150 pg/mL. Plasma: male, <19 pg/mL; female, <14 pg/mL.

3. Other hormones: adrenocorticotropic hormone (ACTH) (lung—oat cell), parathyroid hormone (PTH) (lung—epidermoid), insulin (lung), glucagon (pancreas), gastrin (stomach and other carcinomas), prostaglandins and erythropoietin (kidney).

4. Serotonin (5-hydroxyindole acetic acid [5-HIAA]). Hydroxyacetic acid: used to detect and monitor carcinoid tumors.

1. Increased in primary hepatocellular cancer, embryonal cell (nonseminomatous germ cell) testicular tumors, yolk sac ovarian tumors, teratocarcinoma, gastric, pancreatic, colonic, breast, renal, and lung. >50 ng/mL AFP-producing tumor. Elevations signal recurrence.

2. Increased in Zollinger-Ellison syndrome, pernicious anemia, chronic renal failure, pseudohypoparathyroidism, apudomas, alcoholic cirrhosis, Paget's disease, pregnancy, and benign breast or ovarian disease. Decreased with therapy; an increase after therapy suggests progressive disease

3. See Chap. 6 for specific substance increases.
2. Carcinembryonic antigen (CEA). Initially isolated in endodermally derived adenocarcinoma and fetal gastrointestinal tissue. <2.5–5 ng/mL; up to 5 ng/mL in smokers; <6 ng/mL in spinal fluid. Assess therapy with antineoplastic drugs and following surgery of medullary thyroid cancer, neoplasma of breast, GI tract, lung, and colorectal; monitor cancers of primary colorectal cancer, pancreas, breast, GI, liver, lung, ovaries, prostate.

**PROTEINS**

1. CA 15-3 antigen (cancer-associated antigen breast cystic fluid protein [BCFP]; used in conjunction with CEA); is a marker for breast cancer used for serial testing. <30 U/mL males and females encoded by MUC-1 gene in stage II or III and used with CA 27 and CA 29. Most useful to monitor therapy and disease progression in metastatic disease.

2. NMP22 found in urine, used to manage transitional cell cancer of the urinary tract tissue biopsy obtained by cystoscopy.

3. Bladder tumor associated (BTA) analyte found in urine.

4. CA 27–29 (similar to CA 15-3 serum) is a marker for breast carcinoma. Not used for screening. >38 U/mL in female and male breast cancer. Serial testing for prior stage II or III to detect recurrence.

5. B₂-Microglobulin (β₂) (HLA antigen system). 4–12 mg/L.

6. PSA—more sensitive than PAP—correlates with stage of adenocarcinoma disease and age. Males: 80% <2.0 µg/L; free: total ratio = >0.24 total 2–10 ng/mL = <2.0 ng/mL. Free cascade done. Free PSA to assess risk for cancer with borderline PSA (2–10 ng/mL). PSA screening for prostate cancer recommended only for men >50 yr. Useful for monitoring and staging prostate cancer. PSA not significantly increased until tumor has grown out of prostate gland.

7. CA 19-9 carbohydrate antigen; <37 U/mL. Occurs in serum and tissue. Is a marker for colorectal and pancreatic cancer.

8. CA 125 (ovarian cancer) (glycoprotein antigen) and serum carbohydrate antigen. <35 U/mL. Is a marker for ovarian and endometrial carcinoma. Ovarian and endometrial cancer monitoring.

9. CA 72-4 antigen (serum). Includes melanoma, breast, ovarian, gastrointestinal tract, pancreas, and prostate. Caution: May increase in acute and chronic hepatitis, pulmonary emphysema, bronchitis, peritonitis, pancreatitis, GI disease, Meigs' syndrome, pleural effusion, pulmonary disease, pericarditis, and ovarian cysts.

10. CA 14-3-3 protein (protein encoded by the cell cycle regulatory genes CCNE1 and CCNE2) (serum). Increased in hematopoietic malignant neoplasms; melanoma, breast cancer, colon cancer, thyroid cancer, neuroblastoma, myeloma, other B-cell lymphocytic leukemia, multiple myeloma, and lung cancer.

11. ß-Human chorionic gonadotropin (ß-HCG) and alpha-fetoprotein (AFP) (serum). Increases in early pregnancy, embryonic carcinomas, hepatobiliary carcinoma, acute and chronic hepatitis, pulmonary emphysema, bronchitis, ovarian cysts, Meigs' syndrome, and Meigs' syndrome-like conditions. Decreased with therapy; an increase after therapy suggests progressive disease. Increased in pancreas, liver, colorectal, ovarian, and liver cancer.

12. CA 12–5 antigen (serum). Increased in pancreatic adenocarcinoma. Not used for screening.


15. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.


18. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.


21. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.

22. CA 19-9 antigen (serum). Increased in pancreatic adenocarcinoma. Not used for screening.


24. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.


27. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.


30. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.


32. CA 15-2 antigen (serum). Increased in breast cancer. Not used for screening.

33. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.

34. CA 19-9 antigen (serum). Increased in pancreatic adenocarcinoma. Not used for screening.


36. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.

37. CA 19-9 antigen (serum). Increased in pancreatic adenocarcinoma. Not used for screening.


39. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.

40. CA 19-9 antigen (serum). Increased in pancreatic adenocarcinoma. Not used for screening.

41. CA 15-2 antigen (serum). Increased in breast cancer. Not used for screening.

42. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.

43. CA 19-9 antigen (serum). Increased in pancreatic adenocarcinoma. Not used for screening.

44. CA 15-2 antigen (serum). Increased in breast cancer. Not used for screening.

45. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.

46. CA 19-9 antigen (serum). Increased in pancreatic adenocarcinoma. Not used for screening.

47. CA 15-2 antigen (serum). Increased in breast cancer. Not used for screening.

48. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.

49. CA 19-9 antigen (serum). Increased in pancreatic adenocarcinoma. Not used for screening.


51. CA 12–5 antigen (serum). Increased in ovarian cancer. Not used for screening.

52. CA 19-9 antigen (serum). Increased in pancreatic adenocarcinoma. Not used for screening.
9. CA 50, <17 U/mL. Is a marker for pancreatic and colorectal carcinoma. Used to monitor therapy of GI and pancreatic cancer.

10. CA 72-4 TAG (a micin-like hormone adenocarcinoma associated antigen) <4.0 ng/mL. Is a marker for GI and ovarian cancer. Used in gastric carcinoma monitoring

11. CS549 (acidic glycoprotein), <15.5 U/mL (results correlate with those using CA 15-3). Used in monitoring breast cancer

12. Tissue polypeptide antigen (TPA) 80–100 U/L in serum—may also be detected in urine, washings, and effusions. Is not a specific marker. Monitor GI, genitourinary (GU), breast, lung, and thyroid cancer.

13. Immunoglobulins: monoclonal proteins (M proteins), immunoglobulins produced by B lymphocytes. Normal: Absent. Refer to serum protein electrophoresis (SPEP) or urine protein electrophoresis (UPEP).

14. Tumor-antigen 4 (TA-4), =2.6 ng/mL. Diagnose and monitor squamous cancer.

15. CA 242 is a marker for pancreatic and colorectal cancer.

16. Other antigens: colon mucoprotein antigen (CMA), colon-specific antigen (CSA), zinc glycinate marker (ZGM—colon), pancreatic oncofetal antigen (POA), S-100 protein (malignant melanoma), sialoglycoprotein (wide variety of cancers), B protein (wide variety of antigens), and “Tennessee” antigen glycoprotein (wide variety of cancers)

CYTOKINES

1. Interleukin (IL) (also known as interleukin IL-2) T-cell growth factor I; formed from helper T cells and activated B cells; results highly variable. Monitor therapy in leukemia. Results highly variable

GENETIC MARKERS

Suppressor Genes

1. P-53 gene mutation. No mutation. Most common genetic mutate in cancer used for prognosis

2. Retinoblastoma gene

3. BRCA1 and BRCA2 monitor development of breast and ovarian cancer

4. p21 WAF1 may be clinically useful

5. APC (antigen presenting cells)

1. Increased in leukemias (adult T-cell leukemia)

1. High mutation over expression in breast, BRCA, head and neck, colon, and small cell lung cancer (50%–75%)

2. Found in ocular tumors arising spontaneously; small portion are hereditary

3. Found in hereditary predisposition to developing breast and ovarian cancer

4. Uncertain

5. Increased in patients with hundreds of polyps. Mutations in hereditary colon cancer, also breast and esophageal
6. Neurofibromatosis
6. Inactivating mutations in inherited colorectal cancer, melanoma
6. Information not available

7. Wilm's tumor (nephroblastoma)
7. Mutations in Wilm's tumor
7. Information not available

8. NM2, a marker for metastasis
8. Increased in metastatic breast, colon, and prostate cancer
8. Information not available

Oncogenes
1. Ras oncogene. No mutation
1. Oncogene mutations found in leukemia, neuroblastoma, lymphoma, sarcomas, and endotheliomas
1. Information not available

2. CMYC gene-defect recurrence
2. Found in B- and T-cell lymphoma and small cell lung cancer
2. Information not available

3. C-erb B-2, HER-2/Neu gene, used as a prognostic indicator in breast cancer
3. Found in breast, ovarian, and GI carcinomas
3. Information not available

4. bcl-2 (blocks apoptosis), presence contributes to programmed cell death and survival of cancer cells
4. Found in leukemia and lymphoma. Detection may predict resistance to chemotherapy
4. Information not available

Recently, an inhibitor of apoptosis (programmed cell death), survivin, has been detected in most forms of cancer. Survivin is necessary during mitosis to ensure normal cell division and chromosome distribution. However, in cancerous cells, survivin prolongs their life span, thereby increasing the chances of mutations. Research is being targeted at developing drugs to inhibit survivin expression and thus inhibit tumor growth. Clinical tests are not yet commercially available to measure blood levels of survivin.

3. Refer to Chapter 6 and Chapter 11 and to other tests in Chapter 8 for complete listings of normal values, clinical implications, and safe, effective, informed interpretations, patient preparation, and aftercare.

Interventions
Pretest Patient Preparation
1. Explain purpose and procedure of test.
2. Alleviate fears the patient may have related to cancer test results. Tests for cancer are always anxiety provoking.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test results using the latest knowledge in the field, recognizing that test values, significance, and specificity of tests are continually changing with technology. Generally, tumor markers are not helpful in predicting the site of origin.
2. Provide consultation if test results reveal cancer. Tumor drug resistant assays are performed on tissue obtained in biopsy (see Chap. 11).
3. Provide support through follow-up testing in stages of illness and in forming a therapeutic and diagnostic plan for treating and monitoring the disease.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care regarding shock, denial, and fear as normal response to cancer diagnosis.

BLOOD BANKING OR IMMUNOHEMATOLOGY TESTS
These tests are done to select blood components that will have acceptable survival when transfused and to prevent possible transplant and transfusion reactions; to identify potential problems, such as hemolytic disease of newborns and the need for intrauterine transfusion; and to determine parentage. Immunohematology testing identifies highly reactive antigens on blood cells and their antibodies possibly present in serum.

Donated Blood Testing and Blood Processing
Pretransfusion testing of blood recipient and donor blood:

1. All donated blood, as it is processed, must undergo several measurements. These include tests for the following factors:
   a. ABO groups
   b. Rh type
   c. Antibody screen
d. Hepatitis B surface antigen (HBsAg)
e. Hepatitis B core antigen (HBeAg)
f. Hepatitis C virus (anti-HCV)
g. Syphilis (VDRL)
h. HIV-1 and HIV-2
   i. HTLV-I and HTLV-II
   j. HIV antigen (HIV-1-Ag)
k. Nucleic acid tests (NAT) (narrow window of infection for HIV and hepatitis C)
2. Required testing for whole-blood or red blood cell recipients include the following:
   a. ABO group
   b. Rh type
   c. Antibody screen
   d. Crossmatch for compatibility between donor's cells and recipient's serum

Type and crossmatch, when ordered together, use one 10-mL red-topped tube.

Type and screen when ordered preoperatively, identify ABO and Rh type group and are used for cases that usually do not require transfusion. Even though no crossmatch is needed for plasma administration, compatible ABO typing should be done. Routinely, no crossmatch is needed for platelet administration; compatible ABO and Rh types should be given when possible. If a patient becomes refractory, HLA-matched platelets may be administered. Granulocytes should be tested for HLA compatibility. As a result of previous transfusions or pregnancy, some patients develop antibodies against these antigens and, if given platelets that have these antigens, may have a transfusion reaction. Pretransfusion testing for neonates (younger than 4 months of age) requires determination of ABO group, Rh type, and an antibody screen. The antibody screen may be performed on a specimen obtained from the infant or the mother. If the antibody screen is negative, group O Rh compatible pediatric red blood cells may be used without further crossmatching for the remainder of the neonatal period. If the antibody screen is positive, the antibody is identified, and antigen negative blood will be crossmatched and provided for transfusion. All infants requiring plasma transfusions will receive group AB pediatric fresh-frozen plasma.

A type and screen consists of an ABO group, Rh type, and antibody screen and can be ordered when the need for crossmatched products is unlikely but may be required in an emergency situation. If the patient does have a clinically significant unexpected antibody, at least 2 U of antigen-negative blood will be made available for that patient before surgery. A positive antibody screen will automatically initiate antibody identification to determine the specificity of the antibody detected, if the antibody identified is determined to be clinically significant, antigen-negative blood is required for transfusion.

Procedure

1. Collect a 10-mL sample for hospital pretransfusion testing. Use a 10-mL plain, red-topped tube.
2. Label with the following information:
   a. Patient's full first and last names
   b. Patient's health care record number
   c. Date and time of specimen collection
   d. Initials (if collected by laboratory personnel) or signature (if collected by nonlaboratory personnel) of phlebotomist
   e. Possibly a unique blood bank number (found on special Blood Bank identification band)
3. Attach a special Blood Bank band, at the recipient's bedside, at the time of specimen collection. The Blood Bank band must remain attached to the patient's wrist throughout the transfusion period. The same band may be used throughout one hospital admission as long as the information printed on the band is legible and the band is still securely attached to the patient's wrist.
4. Be aware that a new specimen is required every 3 days if the patient has a history of transfusion or pregnancy during the previous 3 months.

Special Considerations

1. Autologous donations are blood products donated by patients for their own use (ie, blood donor and recipient are the same person). Many patients opt to donate their own blood before scheduled surgery because of the concern regarding transfusion-transmitted diseases. The following are some general guidelines for autologous blood donation.
   a. There is no age limit if donor is healthy.
   b. There are no weight requirements. The volume of blood collected must comply with established weight provisions.
   c. Pregnant women can donate.
   d. Hematocrit should be >33%. If <33%, the patient's physician must approve the phlebotomy, usually in consultation with the blood bank medical director.
   e. Normally, phlebotomy can be done at 3-day intervals; the final phlebotomy can be done no sooner than 72 hours before the time of the scheduled surgery. Two-unit collections using an automated red cellopheresis machine may also be an option. Iron supplements may be prescribed to maintain adequate hemoglobin levels.
   f. Autologous blood is not “crossed over” into the general (allogegenic) blood supply. It is discarded after its expiration date.

2. Allogeneic donations are blood products donated by one individual for use by other individuals (ie, blood donor and blood recipient are not the same person).

3. Direct donations are those in which recipients choose those who donate blood for their transfusions. Laws in several states declare that this request must be honored in nonemergency situations. Standards and testing procedures must be identical to those required for an allogeneic blood donor. (Autologous donors do not need to adhere to the same criteria as do allogeneic blood donors). Directed donor units can be “crossed over” into the general (allogegenic) blood supply. Each establishment must have a policy describing when this can occur.

4. Cytomegalovirus (CMV) testing is done for patients at risk for transfusion-associated CMV infections. These types of CMV infections include pneumonia, hepatitis, retinitis, and disseminated infection. They generally occur in immunosuppressed patients, such as premature infants weighing <1200 g at birth, bone marrow and organ transplant recipients, and certain immunocompromised oncology patients. Therefore, to prevent these infections, CMV antibody testing is done. Patients at risk should receive CMV-seronegative blood and blood products. CMV in blood is associated with leukocytes. Leukocyte reduction using highly efficient leukocyte-reduction filters also appears to be an effective way of reducing CMV infection.

- **CMV antibody testing**
  - Done for patients at risk for transfusion-associated CMV infections.
  - Assesses the presence of CMV antibodies in the blood.
  - Helps identify patients who are CMV-seronegative, reducing the risk of CMV transmission during transfusion.

- **Leukocyte reduction**
  - Utilizes filters to remove leukocytes from blood products.
  - Minimizes the risk of transfusion-associated CMV infection.

- **Pretransfusion testing**
  - Includes crossmatch for compatibility between donor's cells and recipient's serum.
  - Ensures the blood is safe for transfusion.

- **Blood collection**
  - Specimen collection described for autologous and allogeneic donations.
  - Autologous donations are blood products donated by patients for their own use.
  - Allogeneic donations are blood products donated by one individual for use by other individuals.

- **Posttransfusion monitoring**
  - Follow-up testing and observation to monitor for adverse reactions.
  - Ensures patient safety after transfusion.
5. **Irradiation of blood products** is sometimes done before transfusion for certain immunosuppressed patients. Graft-versus-host disease (GVHD) is a rare complication that follows transfusion in severely immunosuppressed patients. GVHD occurs if donor lymphocytes from blood or blood products engraft and multiply in a severely immunodeficient recipient. The engrafted lymphocytes react against host (recipient) tissues. Clinical symptoms include skin rash, fever, diarrhea, hepatitis, bone marrow suppression, and infection, which frequently leads to death. GVHD can be prevented by irradiating blood products with a maximum dose (cesium-137) of 2.5 cGy in the center of the container and a minimum dose of 1.5 cGy delivered to all other parts of the component. This practice renders the T lymphocytes in a unit of blood incapable of replication without affecting platelets or granulocytes. Irradiation does affect the red cell membrane, causing it to “leak” potassium. All irradiated red cells are given a 28-day “outdate” or may keep their original “outdate” of <28 days.

6. **Leukocyte reduction of blood products:** Leukocytes in blood products have long been known to be associated with nonhemolytic febrile transfusion reactions, possibly owing more to cytokines produced by the leukocytes than the leukocytes themselves. Leukocyte reduction may reduce the number of these reactions. It may also decrease the possibility of alloimmunization to the HLA antigens on the leukocytes. Removing leukocytes may effectively reduce the danger of transfusion-transmitted CMV infection.

### Clinical Alert

Before transfusion, there is a requirement for physicians to document that all the alternatives to transfusion, risks of transfusion, and transfusion issues were explained to the patient. Examples of discussion include need for blood or blood components, risks (infection, disease, reactions, alloimmunization), benefits (treatment for active bleeding, anemia, clotting disorders), and alternatives to random allogenic donation (predeposit autologous donation, intraoperative salvage, directed donor donation). The physician must document in the health record that the discussion of these issues took place. The documentation varies from institution to institution. An example of informed consent used before transfusion is included in Appendix H. Allow sufficient time for patient to ask questions, resolve concerns, and to give voluntary consent. There must be documentation in the chart before the patient receives the transfusion.

### Education Alert

1. Information for potential blood donors can be obtained at local blood banks or the American Red Cross.
2. Blood donation begins with meeting requirements: age (at least 17 years), weight (at least 110 pounds), and healthy history that is negative for cancer and infectious diseases (no fever in past 2 weeks).
3. Here is how the process works at a Red Cross Blood Bank: A pint of blood is obtained, filling a bag and four vials. Blood is divided into three parts. Each part (red blood cells, platelets, and plasma) is prepared by spinning and the blood is stored by blood type until the product is used or expired. Blood is shipped to hospitals upon request. Local community blood banks also supply nearby hospitals. The donated pint of blood is also divided into many products that then sell for hundreds of dollars. Bags and products are quarantined until tests show no infection and are checked for antibodies that might cause a transfusion reaction. Unsuitable bags are destroyed.
4. Only donated blood in storage (that has previously been tested) actually helps disaster victims.

### Blood Groups (ABO Groups)

Human blood is grouped according to the presence or absence of specific blood group antigens (ABO). These antigens, found on the surface of red blood cells, can induce the body to produce antibodies. More than 300 distinct antigens have been identified. Compatibility of the ABO group is the foundation for all other pretransfusion testing.

All blood donors and potential blood recipients must be tested for blood type to prevent transfusion with incompatible blood products. Specifically linked sugars determine the antigenic activities named A and B. One sugar, N-acetylgalactosamine, gives the molecule A activity; another sugar, galactose, determines B activity. The backbone molecule, without galactose or N-acetylgalactosamine, has antigenic activity termed H. This H substance, as well as H gene activity, is essential for the function of the ABO antigens. Table 8.10 lists the blood groups and their ABO antigens.

<table>
<thead>
<tr>
<th>Blood Group ABO Antigen</th>
<th>A</th>
<th>B</th>
<th>AB</th>
<th>O</th>
</tr>
</thead>
</table>
In general, patients are transfused with blood of their own ABO group because antibodies against the other blood antigens may be present in their blood serum. These antibodies are designated anti-A or anti-B, depending on the antigen they act against. Under normal conditions, a person's blood serum does not contain the antibody specifically able to destroy its antigen. For example, a person with antigen A will not have anti-A antibodies in the serum; however, anti-B antibodies may be present. Therefore, antigen and antibody testing is necessary to confirm ABO grouping.

### Table 8.10 Antigen Values for Blood Types

<table>
<thead>
<tr>
<th>Blood Group ABO Antigen</th>
<th>A</th>
<th>B</th>
<th>AB</th>
<th>O</th>
</tr>
</thead>
</table>

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Clinical Alert

1. A transfusion reaction can be extremely serious and potentially fatal. Therefore, the blood group must be
determined in vitro before any blood is transfused to an individual. Before blood administration, two health care
professionals (ie, physicians or nurses) must check the recipient’s blood group and type with the donor group and
type to ensure compatibility.
2. A blood group change or suppression may be induced by cancer, leukemia, infection, or bone marrow transplant.

Reference Values

Normal A, B, AB, and O group

<table>
<thead>
<tr>
<th>Relationship Between Blood Antigens and Antibodies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antigen Present on Red Blood Cell</td>
</tr>
<tr>
<td>---------------------------------</td>
</tr>
<tr>
<td>None</td>
</tr>
<tr>
<td>A</td>
</tr>
<tr>
<td>B</td>
</tr>
<tr>
<td>AB</td>
</tr>
</tbody>
</table>

*Called universal donor because no antigens are present on red blood cells; therefore, the person is able to donate to all
blood groups.
†Called universal recipient because no serum antibodies are present; therefore, the person is able to receive blood from
all blood groups.
‡Called universal donor for plasma because no serum antibodies are present; therefore, the plasma can be given to all
blood groups.

Procedure

1. Collect a 7-mL venous clotted blood sample in a red-topped tube. Observe standard precautions.
2. Do not use SST tubes (cell barrier tube).

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure. The following are conditions that at some point may require transfusion:
   a. Malignant tumors (leukemias)
   b. Cardiac surgical procedures
   c. Surgical hip procedures
   d. Anemias
   e. Certain obstetric or gynecologic procedures or complications
   f. Bone and joint diseases
   g. Lung disease
   h. Kidney disease or genitourinary system surgical procedures
   i. Massive trauma
   j. Liver disease
   k. Certain blood dyscrasias
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Inform patient of blood group and interpret meaning. Rh type (see next section) may have implications for the
   pregnant woman and fetus.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Rh Typing

Human blood is classified as Rh positive or Rh negative. This relates to the presence or the absence of the D antigen
on the red cell membrane. The D antigen (now called Rh 1 [D]) is, after the A and B antigens, the next most important
antigen in transfusion practice.

Incidence and Frequency of Blood Group and Rh Type

<table>
<thead>
<tr>
<th>Group and Type Incidence Frequency of Occurrence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>O positive 1 in 3 37.4</td>
</tr>
<tr>
<td>O negative 1 in 15 6.6</td>
</tr>
<tr>
<td>A positive 1 in 3 35.7</td>
</tr>
<tr>
<td>A negative 1 in 16 6.3</td>
</tr>
<tr>
<td>B positive 1 in 12 8.5</td>
</tr>
<tr>
<td>B negative 1 in 67 1.5</td>
</tr>
<tr>
<td>AB positive 1 in 29 3.4</td>
</tr>
</tbody>
</table>
The Rh system is composed of antigens tested for in conjunction with the ABO group. Rh₁ (D) antigen is often the only factor tested for. When this factor is absent, further testing is then done on women of childbearing age to identify if there is Rh₁ (D) antigen present in smaller amounts. This test is called weak D (formally known as D testing). Rh-negative individuals may develop antibodies against Rh-positive antigens if they are challenged through a transfusion of Rh-positive blood or through a fetomaternal bleed from an Rh-positive fetus.

Reference Values

Normal

1. Caucasian
   a. 85% Rh positive [have the Rh(O) antigen]
   b. 15% Rh negative [lack the Rh(O) antigen]
2. African Americans
   a. 90% Rh positive [have the Rh(O) antigen]
   b. 10% Rh negative [lack the Rh(O) antigen]

Procedure

1. Be aware that blood Rh typing must be done for the following reasons:
   a. Rh-positive blood administered to an Rh-negative person may sensitize the person to form anti-D (Rh₁).
   b. Rh₁ (D)-positive blood administered to a recipient having serum anti-D (Rh₁) could be fatal.
2. Identify RhIG (Rh immunoglobulin) candidates. Rh immunoglobulin is a concentrated solution of IgG anti-D (Rh₁) derived from human plasma. A 1-mL dose of RhIG contains 300 µg and is sufficient to counteract the immunizing effects of 15 mL of packed red cells or 30 mL of whole blood.
   a. Rh-negative pregnant women with Rh-positive partners may carry Rh-positive fetuses. Fetal cells may cross the placenta to the mother and cause production of antibodies in the maternal blood. The maternal antibody, in turn, may cross through the placenta into the fetal circulation and cause destruction of fetal blood cells. This condition, called hemolytic disease of the newborn (formerly called erythroblastosis fetalis), may cause reactions that range from anemia (slight or severe) to fetal death in utero. This condition may be prevented if an Rh-negative pregnant woman receives an RhIG dose antepartum at 28 weeks’ gestation and a postpartum injection of RhIG shortly after delivery of an Rh₁ (D)-positive infant. Postpartum Rh immunization can occur despite an injection of RhIG if >30 mL of fetal blood enters the maternal circulation. The American Association of Blood Banks recommends that a postpartum blood specimen of all Rh₁ (D)-negative women (ie, those at risk for immunization) be examined to detect a fetal maternal hemorrhage of >30 mL.
   b. Rh typing and evaluation for RhIG must also be done for patients who have had abortions, miscarriages, accidents, and amniocentesis.
3. Observe standard precautions.

Comparison of Terms Used in Rh System Nomenclatures

<table>
<thead>
<tr>
<th>Weiner</th>
<th>Fisher-Race</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rh₁</td>
<td>D</td>
</tr>
<tr>
<td>Rh₂</td>
<td>C</td>
</tr>
<tr>
<td>Rh₃</td>
<td>E</td>
</tr>
<tr>
<td>Rh₄</td>
<td>c</td>
</tr>
<tr>
<td>Rh₅</td>
<td>e</td>
</tr>
<tr>
<td>Rh₆</td>
<td>f (ce)</td>
</tr>
<tr>
<td>Rh₁₂</td>
<td>G</td>
</tr>
</tbody>
</table>

Clinical Implications

1. The significance of Rh antigens is based on their capacity to immunize as a result of receiving a transfusion or becoming pregnant. The Rh₁ (D) antigen is by far the most antigenic; the other Rh antigens are much less likely to produce isoimmunization. The following general conditions must be met for immunization to Rh antigens to occur:
   a. The Rh blood antigen must be absent in the immunized person.
   b. The Rh blood antigen must be present in the immunizing blood.
   c. The blood antigen must be of sufficient antigenic strength to produce a reaction.
   d. The amount of incompatible blood must be large enough to induce antibody formation.
   e. Factors other than Rh₁ (D) may induce formation of antibodies in Rh-positive persons if the preceding conditions are met.
2. Antibodies for Rh₂ (C) are frequently found together with anti-Rh₁ (D) antibodies in the Rh-negative pregnant woman whose fetus or child is type Rh positive and possesses both antigens.
3. With exceedingly rare exceptions, Rh antibodies do not form unless preceded by antigenic stimulation, as occurs with the following conditions:
   a. Pregnancy and abortions
   b. Blood transfusions
c. Deliberate immunization, most commonly of repeated intravenous injections of blood for the purpose of harvesting a given Rh antibody

Interventions

Pretest Patient Preparation

1. Explain purpose and procedure of the typing.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome. Inform and counsel patient regarding Rh type. Women of childbearing age may need special consideration. See page 634 for incidences of Rh types.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Rh Antibody Titer Test

This antibody study determines the Rh antibody level in an Rh-negative or pregnant woman whose partner is Rh positive. If the Rh-negative woman is carrying an Rh-positive fetus, the antigen from the fetal blood cells causes antibody production in the mother's serum. The firstborn child usually shows no ill effects; however, with subsequent pregnancies, the mother’s serum antibodies increase and eventually destroy the fetal red blood cells, causing hemolytic disease of the newborn.

Reference Values

Normal Negative is 0 (no antibody detected)

Procedure

1. Obtain a 10-mL venous blood sample (plasma or serum) from the mother using a yellow-topped (ACD) and clotted blood (not SST) tube.
2. Observe standard precautions.

Clinical Implications Some institutions have established a critical titer for anti-D below which hemolytic disease of the newborn is considered unlikely. No further investigations are undertaken unless the critical titer level is reached.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel appropriately.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Rosette Test, Fetal Red Cells (Fetal-Maternal Bleed)

This qualitative test detects Rh-positive fetal cells in the Rh-negative maternal circulation. The detection of fetal erythrocytes is important when it is suspected that a severe fetal red cell loss has occurred and when serious risk for the mother becoming immunized against the fetal red cell groups is anticipated. In these instances, the mother's blood sample should be collected immediately after delivery to be examined for fetal cells. This test can be performed only if mother is Rh-negative and newborn is known to be Rh-positive. The Rosette test is 97% accurate for detecting a fetomaternal bleed that exceeds 30 mL of whole blood. This test cannot be performed on patients who have had abortions, miscarriages, accidents, or amniocentesis.

Reference Values

Normal Negative for fetal blood loss No Rh-positive fetal red blood cells detected in maternal blood

Procedure

1. Obtain a 7-mL venous blood EDTA sample from the mother shortly after delivery.
2. Perform this test and examine results for rosettes or mixed field agglutinates. Following manufacturer's guidelines, the presence of rosettes above a predetermined number indicates a fetal bleed that exceeds 30 mL of whole blood.

Clinical Implications When the test sample contains few or new Rh \( ^1 \) positive fetal cells, rosetting or agglutination is absent, and the fetomaternal bleed is <30 mL, one dose of parental RhIG will prevent immunization. If the fetal blood loss into the maternal circulation exceeds 30 mL, a quantitative or semi-quantitative test (ie, Kleihauer-Betke) or a quantitative flow cytometry (if available) must be performed to calculate the amount of RhIG to administer.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcome. Counsel patient regarding RhIG administration and follow-up maternal testing.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Kleihauer-Betke Test (Fetal Hemoglobin Stain)

The Kleihauer-Betke test is a semi-quantitative test to determine the amount of fetomaternal hemorrhage in an Rh₁-negative mother and the amount of RhIG necessary to prevent antibody production. The test is done after full-term delivery if newborn anemia is present or when the mother is Rh negative or weak-negative D. The test is also performed on mothers after invasive procedures (eg, amniocentesis), miscarriages, or traumas.

Reference Values

Negative: No fetal cells in maternal circulation

Procedure

1. A 7-mL maternal venous blood EDTA sample is obtained immediately after delivery, invasive procedure (eg, amniocentesis), miscarriage, or trauma.
2. Examine the specimen immediately or refrigerate until it can be examined.

Clinical Implications

1. Results indicate moderate to great fetomaternal hemorrhage (50%–90% of fetal red blood cells contain fetal hemoglobin [HbF]).
2. With full-term delivery, newborn red blood cells must be Rh-D-positive for the Rh-D-negative mother to be a candidate for RhIG.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately regarding fetal bleed and administration of RhIG to suppress the immunization of fetal red cells or whole-blood hemorrhage (Table 8.11). The calculated dose is as follows:

   \[
   \text{Vials of RhIG} = \frac{\text{mL of fetal blood}}{30}
   \]

<table>
<thead>
<tr>
<th>Fetal Cells (%)</th>
<th>Average</th>
<th>Range</th>
<th>Vials of RhIG to Inject</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.3–0.5</td>
<td>20</td>
<td>&lt;50</td>
<td>2</td>
</tr>
<tr>
<td>0.6–0.8</td>
<td>35</td>
<td>15–80</td>
<td>3</td>
</tr>
<tr>
<td>0.9–1.1</td>
<td>50</td>
<td>22–110</td>
<td>4</td>
</tr>
<tr>
<td>1.2–1.4</td>
<td>65</td>
<td>30–140</td>
<td>5</td>
</tr>
<tr>
<td>1.5–2.0</td>
<td>88</td>
<td>37–200</td>
<td>6</td>
</tr>
<tr>
<td>2.1–2.5</td>
<td>115</td>
<td>52–250</td>
<td>6</td>
</tr>
</tbody>
</table>

   *The range provides for the poor precision of the acid separation elution test. These recommendations are based on one vial needed for each 15 mL of red blood cells or 30 mL of whole blood.

   Many recommend doubling the calculated dose of RhIG. The method of calculating fetal blood is not entirely accurate. The results of undertreatment are serious, but the effects of overtreatment are minor.

2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Crossmatch (Compatibility Test)

The primary purpose of the major crossmatch, or compatibility test, is to prevent a possible transfusion reaction.

Major crossmatch detects antibodies in the recipient's serum that may damage or destroy the cells in the blood donor (Table 8.12). The type and screen determines the ABO and Rh-D type as well as the presence or absence of unexpected antibodies from the recipient. The type and screen is a safe alternative for the routine type and crossmatch ordered.
preoperatively for cases that may, but usually do not, require transfusion (e.g., hysterectomy, cholecystectomy). If blood is needed, a major crossmatch must be done before transfusion.

Table 8.12 **Antibodies Found in Crossmatching**

<table>
<thead>
<tr>
<th>Blood Grouping System</th>
<th>Antibody</th>
<th>Description</th>
</tr>
</thead>
</table>
| Rh-hr                 | Anti-D   | Rh1
|                       |          | May cause severe hemolytic disease of newborn |
|                       | Anti-C   | Rh2
|                       |          | Often found with anti-D, -Ce(rh<sup>+</sup>) or C<sup>W</sup> |
|                       | Anti-E   | Rh3
|                       |          | Often found with anti-
|                       | Anti-c   | Rh4
|                       |          | Often found with anti-E |
|                       | Anti-e   | Rh5
|                       |          | Often found with anti-C |
|                       | Anti-C<sup>W</sup> | Rh8 |
|                       | Anti-V   | Rh10
|                       |          | Alternative antigen names: ce<sup>5</sup>, hr<sup>5</sup> |
| Kell                 | Anti-K   | K1
|                       |          | Strongly immunogenic; some non–red cell immune |
|                       | Anti-k   | K2
|                       |          | Occasional Kell system antibodies may not react |
|                       | Anti-Kp<sup>a</sup> | K3 |
|                       |          | Antigen may be depressed by the presence of Kp<sup>a</sup> |
|                       | Anti-Kp<sup>b</sup> | K4 |
|                       |          | Few non–red cell immune |
|                       | Anti-Js<sup>a</sup> | K6 |
|                       |          | Few non–red cell immune |
|                       | Anti-Js<sup>b</sup> | K7 |
| Duffy                | Anti-Fy<sup>a</sup> | Some antibodies exhibit dosage; quite common and may cause HDN and HTRs |
| Kidd                 | Anti-Fy<sup>b</sup> | Some antibodies may bind complement |
|                     | Anti-Jk<sup>a</sup> | Antibodies may exhibit dosage |
|                     | Anti-Jk<sup>b</sup> | May cause severe delayed hemolytic transfusion reactions |
|                     |          | Antibody titers may drop rapidly below detectable levels |
| Lutheran             | Anti-Lu<sup>a</sup> | Antibodies may require anti-C3 for detection |
|                     | Anti-Lu<sup>b</sup> | Antibody gives mixed field-like agglutination |
| MN                   | Anti-M   | Common antibody |
|                     | Anti-N   | Rare antibody |
|                     | Anti-S   | Formaldehyde-induced anti-N commonly found in dialysis patients |
|                     | Anti-s   | Antibody may be enhanced if incubated below 37°C before AHG |
|                     | Anti-U   | Rarely found in S-, s-patients |
| Lewis                | Anti-Le<sup>a</sup> | Frequently found in serum of pregnant women |
|                     | Anti-Le<sup>b</sup> | Neutralized by soluble antigen |
|                     | -Le<sup>b</sup><sup>h</sup> | Anti-Le<sup>b</sup> often found with anti-Le<sup>a</sup> |
|                     | -Le<sup>b</sup><sup>L</sup> | Anti-Le<sup>b</sup> usually made by Le<sup>a</sup> (a ‘b’ ) individuals |
| P                    | Anti-P<sub>1</sub> | Antigen strength variable; neutralized by soluble antigen |
|                     | Anti-P   | Biphasic hemolytic IgG autoantibody in PCH |
|                     |          | Altoantibody is usually potent IgM hemolysin |
|                     | Anti-P<sup>k</sup> (Anti-T<sub>ja</sub>) | Have caused hemolytic transfusion reactions and occasionally HDN |
| Xg                   | Anti-Xg<sup>a</sup> | X-linked |
| Colton               | Anti-Co<sup>a</sup> | Rare antibodies |
| Dombrock             | Anti-Do<sup>a</sup> | Incidence of Do<sup>a</sup> lower in African Americans, Native Americans, and Asians |
|                     | Anti-Do<sup>b</sup> | Infrequently reported antibodies |
Diego  
Anti-Di $^a$  Di $^a$ antigen frequently higher in Asians and Native Americans  
Anti-Di $^b$  
Wright  
Anti-Wr $^a$  IgM and IgG forms of antibody reported  
Frequently occurring antibody  
Vel  
Anti-Vel  Antibodies usually IgM; antigen strength variable, binds complement  
Sd $^a$  Antigen weaker during pregnancy  
Wide variation of antigen expression  
Agglutinates have refractile, mixed-field appearance  
HLA-associated  
Anti-Bg $^a$  Antigen strength variable  
-Bg $^b$  Antibodies often found in multitransfused multiparous patients  
-Bg $^c$  Antibodies characteristically weakly reactive  
Bg/HLA associations  
Bg $^a$/HLA-B7  
Bg $^b$/HLA-B17  
Bg $^c$/HLA-A28  
Cartwright  
Anti-Yt $^a$  Antibody not uncommon in Yt (a–) individuals  
Anti-Yt $^b$  Rare antibody usually found in combination with other antibodies  
HTLA  
Anti  
-Ch $^a$  Antigen strength variable  
-Kn $^a$  Antibodies characteristically weakly reactive  
-McC $^a$  
-Yk $^a$  
-Cs $^a$  
-Gy $^a$  
-Hy  
-JMH  
Anti-I  Most frequently detected cold autoagglutination  
Anti-I in CHD has wide thermal range, high titer  
Binds complement  
Seen as alloantibody in i adults  
Anti-i  Antibody seen in serum of patients with infectious mononucleosis  
Rare cause of CHD  
Antigen very weakly expressed on the cells of most adults  
HDN, hemolytic disease of the newborn; CHD, cold hemagglutinin disease; HTR, hemolytic transfusion reaction; AHG, antihuman globulin; Ig, immunoglobulin; PCH, paroxysmal cold hemoglobinuria; HTLA, human T-lymphocyte antigen.

Clinical Alert
Even the most carefully performed crossmatch will not detect all possible incompatible sources.

Reference Values
Normal  No cell clumping or hemolysis, and absence of agglutination when serum and cells are appropriately mixed and incubated  The major crossmatch shows compatibility between recipient serum and donor cells.

Procedure
1. Obtain a 10-mL venous blood sample.  
2. Observe standard precautions.

Clinical Implications
1. Crossmatch incompatibility implies that the recipient cannot receive the incompatible unit of blood because antibodies are present.  
2. A transfusion reaction occurs when incompatible blood is transfused, specifically if antibodies in the recipient's serum cause rapid red blood cell destruction in the proposed donor.  
a. Certain antibodies, although not causing immediate red cell destruction and transfusion reaction, may nevertheless reduce the normal life span of transfused incompatible cells; this may necessitate subsequent transfusions.  
b. The patient will derive the most benefit from red cells that survive longest.
Clinical Alert

1. The most common cause of hemolytic transfusion reaction is the administration of incompatible blood to the recipient because of faulty matching in the laboratory, improper patient identification, and/or incorrect labeling of donor blood. If a transfusion reaction is suspected, discontinue the transfusion and notify the blood bank and attending physician immediately.

2. Assess for the following symptoms of transfusion reaction:
   a. Fever
   b. Chills
   c. Chest, abdomen, or flank pain
   d. Hypotension or hypertension
   e. Nausea
   f. Dyspnea
   g. Shock
   h. Oliguria
   i. Back pain
   j. Feeling of heat along vein being transfused
   k. Constricting chest and lumbar back muscles
   l. Facial flushing
   m. Hemoglobinuria
   n. Oozing blood from wounds
   o. Anemia
   p. Allergic reactions such as local erythema, hives, and itching

3. The probable benefits of each blood transfusion must be weighed against the risks, which include the following:
   a. Hemolytic transfusion reactions due to infusion of incompatible blood (can be fatal)
   b. Febrile or allergic reactions
   c. Transmission of infectious disease (eg, hepatitis)

Interventions

Pretest Patient Preparation

1. Explain purpose and procedure of crossmatching.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel patient regarding potential transfusion reactions.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Coombs' Antiglobulin Test

The indirect Coombs' test detects antibodies that react only through a potentiating medium. The direct Coombs' test detects antigen-antibody complexes on the red blood cell membrane in vivo as well as red blood cell sensitization. It is diagnostic for the following conditions:

1. Hemolytic disease of the newborn in which the red cells of the infant are sensitized and exhibit antigen-antibody complexes in vivo
2. Acquired hemolytic anemia in which an antibody is produced that coats the patient's own cells (autosensitization in vivo)
3. Transfusion reaction in which the patient may have received incompatible blood, which in turn has sensitized the donor's and possibly the patient's own red cells
4. Red blood cell sensitization caused by drugs

The indirect Coombs' test detects serum antibodies, reveals maternal anti-Rh antibodies during pregnancy, and can detect incompatibilities not found by other methods.

Reference Values

Normal

Direct Coombs' test: negative for red blood cells
Indirect Coombs' test: negative for serum

Procedure

1. Be aware that the direct Coombs' test is positive (1+ to 4+) in the presence of the following conditions:
   a. Transfusion reactions
   b. Autoimmune hemolytic anemia (most cases)
   c. Cephalothin therapy (75% of cases)
   d. Drugs such as a-methyldopa (Aldomet), penicillin, insulin
   e. Hemolytic disease of newborn
   f. Paroxysmal cold hemoglobinuria (PCH)
2. Be aware that the direct Coombs' test is positive (1+ to 4+) in the presence of specific antibodies, usually from a previous transfusion or pregnancy, or nonspecific antibodies, as in cold agglutinants.
3. Observe standard precautions.

Interfering Factors A number of drugs may cause the direct Coombs' test to be positive.
Clinical Alert

Antibody identification is performed when the antibody screen or direct antiglobulin tests produce positive results and unexpected blood group antibodies need to be classified. Antibody identification tests are an important part of pretransfusion testing so that the appropriate antigen-negative blood can be transfused. These tests are also helpful for diagnosing hemolytic disease of the newborn and autoimmune hemolytic anemia. A 7-mL venous blood sample with added EDTA and 20 mL of clotted blood are studied. Notify the laboratory of diagnosis, history of recent and past transfusions, pregnancy, and any drug therapy.

Interventions

Pretest Patient Preparation

1. Explain purpose and procedure of test.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel appropriately. Hemolytic disease of newborn can occur when the mother is Rh negative and the fetus is Rh positive. Diagnosis is derived from the following information: mother is Rh negative, newborn is Rh positive, and the direct Coombs’ test is positive. Newborn jaundice results from Rh incompatibility, but more often, the jaundice results from an ABO incompatibility.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

TYPES OF TRANSFUSION REACTIONS

Acute Hemolytic Transfusion Reaction (HTR)

HTR reaction is triggered by an antigen-antibody reaction and activates the complement and coagulation systems. These are most always due to ABO incompatibility because of misidentification resulting in the patient receiving incompatible blood. Symptoms include fever, chills, backache, vague uneasiness, and red urine. HTR reactions are potentially fatal.

Bacterial Contamination

Bacteria may enter the blood during phlebotomy. These microbes will multiply faster in components stored at room temperature than in refrigerated components. Although rare, bacteria in blood or its components can cause a septic transfusion reaction. Symptoms include high fever, shock, hemoglobinuria, DIC, and renal failure. Such reactions can be fatal.

Cutaneous Hypersensitivity Reactions

Urticarial reactions are very common, second in frequency only to febrile nonhemolytic reactions and are usually characterized by erythema, hives, and itching. Allergy to some soluble substance in donor plasma is suspected.

Noncardiogenic Pulmonary Reactions (NPR)

Transfusion-related acute lung injury (TRALI) should be considered whenever a transfusion recipient experiences acute respiratory insufficiency and/or x-ray films show findings consistent with pulmonary edema without evidence of cardiac failure. These are possibly reactions between the donor's leukocyte antibodies and the recipient's leukocytes. TRALI produces white cell aggregates that become trapped in the pulmonary microcirculation. The findings on chest x-ray films are typical of acute pulmonary edema. If subsequent transfusions are needed, leukocyte-reduced red cells may prevent NPR reactions.

Febrile Nonhemolytic (FNH) Reactions

FNH reactions are defined as a temperature increase of >1°C. They are seldom dangerous and may be caused by an antibody-antigen reaction.

Anaphylactic Reactions

Anaphylactic reactions occur after infusion of as little as a few milliliters of blood or plasma. Anaphylaxis is characterized by coughing, bronchospasm, respiratory distress, vascular instability, nausea, abdominal cramps, vomiting, diarrhea, shock, and loss of consciousness. Some reactions occur in IgA-deficient patients who have developed anti-IgA antibodies after immunization through previous transfusion or pregnancy.

Circulatory Overload

Rapid increases in blood volume are not tolerated well by patients with compromised cardiac or pulmonary function. Symptoms of circulatory overload include coughing, cyanosis, orthopnea, difficulty breathing, and a rapid increase in systolic blood pressure.

Leukoagglutinin Test

Leukoagglutinins are antibodies that react with white blood cells and sometimes cause febrile, nonhemolytic transfusion reactions. Patients who exhibit this type of transfusion reaction should receive leukocyte-poor blood for any subsequent
transfusions.

This study is done when a blood reaction occurs even though compatible blood has been given. The donor plasma contains an antibody that reacts with recipient white cells to produce an acute clinical syndrome of fever, dyspnea, cough, pulmonary infiltrates, and in more severe cases, cyanosis and hypertension. Patients immunized by previous transfusions, pregnancy, or during allografts often experience these febrile, nonhemolytic transfusion reactions because of incompatible transfused leukocytes. This type of reaction must be confirmed (as compared with hemolytic reactions) before additional transfusions can be safely administered.

Reference Values

**Normal** Negative for leukoagglutinins

**Procedure**

1. Obtain a 10-mL venous blood sample.
2. Observe standard precautions.

**Clinical Implications**

1. Agglutinating antibodies may appear in the donor's plasma.
2. When the agglutinating antibody appears in the recipient's plasma, febrile reactions are common; however, pulmonary manifestations do not occur.
3. Febrile reactions are more common in pregnant women and in individuals with a history of multiple transfusions.

**Clinical Alert**

1. Febrile reactions can be prevented by separating out white cells from the donor blood before transfusion.
2. Patients whose blood contains leukoagglutinins should be instructed that they generally need to be transfused with leukocyte-reduced blood to minimize these reactions.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome and counsel patient regarding future transfusion precautions.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Platelet Antibody Detection Test**

Platelet antibody detection studies are used to diagnose posttransfusion purpura, alloimmune neonatal thrombocytopenic purpura, idiopathic thrombocytopenia purpura, paroxysmal hemoglobinuria, and drug-induced immunologic thrombocytopenia.

**Reference Values**

**Normal** PLAI (negative platelet hyperlysibility): negative ALTP (negative drug-dependent platelet antibodies): negative PAIgG (platelet-associated IgG antibody): negative

**Procedure**

1. Obtain a 10-mL to 30-mL venous blood sample. Obtain 30 mL of venous blood when platelet count is 50,000–100,000/mm$^3$; 20 mL of venous blood when platelet count is 100,000–150,000 mm$^3$; and 10 mL of venous blood when platelet count is >150,000/mm$^3$.
2. Use standard precautions.

**Interfering Factors**

1. Alloantibodies formed in response to previous blood transfusions during pregnancies may produce positive reactions. Such antibodies are usually specific for human leukocyte antigens (HLAs) found in platelets and other cells.
2. Whenever possible, obtain samples for platelet antibody testing before transfusion.

**Clinical Implications**

1. Antibodies to platelet antigens are of two types: Autoantibodies develop in response to one's own platelets as in idiopathic thrombocytopenia purpura, and alloantibodies develop following exposure to foreign platelets during transfusion.
2. Antiplatelet antibody, usually having anti-PLAI specificity, occurs in posttransfusion purpura.
3. A persistent or rising antibody titer during pregnancy is associated with neonatal thrombocytopenia.
4. PLAI incompatibility between mother and fetus appears to account for >60% of alloimmune neonatal thrombocytopenic purpura. A finding of a PLAI-negative mother and a PLAI-positive father provides presumptive diagnostic evidence.
5. Platelet-associated IgG antibody (PAIgG) is present in 95% of both acute and chronic cases of idiopathic
(autoimmune) thrombocytopenic purpura. Patients responding to steroid therapy or undergoing spontaneous remission show increased circulatory times that correlate with decreased PAIgG levels.

6. The platelet hyperlysibility assay measures the sensitivity of platelets to lysis. This test is positive in and specific for paroxysmal hemoglobinuria.

7. In drug-induced immunologic thrombocytopenia, antibodies that react only in the presence of the inciting drug can be detected. Quinidine, quinine, chloroquine, sulfonamides, and diphenylhydantoin most commonly cause this type of thrombocytopenia. Gold-dependent antibodies and heparin-dependent platelet IgG antibodies can be detected by direct assay. Approximately 1% of persons receiving gold therapy develop thrombocytopenia as a side effect. Thrombocytopenia is also a well-known side effect of heparin.

NOTE

Platelet compatibility typing is done to ensure that hemostatically stable platelets can be transfused (eg, for aplastic anemia and malignant disorders). This is important because most patients repeatedly transfused with platelets from random donors become partially or totally refractory to further platelet transfusion because of alloimmunization. Platelet typing also provides diagnostic evidence of posttransfusion purpura. Platelets are routinely typed for PLA1, HLH-A2, and ALEI. Those matched for HLA antigens generally produce satisfactory posttransfusion improvement. A standard platelet count performed 1 hour after the end of a fresh platelet concentrate transfusion is a sensitive indicator for the presence or absence of clinically important antibodies against HLA antigens.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes. Appropriately counsel and monitor patient for bleeding tendencies. Assess for prescribed medications as cause of purpura.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Human Leukocyte Antigen (HLA) Test

The major histocompatibility antigens of humans belong to the HLA system. They are present on all nucleated cells but can be detected most easily on lymphocytes. Each antigen results from a gene that shares a locus on the chromosome with another gene, one paternal and one maternal (two alleles). More than 27 of these antigens have been identified. The HLA complex, located in the short arm of chromosome 6, is a major histocompatibility complex that is responsible for many important immune functions in humans.

This test determines the leukocyte antigens present on human cell surfaces. When tissue or organ transplantation is contemplated, HLA typing identifies the degree of histocompatibility between donor and recipient. By matching donors and potential recipients with compatible lymphocytes and similar HLA types, it is possible to prolong transplant survival and to reduce rejection episodes. The HLA also aids in diagnosis of parentage as well as correlation with certain disease syndromes and rheumatoid diseases, particularly ankylosing spondylitis. HLA-B27, one of the HLA antigens, is found in 90% of patients with ankylosing spondylitis. Generally, the presence of a certain HLA antigen may be associated with increased susceptibility to a specific disease; however, it does not mandate that that person will develop the disease. This test is also done before HLA-matched platelet transfusion.

Reference Values

Normal Requires clinical correlation

Procedure

1. Obtain a 10- to 24 mL (two green-topped tubes) heparinized venous blood sample in three lavender-topped EDTA tubes (14 mL) or two plain red-topped tubes, 10 mL minimum, or 5 mL of clotted blood or two yellow-topped (ACD) tubes.
2. Observe standard precautions.
3. Determine the patient's HLA type by testing the patient's lymphocytes against a panel of defined HLA antisera directed against the currently recognized HLA antigens. The HLA antigens are identified by letter and number. When viable human lymphocytes are incubated with a known HLA cytotoxic antibody, an antigen-antibody complex is formed on a cell surface. The addition of serum that contains complement kills the cells, which are then recognized as possessing a defined HLA antigen.
4. Label carefully with patient's name, date and special laboratory number. Include diagnosis and history.

Clinical Alert

Some of these tests may not be FDA approved.

Clinical Implications

1. Particular HLA antigens are associated with certain disease states:
   a. Ankylosing spondylitis (HLA-B27)
   b. Multiple sclerosis (HLA-B27 + Dw2 + A3 + B18)
   c. Sarcoidosis (HLA-B8)
   d. Psoriasis (HLA-A13 + B17)
e. Reiter’s syndrome (B27)
f. Juvenile insulin-dependent diabetes (Bw15 + B8)
g. Acute anterior uveitis (B27)
h. Graves’ disease (B27)
i. Juvenile RA (B27)
j. Celiac disease (B8)
k. Autoimmune chronic active hepatitis (B8)

2. Four groups of cell surface antigens (HLA-A, HLA-B, HLA-C, and HLA-D) constitute the strongest barriers to tissue transplantation.

3. In parentage determination, if a reputed father presents a phenotype (genotype completely determined by heredity; two haplotypes or gene clusters, one from father and one from mother) with no haplotype or antigen pair identical with one of the child's, he is excluded as the supposed father. If one of the reputed father's haplotypes (gene clusters) is the same as one of the child's, he may be the father. The chances of his being accurately identified as the father increase in direct proportion to the rarity of the presenting haplotype in the general population. Put another way, if the haplotype is very common, there is an increased probability that another man with the same haplotype also could be the father. When the frequency of the haplotype is known, the probability that the nonexcluded man is the father can be calculated. However, the degree of certainty diminishes as the incidence of the haplotype increases.

Interventions

Pretest Patient Preparation

1. Explain HLA test purposes and procedure. It is also used for postmortem testing before a renal transplantation.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately. HLA testing is best used as a diagnostic adjunct and should not be considered as diagnostic by itself. Explain the need for possible further testing.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

ORGAN AND TISSUE TRANSPLANT TESTING

The number of transplanted organs and tissues has been increasing yearly (>23,000 in the U.S. in 2002), with an ever-growing number of patients (>80,000 in the U.S. in 2003) on the waiting list for a transplant. The types and number of required diagnostic tests and procedures on the donor and recipient have been concomitantly increasing (Table 8.13).

Table 8.13 Examples of Laboratory Tests Performed on the Donor and Recipient for Organ Transplant *

<table>
<thead>
<tr>
<th>Tests of Donor and Recipient **</th>
<th>Lung</th>
<th>Liver</th>
<th>Kidney</th>
<th>Pancreas</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blood type</strong></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<tr>
<td><strong>HLA typing</strong></td>
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<tr>
<td><strong>Bilirubin</strong></td>
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<td>X</td>
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<tr>
<td><strong>BUN</strong></td>
<td>X</td>
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<td><strong>WBC</strong></td>
<td>X</td>
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<tr>
<td><strong>CBC</strong></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
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<tr>
<td><strong>Platelets</strong></td>
<td>X</td>
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<tr>
<td><strong>Hgb/Hct</strong></td>
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<tr>
<td><strong>PT/PTT</strong></td>
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<tr>
<td><strong>Electrolytes</strong></td>
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<tr>
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<tr>
<td><strong>Magnesium</strong></td>
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<tr>
<td><strong>Phosphorous</strong></td>
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<tr>
<td><strong>Creatinine</strong></td>
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<td><strong>Serum amylase</strong></td>
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<tr>
<td><strong>Albumin</strong></td>
<td>X</td>
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<tr>
<td><strong>Total protein</strong></td>
<td>X</td>
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<tr>
<td><strong>ALT</strong></td>
<td>X</td>
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<tr>
<td><strong>AST</strong></td>
<td>X</td>
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<tr>
<td><strong>Hepatitis A</strong></td>
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<tr>
<td><strong>Hepatitis B</strong></td>
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<tr>
<td><strong>Hepatitis C</strong></td>
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<tr>
<td><strong>Liver panel</strong></td>
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</tr>
<tr>
<td><strong>Epstein-Barr virus</strong></td>
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<td>X</td>
<td></td>
<td>X</td>
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<tr>
<td><strong>Anti-HIV 1 and 2</strong></td>
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<td>X</td>
<td>X</td>
<td>X</td>
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<tr>
<td><strong>Anti-HTLV 1</strong></td>
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<tr>
<td><strong>PRA</strong></td>
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<tr>
<td><strong>VDRL/RPR</strong></td>
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<td></td>
<td>X</td>
</tr>
</tbody>
</table>
Pretest Donor

Obtain a pertinent history of cancer, foreign travel, collagen and/or immune complex diseases, past exposure to certain infectious diseases, trauma, social history for high-risk behaviors, exposure to drugs, toxic substances, or biological hazards. The goal of testing is to prevent transmission of bacterial, viral, or genetic disease and to provide the best quality donor organ or tissue.

Pretest and Posttest of Recipient

1. A review of health history that includes past and present infection, malignancies, neurodegenerative disease, recipient of human pituitary gland hormones, high-risk behaviors (intravenous drug use, jaundice, and/or external infections).
2. Obtain a profile of the immunological risk factors, identification of HLA antigens to be avoided, or note irrelevant antigens. The HLA match varies for each organ or tissue. The goal is to have a complete match with the donor, prevent rejection, and identify acute rejection so that aggressive antirejection modalities can be started.
3. The presence of cytokines, c-reactive protein, complement, infections (bacterial, viral, or fungal), accelerated vascular disease, metabolic disease and/or diabetes all have an effect on a greater risk of transplant rejection in the recipient.

BIBLIOGRAPHY

Bader TF: Viral Hepatitis Practical Evaluation and Treatment. Seattle, WA, Hogrete & Huber, 1997
Cold agglutinins-positive results occur in other diseases/conditions. ASCP Lab Med 33(11): 8–15, 2001
Department of Health and Human Services: Exposure to Blood: What Health-Care Workers Need to Know. Rockville, MD, Author. 1999
Kiesner F, Brozel M: Transamerica Medicare extends full coverage to the Oncotech extreme drug resistance (EDR) assay. Press Release,
Kiesner F: Oncotech and the Johns Hopkins Cancer Center Institute Research collaboration to eliminate unnecessary exploratory surgery. Press Release, November 2, 2001


Loesmer F: Oncotech announces collaborative research with Southwest Oncology Group (SWOG) to evaluate the recently patented Oncotech radiation resistance assay as a predictor of clinical outcome in cervical carcinoma patients. Press Release, November 9, 2001


Sheehan C: Clinical Immunology, Principles and Laboratory Diagnosis, 2nd ed. Philadelphia, Lippincott-Raven, 1997


OVERVIEW OF NUCLEAR MEDICINE STUDIES

Nuclear medicine is a diagnostic modality that studies the physiology or function of any organ system in the body. Other diagnostic imaging modalities, such as ultrasound, magnetic resonance imaging (MRI), computed tomography (CT), and x-ray, generally visualize anatomic structures.

A pharmaceutical is labeled with a radioactive isotope to form a radiopharmaceutical. The radioisotope emits gamma and positron rays. Radioisotopes are reactor produced (iodine-131 [\(^{131}\)I]), cyclotron produced (fluorine-18 [\(^{18}\)F] for positron emission tomography [PET]), or generator produced (technetium-99m [\(^{99m}\)Tc]).

To visualize the function of an organ system, a radiopharmaceutical is administered. A time delay may be required for the radiopharmaceutical to reach its target site(s), and then the organ of interest is imaged with a gamma camera. Image formation technology involves the detection with very great density of a signal (gamma rays) emanating from the radiopharmaceutical. There is very little signal in the image that does not come from the radiopharmaceutical. The normal background level of radiation within the human body is minimal, with small amounts of radioactive potassium and some cesium. Routes of radiopharmaceutical administration vary with the specific study. Most commonly, a radiopharmaceutical is injected through a vein in the arm or hand. Other routes of administration include the oral, intramuscular, inhalation, intrathecal, subcutaneous, and intraperitoneal routes.

Nuclear medicine studies are performed by certified nuclear medicine technologists, interpreted by radiologists or nuclear medicine physicians, and performed in a hospital or clinic-based nuclear medicine department. The collaborative approach to care is evidenced by interventions from pharmacists, laboratory personnel, and nurses, among others.
Principles of Nuclear Medicine

The radiopharmaceutical is generally made up of two parts—the pharmaceutical, which is targeted to a specific organ; and the radionuclide, which emits gamma rays and allows the organ to be visualized by the gamma camera. Nuclear medicine imaging can yield quantitative as well as qualitative data. A measurement of the ejection fraction of the heart is an example of quantitative data derived from a multigated acquisition (MUGA) or a myocardial stress procedure.

In general, nuclear medicine images visualize the distribution of a particular radiopharmaceutical, with hot spots or cold spots of activity indicating an abnormality. In a hot spot, there is an increased area of uptake of the radiopharmaceutical in diseased tissue compared with the distribution in normal tissue. Examples of this type of uptake can be seen on bone images. In a cold spot, there is an area of decreased uptake of the radiopharmaceutical compared with the distribution in normal tissue. Liver and lung imaging are examples of this type of uptake. Prompt uptake in transplanted organs correlates with 1) adequate perfusion, eg, reperfusion of the transplanted lungs or pancreas; 2) excretory function, eg, in kidney transplants; and 3) evidence of cardiac viability and reinnervation. Poor uptake and nonvisualization of the transplanted organ are evidence of rejection.

Principles of Imaging

Gamma cameras all have basically the same components. The camera may have 1, 2, or 3 heads, with the capability of imaging in multiple configurations. The camera is networked with a multitasking computer capable of acquiring and processing the data.

Several methods of imaging are used: dynamic, static whole-body, and single photon emission computed tomography (SPECT). These imaging capabilities are available on all current camera systems.

Dynamic imaging allows serial display of multiple frames of data, each frame lasting 1 to 3 seconds, to visualize the blood flow associated with a particular organ. Static imaging is also known as planar imaging. The camera acquires one image at a time, covering the field of view. This image is 2-dimensional. Whole-body imaging acquires both anterior and posterior sweeps of the patient's body. This type of imaging also gives 2-dimensional information.

SPECT imaging has revolutionized the field of nuclear medicine. SPECT imaging provides three dimensions of data. SPECT imaging increased the specificity and sensitivity of nuclear imaging through improved resolution and is often combined with CT scans.

General Procedure

1. Alert the patient that he or she may be required to follow a study-specific preparation regimen before imaging determined by the type of nuclear medicine procedure (eg, nothing by mouth, no caffeine for 24 hours, hydration, bowel preparation).
2. Administer a radiopharmaceutical through one of several routes: oral, inhalation, intravenous, intramuscular, intrathecal, or intraperitoneal. On occasion, additional pharmaceuticals may be administered to enhance the function of the organ of interest.
3. Be aware that a time delay may be necessary for the radiopharmaceutical to reach the organ of interest.
4. Remember that imaging time depends on:
   a. Specific study radiopharmaceutical used and the time that must be allowed for concentration in tissues
   b. Type of imaging equipment used
   c. Patient cooperation
   d. Additional views based on patient history and nuclear medicine protocol
   e. Patient's physical size

Benefits and Risks

Benefits and risks should be explained before testing. Patients retain the radioisotope for a relatively short period. The radioactivity decays over time. Some of the radioisotope is eliminated in urine, feces, and other body fluids.

99mTc, the most commonly used radiopharmaceutical, has a radioactive half-life of 6 hours. This means that half of the dose decays in 6 hours. Other radioisotopes, such as iodine, indium, thallium, and gallium, take from 13 hours to 8 days for half of the dose to decay.

1. Benefits
   a. Nuclear medicine yields functional data that are not provided by other modalities.
   b. Nuclear imaging is relatively safe, painless (except for intravenous administration), and noninvasive.

2. Risks
   a. Radiation exposure is minimal; toxicity is nil.
   b. Hematoma at intravenous injection site.
   c. Reactions to the radiopharmaceutical (hives, rash, itching, constriction of throat, dyspnea, bronchospasm, anaphylaxis).

Clinical Considerations The following information should be obtained before diagnostic nuclear imaging:

1. Pregnancy (confirmed or suspected). Pregnancy is a contraindication for most nuclear imaging.
2. Lactating women may be advised to stop nursing for a set period (eg, 2 to 3 days with 99mTc). Most radiopharmaceuticals are excreted in the mother's milk.
3. Radiopharmaceutical uptake from a recent nuclear medicine examination could interfere with interpretation of the current study.
4. The presence of any prostheses in the body must be recorded on the patient’s history because certain devices can shield the gamma rays from imaging.
5. Current medications, treatments, and diagnostic measures (eg, telemetry, oxygen, urine collection, intravenous lines)
6. Age and current weight. This information is used to calculate the radiopharmaceutical dose to be administered. If the patient is younger than 18 years of age, notify the examining department before testing. The amount of radioactive substance administered is adjusted downward for anyone younger than 18 years of age.
7. Allergies. Past history of allergies, especially to contrast substances (eg, iodine) used in diagnostic procedures

**Clinical Alert**
The nuclear medicine department must be notified if the patient may be pregnant or is breast-feeding or is younger than 18 years of age.

**Interventions**

**Pretest Patient Care and Standard Precautions for Nuclear Medicine Scans**

1. Explain the purpose, procedure, benefits, and risks of the nuclear medicine procedure.
2. Assess for allergies to substances such as iodine.
3. Reassure the patient that the procedure is safe and painless.
4. Inform the patient that the procedure is performed in the nuclear medicine department. Contact the department to determine the expected time and length of the procedure.
5. Have the patient appropriately dressed.
6. Obtain an accurate weight because the radiopharmaceutical dose may be calculated by weight.
7. If a female patient is premenopausal, determine whether she may be pregnant. Pregnancy is a contraindication to most nuclear imaging.
8. Irradiation of the fetus should be avoided whenever possible.

**Clinical Alert**
1. Nuclear medicine procedures are usually contraindicated in pregnant women. Lactating women may need to discard their breast milk for several days following the procedure.
2. These precautions are also to be followed for the radionuclide laboratory procedures in Part 2 and PET imaging in Part 3 of this chapter.

**Posttest Patient Aftercare and Standard Precautions for Nuclear Medicine Scans**

1. Use routine disposal procedures for body fluids and excretions unless directed otherwise by the nuclear medicine department. Special considerations for disposal must be followed for therapeutic procedures.
2. Record any problems that may have occurred during the procedure.
3. Monitor the injection site for signs of bruising, hematoma, infection, discomfort, or irritation.
4. Assess for side effects of radiopharmaceuticals.

**Clinical Alert**
These precautions are also to be followed for the radionuclide laboratory procedures in Part 2 and PET imaging in Part 3 of this chapter.

**Pediatric Nuclear Medicine Considerations**

Many of the nuclear medicine procedures that are performed on adults may be indicated in children.

**Interventions**

**Pediatric Pretest Care**

1. Be aware that depending on hospital policy, a valid consent form may be requested to be signed by the parents or legal guardians of the patient.
2. Explain the procedure and its purpose, benefits, and risks to the parents or legal guardians and to the patient. Reassure the patient that the test is safe and painless.
3. Assess for allergy to medications.
4. Have the patient appropriately dressed, ensuring that there are no metal objects on the patient during the procedure.
5. Obtain an accurate weight; the dose is calculated based on the patient’s weight. Because pediatric patients have a different body metabolism than adults, a lower dose is given.
6. Remember that immobilization techniques are often used during the imaging of pediatric patients. Wrapping an infant or small child is often necessary. Head clamps, arm boards, or sandbags may be used for patient immobilization.
7. Administer sedative drugs to reduce patient motion during the examination. Disadvantages of sedation may include nausea and vomiting.
8. Start an intravenous line for administration of radiopharmaceuticals.
9. Do not leave patients unattended during the procedure.
10. Be aware that pediatric patients need constant reassurance and emotional support.
11. Be aware that patient urination is often difficult to control. A urinary catheter may be required.
12. Verify that the adolescent female patient is not pregnant.

**Pediatric Posttest Care**

1. Same as those stated for adults
2. Observe pediatric patients for adverse reactions to radiopharmaceuticals. Infants are more at risk for reactions.

**Part 1 Nuclear Medicine Scans**

**CARDIAC STUDIES**

**Myocardial Perfusion: Rest and Stress**

$^{99m}$Tc sestamibi, thallium-201 (Tl), and $^{99m}$Tc tetrofosmin are the radioactive imaging agents available for myocardial perfusion imaging to diagnose ischemic heart disease and allow differentiation of ischemia and infarction. This test reveals myocardial wall defects and heart pump performance during increased oxygen demands. These scans may also be done before and after streptokinase treatment for coronary artery thrombosis, after surgery for great vessel translocation, and after transplant to detect organ rejection and myocardial viability. Pediatric indications include evaluation for ventricular septal defects and congenital heart disease and postsurgical evaluation of congenital heart disease. $^{201}$Tl is a physiologic analogue of potassium. The myocardial cells extract potassium, as do other muscle cells. The $^{99m}$Tc sestamibi is taken up by the myocardium through passive diffusion, followed by active uptake within the mitochondria. Unlike thallium, technetium does not undergo significant redistribution. Therefore, there are some procedural differences. Myocardial activity also depends on blood flow. Consequently, when the patient is injected during peak exercise, the normal myocardium has much greater activity than the abnormal myocardium. Cold spots indicate a decrease or absence of flow. A completely normal myocardial perfusion study may eliminate the need for cardiac catheterization in the evaluation of chest pain and nonspecific abnormalities of the electrocardiogram (ECG). SPECT imaging can accurately localize regions of ischemia. Administration of dipyridamole (Persantine) or adenosine is indicated in adults and children who are unable to exercise to achieve the desired cardiac stress level and maximum cardiac vasodilation. This medication has an effect similar to that of exercise on the heart. Physical stress testing may be initiated in children beginning at 4 to 5 years. Candidates for drug-induced stress testing are those with lung disease, peripheral vascular disease with claudication, amputation, spinal cord injury, multiple sclerosis, or morbid obesity. Dipyridamole stress testing is also valuable as a significant predictor of cardiovascular death, reinfection, and risk for postoperative ischemic events and to reevaluate unstable angina. In some nuclear medicine departments, an ejection fraction and wall motion can be assessed by computer analysis.

**Reference Values**

**Normal**

Normal stress test: ECG and blood pressure normal Normal myocardial perfusion under both rest and stress conditions

**Procedures**

1. Myocardial perfusion general imaging
   a. Be aware that there are two phases to this procedure: the rest scan and the stress scan. Either $^{201}$Tl, $^{99m}$Tc sestamibi, or $^{99m}$Tc tetrofosmin may be used.
   1. Rest scan
      a. Perform an intravenous injection of the radioisotope. Allow a 30- to 60-minute delay for the radioisotope to localize in the heart.
      b. Perform SPECT imaging.
   2. Stress scan
      a. Be aware that the patient undergoes an exercise or a pharmacologic cardiac stress test. At the peak level of stress, inject the patient with the radioisotope.
      b. SPECT imaging may begin 30 minutes after injection.
   b. Pharmacologic stress tests may be performed with any of three routine stressing agents:
      1. Infuse dipyridamole over 4 to 6 minutes. Inject the radiopharmaceutical to localize in the heart.
      2. Infuse adenosine over 6 minutes. Inject the radiopharmaceutical 3 minutes into the infusion. Infuse the adenosine for 3 additional minutes. Be aware that adenosine has an extremely short half-life: once the infusion has stopped, any symptoms will subside. Contraindications: caffeine and theophylline-based drugs.
      3. Infuse dobutamine until the predicted heart rate is achieved. The infusion protocol lasts 3 minutes at each dose increment.

2. $^{201}$Tl
   a. During the cardiac stress test, the patient is monitored by a cardiologist, a registered nurse, an electrophysiologist, or an ECG technician.
   b. Have the patient begin walking on the treadmill.
   c. When the monitoring person determines that the patient has reached 85% to 95% of maximum heart rate, inject radioactive thallium. Take the patient for immediate imaging.
   d. Be aware that SPECT imaging begins within 5 minutes of injection.
   e. Acquire a second image approximately 3 to 4 hours later, with the patient at rest, to determine redistribution of the thallium.
   f. See Chapter 1 guidelines for safe, effective, informed intratest care.

3. $^{99m}$Tc sestamibi and $^{99m}$Tc tetrofosmin
   a. Follow myocardial perfusion general imaging procedures on page 657.
   b. Observe standard precautions.
**Procedural Alert**

Myocardial perfusion imaging protocols vary among nuclear medicine departments. Some departments use a rest-stress, stress-rest, dual-isotope, or 2-day protocol, separating the phases into 2 different days.

**Procedural Alert**

Some nuclear medicine protocols may require the patient to return 24 hours later for delayed imaging.

**Clinical Implications**

1. Imaging that is abnormal during exercise but remains normal at rest indicates transient ischemia.
2. A scan that is abnormal both at rest and under stress indicates a past infarction.
3. Hypertrophy produces an increase in uptake.
4. The progress of disease can be estimated.
5. The location and extent of myocardial disease can be assessed.
6. Specific and significant abnormalities in the stress ECG usually are indications for cardiac catheterization or further studies.

**Interfering Factors**

1. Inadequate cardiac stress
2. Caffeine intake
3. Injection of dipyridamole in the upright or standing position or with isometric handgrip may increase myocardial uptake.

**Interventions**

**Pretest Patient Care for Stress Testing**

1. Explain test purpose and procedure, benefits, and risks. See standard nuclear scan pretest precautions on page 655.
2. Before the stress test has begun, start an intravenous line and prepare the patient. Perform a resting 12-lead ECG and blood pressure measurement.
3. Advise the patient that the exercise stress period will be continued for 1 to 2 minutes after injection to allow the radiopharmaceutical to be cleared during a period of maximum blood flow.
4. Be aware that the patient should experience no discomfort during the imaging.
5. Alert the patient that fasting may be recommended for at least 2 hours before the stress test. Caffeine intake must be eliminated for 24 hours before the stress test.
6. For dipyridamole administration:
   a. Fasting may be required before the stress test and avoidance of any caffeine products for at least 24 hours before the test is necessary.
   b. Blood pressure, heart rate, and ECG results are monitored for any changes during dipyridamole infusion. Aminophylline may be given to reverse the effects of the dipyridamole.
7. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Clinical Alert**

1. The stress study is contraindicated in patients who:
   a. Have a combination of right and left bundle branch block
   b. Have left ventricular hypertrophy
   c. Are taking digitalis or quinidine
   d. Are hypokalemic (because the results are difficult to evaluate)
2. Adverse short-term effects of dipyridamole may include nausea, headache, dizziness, facial flush, angina, ST-segment depression, and ventricular arrhythmia.

**Posttest Patient Aftercare**

1. Observe the patient for possible effects of dipyridamole infusion.
2. Interpret test outcomes, counsel, and monitor appropriately.
3. Refer to nuclear scan posttest precautions on page 655.
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Myocardial Infarction (PYP) Scan**

\(^{99m}\text{Tc pyrophosphate} (\text{^{99m}\text{Tc-PYP}}) \) is the radioactive imaging agent used to demonstrate the general location, size, and extent of myocardial infarction 24 to 96 hours after suspected myocardial infarction and as an indication of myocardial necrosis, to differentiate between old and new infarcts. In some instances, the test is sensitive enough to detect an infarction 12 hours to 7 days after its occurrence. Acute infarction is associated with an area of increased radioactivity (hot spot) on the myocardial image. This test is useful when ECG and enzyme studies are not definitive.

**Reference Values**

**Normal** Normal distribution of the radiopharmaceutical in sternum, ribs, and other bone structures. No myocardial uptake

**Procedure**

1. Remember that myocardial imaging involves a 4-hour delay before imaging after the intravenous injection of the radionuclide. During this waiting period, the radioactive material accumulates in the damaged heart muscle.
2. Alert the patient that imaging takes 30 to 45 minutes, during which time the patient must lie still on an imaging table.
Clinical Implications

1. Imaging that is entirely normal indicates that an acute infarction is not present and the myocardium is viable.
2. Myocardial uptake of the PYP is compared with the ribs (2+) and sternum (4+). Higher uptake levels (4+) reflect greater myocardial damage.
3. Larger defects have a poorer prognosis than small defects.

Interfering Factors False-positive infarct-avid PYP can occur in cases of chest wall trauma, recent cardioversion, and unstable angina.

Interventions

Pretest Patient Care

1. Be aware that imaging can be performed at the bedside in the acute phase of infarction if the nuclear medicine department has a portable camera.
2. Explain the purpose, procedure, benefits, and risks of the nuclear medicine procedure. See standard pretest precautions on page 655.
3. Remember that imaging must occur within a period of 12 hours to 7 days after the onset of symptoms of infarction. Otherwise, false-negative results may be reported.
4. See Chapter 1 for additional guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret the outcome and monitor appropriately. If heart surgery is needed, counsel the patient concerning follow-up testing after surgery.
2. Refer to standard precautions and posttest care on page 655.
3. Follow additional guidelines in Chapter 1 for safe, effective, informed posttest care.

Multigated Acquisition (MUGA) Imaging: Rest and Stress

The term gated refers to the synchronization of the imaging equipment and computer with the patient's ECG to evaluate left ventricular function. The primary purpose of this test is to provide an ejection fraction (the amount of blood ejected from the ventricle during cardiac cycle). Once injected, the distribution of the radiolabeled red blood cells (RBCs) is imaged by synchronization of the recording of cardiac images with the ECG. This technique provides a means of obtaining information about cardiac output, end-diastolic volume, end-systolic volume, ejection fraction, ejection velocity, and regional wall motion of the ventricles. Computer-aided imaging of wall motion of the ventricles can be portrayed in the cinematic mode to visualize contraction and relaxation. This procedure may also be performed as a stress test. MUGA scans are not often performed on children.

Reference Values

Normal Normal myocardial wall motion and ejection fractions under conditions of stress and rest

Procedure

1. Remember that this procedure may be performed with or without stress. A MUGA with the patient at rest could be performed at the bedside if necessary, if the nuclear medicine department has a portable camera.
2. Label the patient's own RBCs with $^{99m}$Tc-PYP by any of several methods. Inject the blood once it is labeled. In children and adults, administer the $^{99m}$Tc-labeled RBCs slowly through an intravenous line. For children younger than 3 years of age, sedation may be required for the injection and to allow the pediatric patient to hold still for the required 20 to 30 minutes. Alternatively, perform a cardiac flow study.
3. Be aware that during an ECG, the patient's R wave signals the computer and camera to take several image frames for each cardiac cycle.
4. Image the patient immediately after injection of the labeled RBCs.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications Abnormal MUGA procedures as associated with:

1. Congestive cardiac failure
2. Change in ventricular function due to infarction
3. Persistent arrhythmias from poor ventricular function
4. Regurgitation due to valvular disease
5. Ventricular aneurysm formation

Interfering Factors If a reliable ECG cannot be obtained because of arrhythmias, the test cannot be performed.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks.
2. Follow standard nuclear scan pretest precautions on page 655.
3. See Chapter 1 for additional guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret MUGA outcomes and monitor appropriately for cardiac disease.
2. Refer to standard nuclear scan posttest precautions on page 655.
3. Follow basic Chapter 1 guidelines for safe, effective, informed posttest care.
Cardiac Flow Study (First-Pass Study; Shunt Imaging) The cardiac flow study is performed to check for blood flow through the great vessels and after vessel surgery; it is useful in the determination of both right and left ventricular ejection fractions. Immediately after the injection, the camera traces the flow of the radiopharmaceutical in its “first pass” through the cardiac chambers in multiple rapid images. The first-pass study uses a jugular or antecubital vein injection of the radiopharmaceutical. A large-bore needle is used. This study is useful in examining heart chamber disorders, especially left-to-right and right-to-left shunts. Children are commonly candidates for this procedure. Indications for pediatric patients include evaluation for congenital heart disease, transposition of great vessels, and atrial or ventricular septal defects and quantitative assessment of valvular regurgitation. In neonates, the cardiac flow study can be used in conjunction with computer software for the quantitative assessments. These quantitative values are useful in determining the degree of cardiac shunting with septal defects in the atria or ventricles.

Reference Values

**Normal**
- Normal wall motion and ejection fraction
- Normal pulmonary transit times and normal sequence of chamber filling

**Procedure**

1. Use a three-way stopcock with saline flush for radionuclide injection into the jugular vein or the antecubital fossa. For a shunt evaluation, inject the radionuclide into the external jugular vein to ensure a compact bolus. Be aware that with pediatric patients, it is important that the child not cry because this disrupts the flow of the radiopharmaceutical and negates the results of the test.
2. Have the patient lie supine with the head slightly raised.
3. Be aware that the total patient time is approximately 20–30 minutes; the actual imaging time is only 5 minutes.
4. Perform resting MUGA imaging with a shunt study.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. Abnormal first-pass ejection fraction values are associated with:
   - Congestive heart failure
   - Change in ventricular function due to infarction
   - Persistent arrhythmias from poor ventricular function
   - Regurgitation due to valvular disease
   - Ventricular aneurysm formation
2. Abnormal heart shunts reveal:
   - Left-to-right shunt
   - Right-to-left shunt
   - Mean pulmonary transit time
   - Tetralogy of Fallot (seen most often in children)

Interfering Factors
- Inability to obtain intravenous access to the jugular vein or large-bore antecubital access.

Interventions

**Pretest Patient Care**

1. Explain the purpose, procedure, benefits, and risks. An intravenous line is required.
2. See Chapter 1 for additional guidelines for safe, effective, informed pretest care.
3. Refer to standard nuclear scan precautions on page 655.
4. Obtain a signed, witnessed consent form if stress testing is to be done.

**Posttest Patient Aftercare**

1. Interpret test outcomes, monitor injection site, and counsel appropriately.
2. Refer to standard nuclear scan precautions on page 655.
3. Follow basic Chapter 1 guidelines for safe, effective, informed posttest care.

ENDOCRINE STUDIES

**Thyroid Imaging** The thyroid imaging test systematically measures the update of radioactive iodine (either $^{131}$I or $^{123}$I) by the thyroid. Iodine (and, consequently, radioiodine) is actively transported to the thyroid gland and is incorporated into the production of thyroid hormones. The test is required for the evaluation of thyroid size, position, and function. It is used in the differential diagnosis of masses in the neck, base of the tongue, or mediastinum. Thyroid tissue can be found in each of these three locations. Benign adenomas may appear as nodules of increased uptake of iodine (“hot” nodules), or they may appear as nodules of decreased uptake (“cold” nodules). Malignant areas generally take the form of cold nodules. The most important use of thyroid imaging is the functional assessment of these thyroid nodules. Pediatric indications include evaluation of neonatal hypothyroidism or thyrocarcinoma (lower incidence than adults). Thyroid imaging performed with iodine is usually acquired in conjunction with a radioactive iodine uptake study, which is usually performed 4 to 6 hours and 24 hours after dosing. For a complete thyroid workup, in both adults and children, thyroid hormone blood levels are usually measured. A thyroid ultrasound examination also may be performed.

**Reference Values**
- Normal or evenly distributed concentration of radioactive iodine
- Normal size, position, shape, site, weight, and function of the thyroid gland
- Absence of nodules

**Procedure**

1. Have the patient swallow radioactive iodine in a capsule or liquid form.
2. Determine an uptake 4 to 6 hours and 24 hours after dosing. Four hours after dosing, the thyroid (neck area) is imaged.
3. Alert patient that normal scan time is 45 minutes.
4. See Chapter 1 guidelines for safe, effective, informed intratest care.

**Clinical Implications**

1. Cancer of the thyroid most often manifests as a nonfunctioning cold nodule, indicated by a focal area of decreased uptake.
2. Some abnormal results are:
   a. Hyperthyroidism, represented by an area of diffuse increased uptake
   b. Hypothyroidism, represented by an area of diffuse decreased uptake
   c. Graves’ disease, represented by an area of diffuse increased uptake
   d. Autonomous nodules, represented by focal area of increased uptake
   e. Hashimoto’s disease, represented by mottled areas of decreased uptake
3. Imaging alone cannot definitively determine the diagnosis; uptake information is essential for a definitive diagnosis.

**Interfering Factors**

1. Thyroid imaging needs to be completed before radiographic examinations using contrast media (e.g., intravenous pyelogram, cardiac catheterization myelogram) are performed.
2. Any medication containing iodine should not be given until nuclear thyroid medicine procedures are concluded. Notify the attending physician if thyroid studies have been ordered or if there are interfering radiographs or medications.

**Interventions**

**Pretest Patient Care**

1. Instruct the patient about nuclear medicine imaging purpose, procedure, and special restrictions. Refer to standard nuclear scan pretest precautions on page 655.
2. Be aware that because the thyroid gland responds to small amounts of iodine, the patient may be requested to refrain from iodine intake for at least 1 week before the test. Patients should consult with a physician. Restricted items include the following:
   a. Certain thyroid drugs
   b. Weight-control medicines
   c. Multiple vitamins
   d. Some oral contraceptives
   e. X-ray contrast materials containing iodine
   f. Cough medicine
   g. Iodine-containing foods, especially kelp and other “natural” foods
3. Alleviate any fears the patient may have about radionuclide procedures.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Clinical Alert**

1. Nuclear medicine thyroid imaging is contraindicated in pregnancy. Thyroid testing in pregnancy is routinely limited to blood testing.
2. This study should be completed before thyroid-blocking radiographic contrast agents are administered and before thyroid or iodine drugs are given.
3. Occasionally, tests are performed purposely with iodine or some thyroid drug in the body. In these cases, the physician is testing the response of the thyroid to these drugs. These stimulation and suppression tests are usually done to determine the nature of a particular nodule and whether the tissue is functioning or nonfunctioning.

**Posttest Patient Aftercare**

1. If iodine has been administered, observe the patient for signs and symptoms of allergic reaction as needed.
2. Explain test outcomes and possible treatment.
3. Refer to standard nuclear scan posttest precautions on page 655.
4. Interpret test outcomes and counsel appropriately.
5. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Radioactive Iodine (RAI) Uptake Test** This direct test of the function of the thyroid gland measures the ability of the gland to concentrate and retain iodine. When radioactive iodine is administered, it is rapidly absorbed into the bloodstream. This procedure measures the rate of accumulation, incorporation, and release of iodine by the thyroid. The rate of absorption of the radioactive iodine, which is determined by the increase in radioactivity of the thyroid gland, is a measure of the ability of the thyroid to concentrate iodine from blood plasma. The radioactive isotopes of iodine used are $^{131}$I and $^{123}$I. This procedure is indicated in the evaluation of hypothyroidism, hyperthyroidism, thyroiditis, goiter, and pituitary failure and for posttreatment evaluation. The patient who is a candidate for this test may have a lumpy or swollen neck or complain of pain in the neck; the patient may be jittery and ultrasensitive to heat or sluggish and ultrasensitive to cold. The test is more useful in the diagnosis of hyperthyroidism than hypothyroidism.

**Reference Values**

*Normal* Absorption (uptake) by the thyroid gland: 1% to 13% after 2 hours 5% to 20% after 6 hours 15% to 40% after 24 hours (Values are laboratory dependent.)

**Procedure**

...
NOTE
The test usually is done in conjunction with thyroid imaging and assessment of thyroid hormone blood levels (see page 602).

1. Be aware that a fasting state is preferred. A complete history and listing of all medications is a must for this test. This history should include nonprescription medications and patient dietary habits.
2. Administer a liquid form or a tasteless capsule of radioactive iodine orally.
3. Measure the amount of radioactivity by an uptake calculation of the thyroid gland 4 to 6 and 24 hours later. There is no plan or discomfort involved.
4. Have the patient return to the laboratory at the designated time because the exact time of measurement is crucial in determining the uptake.

Clinical Implications

1. Increased uptake (eg, 20% in 1 hour, 25% in 6 hours, 45% in 24 hours) suggests hyperthyroidism but is not diagnostic for it.
2. Decreased uptake (eg, 0% in 2 hours, 3% in 6 hours, 10% in 24 hours) may be caused by hypothyroidism but is not diagnostic for it.
   a. If the administered iodine is not absorbed, as in severe diarrhea or intestinal malabsorption syndromes, the uptake may be low even though the gland is functioning normally.
   b. Rapid diuresis during the test period may deplete the supply of iodine, causing an apparently low percentage of iodine uptake.
   c. In renal failure, the uptake may be high even though the gland is functioning normally.

Clinical Alert

1. This test is contraindicated in pregnant or lactating women, in children, in infants, and in persons with iodine allergies.
2. Whenever possible, this test should be performed before any other radionuclide procedures are done, before any iodine medications are given, and before any radiographs using iodine contrast media are taken.

Interfering Factors

1. The chemicals, drugs, and foods that interfere with the test by lowering the uptake are:
   a. Iodized food and iodine-containing drugs such as Lugol solution, expectorants, cough medications, saturated solutions of potassium iodide, and vitamin preparations that contain minerals. The duration of the effects of these substances in the body is 1 to 3 weeks.
   b. Radiographic contrast media such as iodopyracet (Diodrast), sodium diatrizoate (Hypaque, Renografin), poppy-seed oil (Lipiodol), ethiodized oil (Ethiodol), iophendylate (Pantopaque), and iopanoic acid (Telepaque). The duration of the effects of these substances is 1 week to 1 year or more; consult with the nuclear medicine laboratory for specific times.
   c. Antithyroid drugs such as propylthiouracil (PTU) and related compounds (duration, 2 to 10 days)
   d. Thyroid medications such as lithium (duration, 1 to 2 weeks)
   e. Miscellaneous drugs such as thiocyanate, perchlorate, nitrates, sulfonamides, tolbutamide (Orinase), corticosteroids, parabenzamides (isoniazid, phenylbutazone (Butazolidin), thiopental (Pentothal), antihistamines, adrenocorticotropic hormone, aminosalicylic acid, cobalt, and coumarin anticoagulants. Consult with the nuclear medicine department for duration of effects of these drugs as they vary widely.
2. The compounds and conditions that interfere by enhancing the uptake are:
   a. Thyroid-stimulating hormone (thyrotropin)
   b. Pregnancy
   c. Cirrhosis
   d. Barbiturates
   e. Lithium carbonate
   f. Phenothiazines (duration, 1 week)
   g. Iodine-deficient diet
   h. Renal failure

Interventions

Pretest Patient Care

1. Explain test purpose and procedure; the test takes 24 hours to complete. Assess and record pertinent dietary and medication history.
2. Advise that iodine intake is restricted for at least 1 week before testing.
3. Refer to standard nuclear scan pretest precautions on page 655.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Explain test outcomes and possible treatment.
2. Refer to standard nuclear scan posttest precautions on page 655.
3. Interpret test outcomes and counsel appropriately.
4. Follow Chapter 1 guidelines for safe, effective, informed, posttest care.

Adrenal Gland (MIBG) Imaging The adrenal gland is divided into two different components: cortex and medulla. The scope of adrenal imaging is limited to the medulla. Testing can be performed in both adults and children. The purpose of adrenal medulla imaging is to identify sites of certain tumors that produce excessive amounts of catecholamines. Pheochromocytomas develop in cells that make up the adrenergic portion of the autonomic nervous system. A large number of these well-differentiated cells are found in adrenal medullas. Adrenergic tumors have been called paragangliomas when they are found outside the adrenal medulla, but many practitioners refer to all neoplasms that secrete norepinephrine and epinephrine as pheochromocytomas. Because the only definite and effective therapy is surgery to remove the tumor, identification of the site using this test, computed tomography, and ultrasound, is an essential goal of treatment.

**Reference Values**

**Normal** No evidence of tumors or hypersecreting hormone sites Normal salivary glands, urinary bladder, and vague shape of liver and spleen can be seen.

**Procedure**

1. Inject intravenously the radionuclide $^{131}$I metaiodobenzylguanidine (MIBG).
2. Take sequential images at the physician's discretion, usually beginning 24 hours after injection.
3. Alert patient that imaging may take 2 hours.
4. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. Abnormal results give substance to the "rough rule of 10" for these tumors:
   a. Ten percent are in children.
   b. Ten percent are familial.
   c. Ten percent are bilateral in the adrenal glands.
   d. Ten percent are malignant.
   e. Ten percent are multiple, in addition to bilateral.
   f. Ten percent are extraluminal.
2. More than 90% of primary pheochromocytomas occur in the abdomen.
3. Pheochromocytomas in children often represent a familial disorder.
4. Bilateral adrenal tumors often indicate a familial disease, and vice versa.
5. Multiple extraluminal pheochromocytomas are often malignant.
6. The presence of two or more pheochromocytomas strongly indicates malignant disease.

**Interfering Factors** Barium interferes with the test.

**Interventions**

**Pretest Patient Care**

1. Explain nuclear medical imaging purpose, procedure, benefits, and risks.
2. Give Lugol solution (potassium iodine) for 1 week before the injection to prevent uptake of radioactive iodine by the thyroid gland.
3. Refer to standard nuclear scan pretest precautions on page 655.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome and counsel appropriately about the need for possible follow-up tests. Follow-up tests include:
   a. Kidney and bone imaging to give further orientation to abnormalities discovered by MIBG scan.
   b. Computed tomography procedure if MIBG imaging failed to locate the tumor.
   c. Ultrasound of the pelvis if the tumor produces urinary symptoms.
2. Refer to standard nuclear scan posttest precautions on page 655.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Parathyroid Imaging Parathyroid imaging is done to localize parathyroid adenomas in clinically proven cases of primary hyperparathyroidism. It is helpful in demonstrating intrinsic or extrinsic parathyroid adenoma. Two tracers, $^{99m}$Tc sestamibi and $^{123}$I capsules, are administered. In children, this scan is done to verify presence of the parathyroid gland after thyroidectomy.

**Reference Values**

**Normal** No areas of increased perfusion or uptake in parathyroid or thyroid

**Procedure**

1. Administer $^{123}$I. Four hours later, image the neck.
2. Inject $^{99m}$Tc sestamibi without moving the patient; after 10 minutes, acquire additional images. Computer processing involves subtracting the technetium-visualized thyroid structures from the $^{123}$I accumulation in a parathyroid adenoma.
3. Alert patient that total examination time is 1 hour.
4. See Chapter 1 guidelines for safe, effective, informed intratest care.

**Clinical Implications** Abnormal concentrations of the radiopharmaceuticals reveal parathyroid adenoma, both intrinsic and extrinsic, but cannot differentiate between benign and malignant adenomas.

**Interfering Factors** Recent ingestion of iodine in food or medication and recent tests with iodine contrast are
Clinical Considerations Pregnancy is a relative contraindication. However, if primary hyperparathyroidism is suspected and surgical exploration is essential before delivery, the study may be performed.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of parathyroid imaging.
2. Assess for the recent intake of iodine. However, this finding is not a specific contraindication to performing the study.
3. Palpate the thyroid carefully.
4. Refer to standard nuclear scan pretest precautions on page 655.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Refer to standard nuclear scan posttest precautions on page 655.
2. Interpret test outcome and monitor appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

GENITOURINARY STUDIES

Renogram: Kidney Function and Renal Blood Flow Imaging (With Furosemide or Captopril)

The renogram is performed in both adult and pediatric patients to study the function of the kidneys and to detect renal parenchymal or vascular disease or defects in excretion. The radiopharmaceutical of choice, $^{99m}$Tc mertiatide (MAG-3), permits visualization of renal clearance. In pediatric patients, this procedure is done to evaluate hydronephrosis, obstruction, reduced renal function (premature neonates), renal trauma, and urinary tract infections. The renogram is ideal for pediatric evaluation because of the nontoxic nature of the radiopharmaceuticals, compared with the contrast media used in radiology procedures. Post–kidney transplant scans, which assess perfusion and excretory function as a reflection of GFR, are done when the serum creatinine level increases and determine kidney damage leading to acute tubular necrosis (ATN).

Reference Values

Normal Equal blood flow in right and left kidneys in 10 minutes, 50% of the radiopharmaceutical should be excreted.

Indications

1. To detect the presence or absence of unilateral kidney disease
2. For long-term follow-up of hydroureteronephrosis
3. To study the hypertensive patient to evaluate for renal artery stenosis. The captopril test is a first-line study to determine a renal basis for hypertension.
4. To study the azotemic patient when urethral catheterization is contraindicated or impossible
5. To evaluate upper urinary tract obstruction
6. To assess renal transplant efficacy

Procedure

1. Place the patient in either an upright sitting or supine position for imaging; the supine position is preferred for pediatric patients.
2. Inject the radiopharmaceutical intravenously. An intravenous diuretic (furosemide [Lasix]) or angiotensin-converting enzyme (ACE) inhibitor (captopril) may also be administered during a second phase of the renogram.
3. Start imaging immediately after injection.
4. Alert patient that total examination time is approximately 45 minutes for a routine, one-phase renogram.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Alert

1. A renogram may be performed in a pregnant woman if it is imperative that renal function be ascertained.
2. Some renal transplant recipients may have more than two kidneys, eg, the transplant, their own two kidneys, and an older failing transplant. Sometimes, two pediatric kidneys will both be transplanted.

Clinical Implications Abnormal distribution patterns may indicate:

1. Hypertension
2. Obstruction due to stones or tumors
3. Renal failure
4. Decreased renal function
5. Diminished blood supply
6. Renal transplant rejection
7. In pediatric patients, urinary tract infections in male neonates; the finding shifts to females after 3 months of age.

Interfering Factors Diuretics, ACE inhibitors, and beta blockers are medications that may interfere with the test results.

Interventions

Pretest Patient Care
1. Explain the purpose, procedure, benefits, and risks of the procedure. Pediatric patients have a detectible glomerular filtration rate after 6 months of age. In the neonate, ultrasound is used in combination with nuclear medicine procedures for a more complete renal assessment. Refer to standard nuclear scan pretest precautions on page 655. An intravenous line is placed before imaging. Check for history of previous transplant.

2. Unless contraindicated, ensure that the patient is well hydrated with two to three glasses of water (10 mL per kilogram of body weight) before undergoing the test.

3. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Encourage fluids and frequent bladder emptying to promote excretion of radioactivity.
2. Interpret test outcome and counsel appropriately.
3. Refer to standard nuclear scan posttest precautions on page 655.
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

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**Testicular (Scrotal) Imaging**

This test is performed on an emergency basis to evaluate acute, painful testicular swelling. It also is used in the differential diagnosis of torsion or acute epididymitis and in evaluation of injury, trauma, tumors, and masses. The radiopharmaceutical $^{99m}$Tc pertechnetate is injected intravenously. The images obtained differentiate lesions associated with increased perfusion from those that are primarily ischemic. In pediatric patients, the procedure is done to diagnose acute or latent testicular torsion, epididymitis, or testicular hydrocele and for evaluation of testicular masses such as abscesses and tumors.

**Reference Values**

**Normal** Normal blood flow to scrotal structures, with even distribution and concentration of the radiopharmaceutical

**Procedures**

1. Have the patient lie supine under the gamma camera. Tape the penis gently to the lower abdominal wall. For proper positioning, use towels to support the scrotum. Place lead shielding in the perineal area to reduce any background activity.
2. Inject the radionuclide intravenously. In pediatric patients, do not inject the radiopharmaceutical through veins in the legs because this interferes with the study.
3. Perform imaging in two phases: first, as a dynamic blood flow study of the scrotum; and second, as an assessment of distribution of the radiopharmaceutical in the scrotum.
4. Alert patient that total examining time is 30–45 minutes.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

**Clinical Implications**

1. Abnormal concentrations reveal:
   a. Tumors
   b. Hematomas
   c. Infection
   d. Torsion (with reduced blood flow). In the neonatal patient, torsion is caused primarily by developmental anomalies.
   e. Acute epididymitis
2. The nuclear scan is most specific soon after the onset of pain, before abscess is a clinical consideration.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose, procedure, benefits, and risks of the test. There is no discomfort involved in testing.
2. If the patient is a child, a parent should accompany the boy to the department.
3. Tape the penis to the lower abdominal wall.
4. Refer to standard nuclear scan pretest precautions, page 655.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Refer to standard nuclear scan posttest precautions, page 655.
2. Interpret test outcome and monitor appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Vesicoureteric Reflux (Bladder and Ureters) Imaging
Vesicoureteric reflux imaging usually is done on pediatric patients to assess abnormal bladder filling and possible reflux into the ureters. $^{99m}$Tc pentetate (DTPA) is administered through a urinary catheter, followed by sufficient saline until the patient has an urge to urinate. The ureters and kidneys are scanned by the camera during administration to detect the reflux.

Reference Values

**Normal** Normal bladder filling without any reflux into the ureters

Procedure

1. Place the patient in the supine position. Use a special urinary catheter kit, and insert a urinary catheter.
2. Start the camera immediately for dynamic acquisition while the radiopharmaceutical and saline are administered until the bladder is full or there is patient discomfort.
3. Remove the catheter once the imaging is complete.

Clinical Implications Abnormal vesicoureteric reflux may be either congenital (immature development of the urinary tract) or caused by infection.

Interventions

**Pretest Patient Care**

1. See standard pretest care for nuclear scan of pediatric patients (see page 656).
2. Place a urinary catheter with sterile saline. Place an absorbent, plastic-backed pad under the patient to absorb any leakage of radioactive material. If a urinary catheter is contraindicated for the patient, use an alternative indirect renogram method.

**Posttest Patient Aftercare**

1. Refer to standard nuclear scan posttest precautions (see page 655), the same as for adults.
2. Be aware that depending on cause and severity, antibiotic therapy or surgery is used to treat the condition.
3. Remember that special handling of the patient's urine (gloves and handwashing before and after gloves are removed) is necessary for 24 hours after completion of the test.

GASTROINTESTINAL STUDIES

Hepatobiliary (Gallbladder, Biliary) Imaging With Cholecystokinin

This study, using $^{99m}$Tc disofenin or mebrofenin, is performed to visualize the gallbladder and determine patency of the biliary system. In pediatric patients, this test is done to differentiate biliary atresia from neonatal hepatitis and to assess liver trauma, right upper quadrant pain, and congenital malformations.

A series of images traces the excretion of the radionuclide. Through computer analysis, the activity in the gallbladder is quantitated, and the amount ejected (ejection fraction) is calculated.

Indications for Testing

1. To evaluate cholecystitis
2. To differentiate between obstructive and nonobstructive jaundice
3. To investigate upper abdominal pain
4. Biliary assessment after surgery
5. Evaluation of biliary atresia

Reference Values

**Normal** Rapid transit of the radionuclide through the liver cells to the biliary tract (15 to 30 minutes) with significant uptake in the normal gallbladder Normal distribution patterns in the biliary system, from the liver, through the gallbladder, to the small intestines

Procedure

1. Inject the radionuclide intravenously. In adults and older children, give cholecystokinin (CCK) to stimulate gallbladder contraction. In infants, give phenobarbital to distinguish between biliary atresia and neonatal jaundice.
2. Start imaging immediately after injection. Take a series of images at 5-minute intervals for as long as it takes to visualize the gallbladder and small intestine.
3. In the event of biliary obstruction, obtain delayed views.
4. Remember that if CCK is administered, computer-assisted quantitative measurements can determine an ejection fraction.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. Abnormal concentration patterns reveal unusual bile communications.
2. Gallbladder visualization excludes the diagnosis of acute cholecystitis with a high degree of certainty.

Interfering Factors
1. Patients with high serum bilirubin levels (>10 mg/dL or >171 µmol/L) have less reliable test results.
2. Patients receiving total parenteral nutrition or with long-term fasting may not have gallbladder visualization.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of the procedure.
2. Ensure that the patient is NPO for at least 4 hours (3–4 hours for pediatric patients) before testing. In case of prolonged fasting (>24 hours), notify the nuclear medicine department. Fasting does not apply when the indication is for biliary atresia or jaundice.
3. Discontinue opiate- or morphine-based pain medications 2 to 6 hours before the test to avoid interference with transit of the radiopharmaceutical.
4. Refer to standard nuclear scan pretest precautions on page 655.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately.
2. Refer to standard nuclear scan posttest precautions on page 655.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Gastroesophageal Reflux Imaging

This test is indicated for both adult and pediatric patients to evaluate esophageal disorders such as regurgitation and to identify the cause of persistent nausea and vomiting. In infants, the study is used to distinguish between vomiting and reflux (for those with more severe symptoms). A certain amount of reflux occurs naturally in infants. If timely diagnosis and treatment of gastrointestinal reflux does not occur, additional complications may result, such as recurrent respiratory infections, apnea, or sudden infant death syndrome (SIDS).

After oral administration of the radioisotope $^{99m}$Tc sulfur colloid in orange juice or scrambled eggs, the patient is immediately imaged to verify that the dose is in the stomach. Images are acquired in 2 hours. A computer analysis is used to calculate the percentage of reflux into the esophagus for each image.

Reference Values

Normal Less than 40% gastric reflux across the esophageal sphincter

Procedure

1. Have the patient ingest the radionuclide in orange juice or in scrambled eggs. For infants, perform the test at the normal infant feeding time to determine esophageal transit. Have the infant drink $^{99m}$Tc-labeled sulfur colloid mixed with milk. Give a portion of the milk containing the radioisotope, and burp the infant before the remainder is given. Give some unlabeled milk to clear the esophagus of the radioactive material. If a nasogastric tube is required for radiopharmaceutical administration, remove it before the imaging occurs to avoid a false-positive result.
2. Be aware that images are obtained in 2 hours.
3. Remember that a computer analysis generates a time-activity curve to calculate the reflux.
4. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications More than 4% reflux is abnormal. The percentage of reflux is used to evaluate patients before and after surgery for gastroesophageal reflux.

Clinical Alert

Patients who have esophageal motor disorders, hiatal hernias, or swallowing difficulties should have an endogastric tube inserted for the procedure.

Interfering Factors Previous upper gastrointestinal radiographic procedures may interfere with this test.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks. See standard nuclear scan pretest precautions on page 655.
2. Perform imaging with the patient in a supine position.
3. Ensure that the patient is fasting from midnight of the previous night until the examination.
4. Monitor oral intake of the orange juice or scrambled eggs containing $^{99m}$Tc sulfur colloid.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Remove endogastric tubes, if placed for the examination, after the radiopharmaceutical is administered.
2. Refer to standard nuclear scan posttest precautions on page 655.
3. Interpret test outcome and monitor appropriately.
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Gastric Emptying Imaging

Gastric emptying imaging is used in both adult and pediatric patients to assess gastric motility disorders and in patients
with unexplained nausea, vomiting, diarrhea, and abdominal cramping. The emptying of food by the stomach is a complex process that is controlled by food composition (fats, carbohydrates), food form (liquid, solid), hormone secretion (gastrin, CCK), and nervous innervation. Because clearance of liquids and clearance of solids vary, the imaging procedure traces both food forms. Indications for imaging include both mechanical and nonmechanical gastric motility disorders. Mechanical disorders include peptic ulcerations, gastric surgery, trauma, and cancer. Nonmechanical disorders include diabetes, uremia, anorexia nervosa, certain drugs (opiates), and neurologic disorders. Clearance of liquids, solids, or a combination (dual-phase examination) may be studied.

Reference Values
Normal
Normal half-time clearance ranges: 45–110 minutes for solids 10–65 minutes for liquids

Procedure

1. Have the fasting patient consume the solid phase (99mTc sulfur colloid, usually in scrambled eggs or oatmeal or chicken livers) followed by the liquid phase (indium-111 [111In]-DTPA in 300 mL water). For infants, perform the test at the normal feeding time. Have the infant drink 99mTc sulfur colloid mixed with milk. Provide older children solids such as scrambled eggs mixed with 99mTc sulfur colloid.
2. Perform imaging immediately, with the patient in the supine position.
3. Obtain subsequent images over the next 2 hours.
4. Use computer processing to determine the half-time clearance for both liquid and solid phases of gastric emptying.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. Slow or delayed emptying is usually seen in the following conditions:
   a. Peptic ulceration
   b. Diabetes
   c. Smooth muscle disorders
   d. After radiation therapy
   e. In pediatric patients, hypomotility of the antrum portion of the stomach is the primary cause of delayed gastric emptying. However, all abnormal functions of the stomach do contribute to the delay.
2. Accelerated emptying is often seen in the following conditions:
   a. Zollinger-Ellison syndrome
   b. Certain malabsorption syndromes
   c. After gastric or duodenal surgery

Interfering Factors
Administration of certain medications (eg, gastrin, CCK) interferes with gastric emptying.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of the procedure.
2. Have the adult patient fast for 8 hours before the test.
3. Refer to standard nuclear scan pretest precautions on page 655.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Be aware that the patient may eat and drink normally.
2. Interpret test outcomes and counsel appropriately.
3. Refer to standard nuclear scan posttest precautions on page 655.
4. Follow Chapter 1 basic guidelines for safe, effective, informed posttest care.

Gastrointestinal Bleeding Imaging

This test is very sensitive in the detection and location of acute gastrointestinal bleeding that occurs distal to the ligament of Treitz. (Gastroscopy is the procedure of choice for diagnosis of upper gastrointestinal bleeding.) Before this diagnostic technique was refined, barium enemas were used to identify lesions reflecting sites of bleeding, but that test was not specific and frequently missed small sites of bleeding. This procedure is also indicated for detection and localization of recent hemorrhage, both peritoneal and retroperitoneal. The radiopharmaceutical of choice for suspected active bleeding is 99mTc-labeled RBCs.

Reference Values
Normal
No sites of active bleeding

Procedure

1. Inject 99mTc-labeled RBCs intravenously.
2. Begin imaging immediately after injection and continue every few minutes. Obtain images anteriorly over the abdomen at 5-minute intervals for 60 minutes or until a bleeding site is located. If the study is negative after 1 hour, obtain delayed images 2, 6, and sometimes 24 hours later, when necessary, to identify the location of difficult-to-determine bleeding sites.
3. Be aware that total examining times varies.
4. See Chapter 1 guidelines for safe, effective, informed intratest care.
Clinical Alert

1. This test is contraindicated in patients who are hemodynamically unstable. In these instances, angiography or surgery should be the procedure of choice.
2. Assess the patient for signs of active bleeding during the examining period.
3. Recent blood transfusion may be a contraindication for this study.

Clinical Implications: Abnormal concentrations of RBCs (hot spots) are associated with active gastrointestinal bleeding sites, both peritoneal and retroperitoneal.

Interfering Factors: Presence of barium in gastrointestinal tract may obscure the site of bleeding because of the high density of barium and the inability of the technetium to penetrate the barium.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of the gastrointestinal blood loss imaging.
2. Determine whether the patient has received barium as a diagnostic agent within the past 24 hours. If the presence of barium in the gastrointestinal tract is questionable, an abdominal radiograph may be ordered.
3. Advise the patient that delayed images may be necessary. Also, if active bleeding is not seen on initial imaging, additional images must be obtained for up to 24 hours after injection in a patient with clinical signs of active bleeding.
4. Refer to standard nuclear scan pretest precautions on page 655.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Refer to standard nuclear scan posttest precautions on page 655.
2. Interpret test outcome and monitor appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Parotid (Salivary) Gland Imaging

This study is helpful in the evaluation of swelling masses in the parotid region. This imaging is done to detect blocked tumors of parotid or salivary glands and to diagnose Sjögren's syndrome. The radionuclide injected intravenously is $^{99m}$Tc pertechnetate. One of the limitations of the test is that it cannot furnish an exact preoperative diagnosis.

Reference Values

Normal: No evidence of tumor-type activity or blockage of ducts. Normal size, shape, and position of the glands.

Procedure

1. Inject the radionuclide pertechnetate intravenously. Perform imaging immediately. There are three phases to imaging: blood flow, uptake or trapping mechanism, and secreting capability.
2. Take images of the gland every minute for 30 minutes.
3. If a secretory function test is being performed to detect blockage of the salivary duct, three fourths of the way through the test, ask the patient to suck on a lemon slice. If the salivary duct is normal, this causes the gland to empty. This is not done in studies undertaken for tumor detection.
4. Alert patient that total test time is 45 to 60 minutes.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. The reporting of a hot nodule amidst normal tissue that accumulates the radionuclide is associated with tumors of the ducts, as in:
   a. Warthin's tumor
   b. Oncocytoma
   c. Mucocoeiform tumor
2. The reporting of a cold nodule amidst normal tissue that does not accumulate the radionuclide is associated with:
   a. Benign tumors, abscesses, or cysts, which are indicated by smooth, sharply defined outlines
   b. Adenocarcinomas, which are indicated by ragged, irregular outlines
3. Diffuse decreased activity occurs in obstruction, chronic sialadenitis, or Sjögren's syndrome.
4. Diffuse increased activity occurs in acute parotitis.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks.
2. No pain or discomfort is involved.
3. Lemon may be given to the patient to stimulate parotid secretion.
4. Refer to standard nuclear scan pretest precautions on page 655.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and monitor appropriately.
2. Refer to standard nuclear scan posttest precautions on page 655.
Liver/Spleen Imaging and Liver RBC Imaging

This test is used to demonstrate the anatomy and size of the liver and spleen. It is helpful in determining the cause of right upper quadrant pain and in the detection of metastatic disease, cirrhosis, ascites, infarction due to trauma, and liver damage due to radiation therapy. Most liver and spleen imaging evaluates for metastatic disease and for the differential diagnosis of jaundice. Post–liver transplant scans detect bile and anastomoses leaks and rule out abnormal perfusion as a sign of rejection.

The radioactive material, $^{99m}$Tc-labeled sulfur colloid, is injected intravenously. Liver/ spleen SPECT imaging provides 3-dimensional images of radiopharmaceutical uptake. The radiopharmaceutical most specific for detection of hemangioma in the liver is $^{99m}$Tc labeled to a patient’s own RBCs. In many instances, ultrasound imaging replaces this test.

Reference Values

| Normal | Normal liver size, shape, and position within the abdomen | Normal spleen size, cell function, and blood flow in the spleen | Normally functioning liver and spleen reticuloendothelial system |

NOTE

The amount of uptake in the spleen should always be less than in the liver.

Procedure

1. Inject the radiopharmaceutical intravenously.
2. Perform a SPECT study and planar images.
3. Be aware that the entire study usually takes 60 minutes from injection to finish.
4. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. Abnormal liver and spleen scan patterns occur in:
   a. Cirrhosis
   b. Hepatitis
   c. Trauma
   d. Hepatomas
   e. Sarcoïdosis
   f. Metastasis
   g. Cysts
   h. Perihepatic abscesses
   i. Hemangiomas
   j. Adenomas
   k. Ascites
2. Abnormal splenic concentrations reveal:
   a. Unusual splenic size
   b. Infarct
   c. Ruptured spleen
   d. Accessory spleen
   e. Tumors
   f. Metastatic spread
   g. Leukemia
   h. Hodgkin’s disease
3. Spleens more than 14 cm are abnormally enlarged; those less than 7 cm are abnormally small. Areas of absent radioactivity or holes in the spleen scan are associated with abnormalities that displace or destroy normal splenic pulp.
4. About 30% of persons with Hodgkin’s disease with splenic involvement have a normal splenic image.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks.
2. Be aware that this test can be performed in cases of trauma or suspected ruptured spleen, at bedside or in the emergency room.
3. Refer to standard nuclear scan pretest precautions on page 655.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Refer to standard nuclear scan posttest precautions on page 655.
2. Interpret test outcome and monitor appropriately. Explain need for medical treatment or surgery.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Meckel’s Diverticulum Imaging

3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.
The test for Meckel's diverticulum usually is done in pediatric patients diagnosed with congenital abnormality of the ileum, which sometimes continues to the umbilicus with fistula formation. The uptake of $^{99m}$Tc pertechnetate occurs in the parietal cells of the gastric mucosa and is detected by the gamma camera. Meckel's diverticulum shows uptake in the distal portion of the ileum. This anomaly contains secretory cells similar to those of gastric mucosa. An alternative radiopharmaceutical, $^{99m}$Tc-labeled RBCs, may be considered in cases of suspected bleeding sites associated with the diverticulum.

Reference Values

**Normal** Normal blood pool distribution and clearance of the radioactive tracer into the duodenum and jejunum

**Procedure**
1. Have the patient lie supine and inject with the radiopharmaceutical.
2. Start the camera immediately with a series of static images obtained at 5-minute intervals of 30 minutes.
3. Be aware that extra spot views may be requested by the physician.
4. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications
1. Abnormal results reveal rectal bleeding, the most common symptom of Meckel's diverticulum. Meckel's diverticulum can occur with or without abdominal symptoms.
2. If it is left undetected and untreated, ulceration of the ilium may occur, and strangulation may cause intestinal obstruction.

Interventions

**Pretest Patient Care**
1. See standard pretest care for nuclear scan of pediatric patients (see page 656). Explain the purpose and procedures of the examination. Patients should be fasting. Other diagnostic procedures involving the gastrointestinal tract and medications affecting the intestines should be avoided for 2 to 3 days before the examination. This is especially true of lower and upper gastrointestinal radiographic procedures.
2. Have patients void immediately before the examination.

**Posttest Patient Aftercare**
1. Refer to standard posttest precautions (see page 655), the same as for adults. Special handling of the patient's urine (gloves and handwashing before and after glove removal) is necessary for 24 hours after test completion.
2. Interpret test outcome and monitor appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**NEUROLOGIC STUDIES**

**Brain Scan and Cerebral Blood Flow Imaging**

Brain imaging provides information about regional perfusion and brain function, whereas CT and MRI show structural changes. Recent developments in radiopharmaceuticals and SPECT have rejuvenated brain imaging. Newer technetium complexes, such as $^{99m}$Tc bicisate (ECD) and $^{99m}$Tc exametazime, are radiopharmaceuticals that cross the blood-brain barrier. The blood-brain barrier is not an anatomic structure but a complex system of select mechanisms that oppose the passage of most ions and large molecular-weight compounds from the blood to the brain tissue, that includes capillary endothelium with closed intracellular clefts, a small or absent extracellular fluid space between endothelium and glial sheaths, and the membrane of the neurons themselves. SPECT technology allows for 3-dimensional slices, providing depth resolution from different angles. Although PET imaging is more effective in functional diagnosis, SPECT is less expensive and more readily available. This test is indicated in both adults and children to determine brain death or the presence of encephalitis; it is also used in children with hydrocephalus, to localize epileptic foci, to assess metabolic activity, to evaluate brain tumors, and for the assessment of childhood development disorders.

Reference Values

**Normal** Normal extracranial and intracranial blood flow Normal distribution, with highest uptake in the gray matter, basal ganglia, thalamus, and peripheral cortex and less activity in the central white matter and ventricles

**Procedure**
1. Inject the radionuclide intravenously. During the injection, have the patient in a relaxed, controlled environment to minimize anxiety. In uncooperative children, do not use sedation until after the injection because it may affect brain activity. Secure the patient's head during the examination.
2. Begin imaging immediately after administration of the radiopharmaceutical or after a 1-hour delay. It takes about 1 hour to complete.
3. With the patient in the supine position, obtain SPECT images around the circumference of the head.
4. Be aware that with administration of iodoamphetamine, some departments require a dark and quiet environment.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications
1. Abnormal radionuclide distribution patterns indicate:
   a. Alzheimer's disease
b. Stroke  
c. Dementia  
d. Seizure disorders  
e. Epilepsy  
f. Systemic lupus erythematosus  
g. Huntington's disease  
h. Parkinson's disease  
i. Psychiatric diagnosis (schizophrenia)

2. The cerebral blood flow in a patient with brain death shows a very distinct image: there is a lack of tracer uptake in the anterior and middle cerebral arteries and in the cerebral hemisphere, but perfusion is present in the scalp veins.

Interfering Factors

1. Any patient motion (eg, coughing, leg movement) can alter cerebral alignment.  
2. Sudden distractions or loud noises can alter the distribution of the radionuclide.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risk.  
2. Refer to standard nuclear scan pretest precautions in page 655.  
3. Because precise head alignment is crucial, advise the patient to remain quiet and still.  
4. Obtain a careful neurologic history before testing.  
5. See Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Refer to standard nuclear scan posttest precautions on page 655.  
2. Interpret test outcome and monitor appropriately, especially if sedation is used.  
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Cisternography (Cerebrospinal Fluid Flow Imaging)

This study, in which the radiopharmaceutical $^{111}$In DTPA is injected intrathecally during a lumbar puncture, is a sensitive indicator of altered flow and reabsorption of CSF. Congenital malformations are the most common causes of hydrocephalus in the neonate. In older patients and in cases of trauma, CT or MRI is often used to identify anatomic origins of obstructive hydrocephalus. In the treatment of hydrocephalus, this test aids in selection of the type of shunt and pathway and in determining the prognosis of both shunting and hydrocephalus.

Reference Values

Normal
Unobstructed flow of cerebrospinal fluid and normal reabsorption

Procedure

1. Perform a sterile lumbar puncture after the patient has been positioned and prepared (see Chapter 5 for lumbar puncture procedure). At this time, inject the radionuclide into the cerebrospinal circulation.  
2. Have the patient lie flat after the puncture; the length of time depends on the physician's order.  
3. Perform imaging 2 to 6 hours after injection and repeat after 24 hours, 48 hours, and 72 hours if the physician so directs.  
4. Be aware that examining time is 1 hour for each imaging.  
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications
Abnormal filling patterns reveal:

1. Cause of hydrocephalus (eg, trauma, inflammation, bleeding, intracranial tumor)  
2. Subdural hematoma  
3. Spinal mass lesions  
4. Posterior fossa cysts  
5. Parenchymal and subarachnoid cysts  
6. Communicating versus noncommunicating hydrocephalus  
7. Shunt patency  
8. Diagnosis and localization of rhinorrhea and otorrhea

Interventions

Pretest Patient Care

1. Explain the purposes, procedures, benefits, and risks of both lumbar puncture and cisternography.  
2. Refer to standard nuclear scan pretest precautions on page 655.  
3. Advise the patient that it may take as long as 1 hour for each imaging session.  
4. Because of the lumbar puncture, take the patient by cart to the nuclear medicine department for the first imaging session.  
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Follow instructions for lumbar puncture (see Chapter 5) and standard nuclear scan posttest precautions on page 655.
2. Be alert to complications of lumbar puncture, such as meningitis, allergic reaction to anesthetic, bleeding into spinal canal, herniation of brain tissue, and mild to severe headache.
3. Interpret test outcome and monitor appropriately.
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

PULMONARY STUDIES

Lung Scan (Ventilation and Perfusion Imaging)

The lung imaging is performed for three major purposes:

1. To diagnose and locate pulmonary emboli
2. To detect the percentage of the lungs that is functioning normally
3. To assess the pulmonary vascular supply by providing an estimate of regional pulmonary blood flow

Lung imaging in both adults and children is done to assess pneumonia, cystic fibrosis, cyanosis, asthma, airway obstruction, infection, inflammation, and acquired immunodeficiency syndrome (AIDS)-related pulmonary diseases. It is a simple method for monitoring the course of embolic disease because an area of ischemia persists after apparent resolution on chest radiographs. In the case of pulmonary embolus, the blood supply beyond an embolus is restricted. Imaging results in poor or no visualization of the affected area. Assessment of the adequacy of pulmonary artery perfusion in areas of known disease can also be done reliably, as well as after lung transplant to detect reperfusion of lung and bronchiolitis obliterans.

There are two parts to the lung imaging: the ventilation ($\bar{V}$) imaging and the perfusion ($\bar{Q}$) imaging. The ventilation imaging reveals the movement or lack of air in the lungs. An aerosol of $^{99m}$Tc DTPA or xenon-133 ($^{133}$Xe) gas demonstrates the ventilation properties of the patient's lungs. The perfusion imaging demonstrates the blood supply to the tissues in the lungs.

When inhaled, the radioactive gas or aerosol follows the same pathway as the air in normal breathing. In some pathologic conditions affecting ventilation, there is significant alteration in the normal ventilation process. The $\bar{V}/\bar{Q}$ is significant in the diagnosis of pulmonary emboli. It is also helpful in diagnosing bronchitis, asthma, inflammatory fibrosis, pneumonia, chronic obstructive pulmonary disease, and lung cancer.

The lung perfusion study can be performed after the ventilation test. A macroaggregated albumin (MAA) labeled with technetium is injected intravenously, and assessment of the pulmonary vascular supply is achieved by imaging.

Certain limitations exist with these tests. With a positive chest film and a positive $\bar{V}/\bar{Q}$, the differential possibilities are multiple: pneumonia, abscess, bullae, atelectasis, and carcinoma, among others. A pulmonary arteriogram is still necessary before an embolectomy can be attempted. Pulmonary embolism (PE) is determined by a mismatch between the ventilation and perfusion images. In other words, a normal ventilation image and an abnormal perfusion image with segmental defects indicate PE.

Clinical Alert

Pulmonary perfusion imaging is contraindicated in patients with primary pulmonary hypertension.

Reference Values

Normal Normal functioning lung Normal pulmonary vascular supply Normal gas exchange

Procedure

1. Ask the patient to breathe for approximately 4 minutes through a closed, nonpressurized ventilation system. During this time, administer a small amount of radioactive gas or aerosol. It is important that the patient not swallow the radioactive aerosol during the ventilation portion of the lung imaging. Doing so causes radioactive interference with the lower lobes of the lung and makes an accurate diagnostic interpretation difficult. Also, take care that the patient does not aspirate the aerosol.
2. Alert the patient that breath holding will be required for a brief period at some time during the imaging.
3. Be aware that the imaging time is 10 to 15 minutes. When the ventilation imaging is performed with lung perfusion imaging (eg, in differential diagnosis of PE), the testing time is 30 to 45 minutes.
4. Perform the perfusion imaging immediately after the ventilation study.
5. In the pediatric patient, reduce the number of particles given in the MAA dose because of the smaller size of the capillary beds. Use caution with MAA in patients with atrial and ventricular septal defects.
6. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. Abnormal ventilation and perfusion patterns indicate possible:
   a. Tumors
   b. Emboli
   c. Pneumonia
   d. Atelectasis
e. Bronchitis  
f. Asthma  
g. Inflammatory fibrosis  
h. Chronic obstructive pulmonary disease  
i. Lung cancer  

2. In pediatric patients, there is an increased incidence of an airway obstruction caused by mucus plugs or foreign bodies. However, pulmonary emboli do not occur in children as often as in adults.

Interfering Factors

1. False-positive imagings occur in vasculitis, mitral stenosis, and pulmonary hypertension and when tumors obstruct a pulmonary artery with airway involvement.  
2. During the injection of MAA, care must be taken that the patient’s blood does not mix with the radiopharmaceutical in the syringe. Otherwise, hot spots may be seen in the lungs.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of the test.  
2. Alleviate any fears the patient may have concerning nuclear medicine procedures.  
3. Be aware that it is important that a recent chest radiograph be available.  
4. Remember that the patient must be able to follow directions for breathing and holding the breath, including breathing through a mouthpiece or into a face mask.  
5. Refer to standard nuclear scan pretest precautions on page 655.  
6. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Refer to standard nuclear scan posttest precautions on page 655.  
2. Interpret test outcome and monitor appropriately for postprocedural signs of aspiration.  
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

ORTHOPEDIC STUDIES

Bone Imaging

This test is used primarily to evaluate and monitor persons with known or suspected metastatic disease. Breast cancers, prostate cancers, lung cancers, and lymphomas tend to metastasize to bone. Bone imagings visualize lesions 6 to 12 months before they appear on radiographs. Bone imaging may also be performed to evaluate patients with unexplained bone pain, primary bone tumors, arthritis, osteomyelitis, abnormal healing of fractures, fractures, shin splints, or compression fractures of the vertebral column; to evaluate pediatric patients with hip pain (Legg-Calvé-Perthes disease); and to assess child abuse, bone growth plates, sports injuries, and stress fractures. It is also performed to determine the age and metabolic activity of traumatic injuries and infections.

Other indications are evaluation of candidates for knee and hip prostheses, diagnosis of aseptic necrosis and vascularity of the femoral head, presurgical and postsurgical assessment of viable bone tissue, and evaluation of prosthetic joints and internal fixation devices to rule out loosening of prosthesis or infection.

Bone imaging has greater sensitivity in the pediatric patient than in the adult and is used for early detection of trauma. Normally, there is increased activity in the growth plates of the long bones. The child’s history is significant for correlation and diagnostic differentiation. In older children with unexplained pain, who participate in sports, stress fractures are often found on bone imaging.

A bone-seeking radiopharmaceutical is used to image the skeletal system. An example is $^{99m}$Tc-labeled phosphate injected intravenously. Imaging usually begins 2 to 3 hours after injection. Abnormal pathology, such as increased blood flow to bone or increased osteocytic activity, concentrates the radiopharmaceutical at a higher or lower rate than the normal bone does. The radiopharmaceutical mimics calcium physiologically; therefore, it concentrates more heavily in areas of increased metabolic activity.

Reference Values

Normal Homogenous distribution of radiopharmaceutical

Procedure

1. Inject radioactive $^{99m}$Tc methylene diphosphonate (MDP) intravenously.  
2. Be aware that a 2- to 3-hour waiting period is necessary for the radiopharmaceutical to concentrate in the bone. During this time, the patient may be asked to drink 4 to 6 glasses of water.  
3. Before the imaging begins, ask the patient to void because a full bladder masks the pelvic bones.  
4. Remember that imaging takes about 30 to 60 minutes to complete. The patient must be able to lie still during the imaging.  
5. Be aware that additional spot views of a specific area or 3-dimensional SPECT imaging may be requested by the physician.  
6. See Chapter 1 guidelines for safe, effective, informed intratest care.
Clinical Alert
For osteomyelitis, images are acquired during the injection of the radiopharmaceutical, thus giving the image of the blood flow to the bone.

Clinical Implications Abnormal concentrations indicate the following:

1. Very early bone disease and healing is detected by nuclear medicine bone images long before it is visible on radiographs. Radiographs are positive for bone lesions only after 30% to 50% decalcification (decrease in bone calcium) has occurred.
2. Many disorders can be detected but not differentiated by this test (eg, cancer, arthritis, benign bone tumors, fractures, osteomyelitis, Paget's disease, aseptic necrosis). The findings must be interpreted in light of the whole clinical picture because any process inducing an increased calcium excretion rate will be reflected by an increased uptake in the bone.
3. In patients with breast cancer, the likelihood of a positive bone image finding in the preoperative period depends on the staging of the disease, and imaging tests are recommended before initial therapy. Stages 1 and 2: 40% have a positive bone image. Stage 3: 19% have a positive bone image. Yearly nuclear medicine bone imaging should be done for follow-up.
4. Multiple myeloma is the only tumor that shows better detectability with a plain radiograph than a radionuclide bone procedure.
5. Multiple focal areas of increased activity in the axial skeleton are commonly associated with metastatic bone disease. The reported percentage of solitary lesions due to metastasis varies on a site-by-site basis. With a single lesion in the spine or pelvis, the cause is more likely to be metastatic disease than with a single lesion occurring in the extremities or ribs.

Clinical Alert
The “flare phenomenon” occurs in patients with metastatic disease who are receiving a new treatment. The bone imaging may show increased activity or new lesions in patients with clinical improvement. This is caused by a healing response in patients with prostate or breast cancer within the first few months of starting a new treatment. These lesions should show marked improvement on imagings taken 3 to 4 months later.

Interfering Factors
1. False-negative bone images occur in multiple myeloma of the bone. When this condition is known or suspected, the bone image is an unreliable indicator of skeletal involvement.
2. Patients with follicular thyroid cancer may harbor metastatic bone marrow disease, but these lesions are often missed by bone scans.

Interventions

Pretest Patient Care
1. Instruct the patient about the purpose and procedure of the test. Alleviate any fears concerning the procedure. Advise the patient that frequent drinking of fluids and activity during the first 6 hours help to reduce excess radiation to the bladder and gonads.
2. Remember that the patient can be up and about during the waiting period. There are no restrictions during the day before imaging.
3. Remind the patient to void before the imaging. If the patient is in pain or debilitated, offer assistance to the restroom.
4. Order and administer a sedative to any patient who will have difficulty lying quietly during the imaging period.
5. Refer to standard nuclear scan pretest precautions on page 655. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Advise the patient to empty the bladder when imaging is completed, to decrease radiation exposure time.
2. Refer to standard nuclear scan posttest precautions on page 655.
3. Interpret test outcome and monitor appropriately.
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Bone Mineral Density (Bone Densitometry; Osteoporosis Imaging)
Bone densitometry enables the clinician to obtain a diagnosis of osteoporosis or osteopenia, often before fractures occur, by measuring bone mineral density. No radiopharmaceuticals are used in this procedure, but special imaging techniques are used. X-ray absorptiometry for measuring bone mineral density includes these special modalities:

1. Dual-energy absorptiometry (DEXA or DXA) to measure spine, hip, and forearm density
2. Peripheral dual-energy absorptiometry (pDXA) to measure forearm density
3. Single-energy x-ray absorptiometry (SXA) to measure the heel and forearm density
4. Radiographic absorptiometry (RA) to measure the density of the phalanges

DEXA is the most common and preferred method of measuring bone mineral density because of its precision and low
radiation exposure. With the use of laser x-ray imaging and specific computer software, DEXA can assess fracture risk with relative ease and patient comfort. Fracture risk is measured in standard deviations (SDs) by comparing the patient's bone mass to that of healthy 25- to 35-year-old persons. Test scores are printed out and reported with a T-score and a Z-score. The T-score is the number of SDs for the patient compared with normal young adults with mean peak bone mass. Fracture risk increases about 1.5 to 2.5 times for every SD. According to the World Health Organization, T-scores of less than 2.5 may confirm a diagnosis of osteoporosis; scores of 2.5 to 1.0 are associated with osteopenia; and scores of 1.0 or greater are considered normal. The Z-score is defined as the number of SDs for the patient compared with normal persons in the same age category. The T-score is the score most commonly reported and currently is the preferred reference point for diagnosing osteoporosis.

Reference Values

**Normal** Absence of osteoporosis or osteopenia T-score: <1.0 SD below normal (>1.0) Osteopenia 1.0 to 2.5 SD below normal (-1.0 to -2.5) Osteoporosis >2.5 SD below normal (<-2.5)

**Procedure**

1. Position the patient in such a way as to keep the area being imaged immobile.
2. Place a foam block under both knees during the spine imaging. Use a leg brace immobilizer during the femur imaging, and use an arm brace when imaging the forearm.
3. Be aware that DEXA images of the spine and hip take approximately 20 minutes to complete. An additional 15 minutes is needed to image the forearm.
4. See Chapter 1 guidelines for safe, effective, informed intratest care.

**Clinical Implications** Abnormal imagings may be associated with the following:

1. Estrogen deficiency in postmenopausal women
2. Vertebral abnormalities
3. Patients with radiographic osteopenia
4. Hyperparathyroidism
5. Patients receiving long-term corticosteroid therapy

**Interfering Factors** False readings may occur with the following:

1. Nuclear medicine imagings within the previous 72 hours (longer for gallium or indium imagings) may cause residual emission that can be misinterpreted.
2. Barium studies within the previous 7 to 10 days may interfere with the spine imaging.
3. Prosthetic devices or metallic objects surgically implanted in areas of interest may interfere with the image.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure for measuring bone density of spine, hip, forearm, heel, and phalanges. No radiopharmaceuticals are administered.
2. Encourage patients to wear cotton garments that are free of metal or plastic zippers or buttons.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret abnormal test outcome. If needed, serial studies may be ordered to measure the effectiveness of treatment.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Clinical Alert**

1. Bone densitometry tests use x-rays: the precautions outlined in Chapter 10 should be noted.
2. Additional means, of measuring bone mineral density arcs
   a. Quantitative computed tomography (QCT) to measure spine density
   b. Peripheral quantitative computed tomography (pQC) to measure forearm density

**TUMOR IMAGING STUDIES**

**Gallium (^{67}Ga) Imaging**

This image is used to detect the presence, location, and size of lymphoma, to detect chronic infections and abscesses, to differentiate malignant from benign lesions, and to determine the extent of invasion of known malignancies. The entire body is imaged looking for lymph node involvement. In both adult and pediatric patients, these studies are used to help stage bronchogenic cancer, Hodgkin's lymphomas, and non-Hodgkin's lymphomas. Gallium imagings may also be used to record tumor regression after radiation or chemotherapy. The radionuclide used in this study is gallium citrate (^{67}Ga).

The underlying mechanism for the uptake of ^{67}Ga is not well understood. Uptake in some neoplasms may depend on the presence of transferrin receptors in tumor cells, but this is only speculation. Once gallium enters a tissue, it remains there until radioactive decay dissipates the isotope.

**Reference Values**
No evidence of tumor-type activity or infection

Procedure

1. Give a laxative the evening before the imaging.
2. Be aware that laxatives, suppositories, and/or tap water enemas are often ordered before imaging. The patient may eat breakfast on the day of imaging.
3. Inject the radionuclide 24 to 96 hours before imaging.
4. Have the patient lie quietly without moving during the imaging procedure. Take anterior and posterior views of the entire body.
5. Remember that additional imaging may be done at 24-hour intervals to differentiate normal bowel activity from pathologic concentrations.
6. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. Abnormal gallium concentration usually implies the existence of underlying pathology:
   a. Malignancy, especially lung, testes, and mesothelioma
   b. Stages of lymphoma, Hodgkin’s disease, melanoma, hepatoma, soft tissue sarcoma, primary tumor of bone or cartilage, neuroblastoma, and leukemia
   c. Abscesses
   d. Tuberculosis
   e. Thrombosis
   f. Abscessed sarcoidosis
   g. Chronic infection
   h. Interstitial pulmonary fibrosis
2. Further diagnostic studies usually are performed to distinguish benign from malignant lesions.
3. Tumor uptake of $^{67}$Ga varies with tumor type, among persons with tumors of the same histologic type, and even among tumor sites of a given patient.
4. Tumor uptake of $^{67}$Ga may be significantly reduced after effective treatment.
5. Although $^{111}$In-labeled leukocyte imaging is more specific for acute abscess localization, gallium imaging may be used as a multipurpose screening procedure for chronic infection.

Interfering Factors

1. A negative study cannot be definitely interpreted as ruling out the presence of disease. (The rate of false-negative results in gallium studies is 40%.)
2. It is difficult to detect a single, solitary nodule (eg, adenocarcinoma). Lesions smaller than 2 cm can be detectable. Tumors near the liver are difficult to detect, and interpretation of iliac nodes is difficult.
3. Because gallium does collect in the bowel, there may be an abnormal concentration in the lower abdomen. For this reason, laxatives and enemas may be ordered.
4. Degeneration or necrosis of tumor and use of antineoplastic drugs immediately before imaging cause false-negative results.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of the gallium imaging.
2. Remember that usually, no change in eating habits is required before testing. However, some departments request that their patients eat a low-residue lunch and a clear-liquid supper the day before the examination.
3. See standard nuclear scan pretest precautions on page 655.
4. Be aware that the usual preparation includes oral laxatives taken on the night before the first imaging session and again on the night before each imaging session. Enemas or suppositories may also be given. These preparations clean normal gallium activity from the bowel.
5. Be aware that actual imaging time is 45 to 90 minutes per imaging session.
6. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Refer to standard nuclear scan posttest precautions on page 655.
2. Interpret test outcome and monitor appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

Breastfeeding should be discontinued for at least 4 weeks after testing.

OVERVIEW OF MONOCLONAL ANTIBODY TUMOR IMAGING (ONCOSCINT, PROSTASCINT, CARCINOEMBRYONIC ANTIGEN, OCTREOTIDE AND OTHER PEPTIDES)

These classes of tumor imaging have revolutionized the foundation of radiopharmaceutical production. Like other radiopharmaceuticals, monoclonal antibodies (MABs) and peptides have two parts: a radioisotope linked to a substance specific to a target organ. In the case of MABs, that substance is an antibody that has been cloned and mass produced.
Because all the daughter antibodies are identical, a high yield of very specific antibodies can be produced.

1. OncoScint MAB. This was the first monoclonal antibody radiopharmaceutical to be approved by the U.S. Food and Drug Administration and mass marketed. OncoScint was approved for the detection of ovarian and colon cancer. OncoScint is an antibody linked to $^{111}$In.

2. ProstaScint MAB. ProstaScint is a monoclonal antibody approved for the detection of lymph node metastasis from prostate cancer.

3. Carcinoembryonic antigen (CEA) MAB. This monoclonal antibody is similar to OncoScint in that an antibody to CEA is used. The radioisotope, however, is $^{99m}$Tc pertechnetate. This anti-CEA antibody produces a different pattern of biodistribution than OncoScint does. Each radiopharmaceutical has its own indications.

4. Octreotide-peptide. This radiopharmaceutical peptide is used for localizing neuroendocrine tumors.

Antibody and Peptide Tumor Imagings

Antibody and peptide tumor imaging is used to detect the location and size of known extrahepatic malignancies. These procedures are not screening techniques.

Reference Values

Normal Distribution occurs in the normal liver, spleen, bone marrow, and bowel.

Procedure

1. Inject the patient with the radioisotope over a period of 5 minutes. Observe the patient for any reaction to the radiopharmaceutical.
2. Remember that optimal whole-body images are obtained between 2 and 4 days after injection; additional images may be obtained at 24 hours and at 5 days.
3. Perform SPECT imaging if necessary.
4. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. Abnormal distributions are found in tumors. Any change in the distribution provides information regarding the effectiveness of surgery or therapy.
2. Abnormal results have been observed in nonspecific areas such as inflammatory bowel disease, colostomy sites, and postoperative bowel adhesions.
3. The patient's medical history should be reviewed carefully.

Interfering Factors Radioactivity in the bowel may interfere with colorectal assessment. Follow-up imaging is useful after administration of a cathartic to clarify equivocal findings.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks.
2. Refer to standard nuclear scan pretest precautions on page 655.
3. Establish an intravenous line before injecting the radiopharmaceutical.
4. Be aware that a cathartic is required to differentiate bowel activity from abnormal pathology.
5. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Refer to standard nuclear scan posttest precautions on page 655.
2. Interpret test outcome and monitor appropriately.
3. Observe the patient for 1 hour after injection of OncoScint for antibody reactions (eg, chills, fever, nausea).
4. Realize that some patients develop human antimouse antibody (HAMA) titers after OncoScint injection.
5. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

Following OncoScint imaging, HAMA titers may result in falsely elevated immunoassay levels for CA-125 and CEA.

Iodine-131 Whole-Body (Total-Body) Imaging

Whole-body imaging using $^{131}$I can identify functioning thyroid tissue throughout the body. It is useful to determine the presence of metastatic thyroid cancer and the amount and location of residual tissue after thyroidectomy. The procedure is routinely performed in conjunction with thyroid therapy using $^{131}$I for thyrocarcinoma.

Reference Values

Normal No functioning extrathyroid tissues outside of the thyroid gland

Procedure

1. Administer radionuclide orally in a capsule form.
2. Perform imaging 24 to 72 hours after administration of the radiopharmaceutical. Imaging may take as long as 2 hours to perform.
3. Remember that sometimes, thyrotropin (thyroid-stimulating hormone, or TSH) is administered intravenously before the radionuclide is given. This stimulates any residual thyroid tissue and enhances $^{131}$I uptake.

4. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications Abnormal uptake of iodine reveals:

1. Areas of extrathyroid tissue such as:
   a. Stroma ovarii
   b. Substernal thyroid
   c. Sublingual thyroid
2. Residual tissue after thyroidecтомy
3. Metastatic thyroid cancer

Clinical Alert

1. If possible, this test should be performed before any other radionuclide procedures and before use of antiodine contrast medium, surgical preparation, or other form of iodine.
2. The test is most effective when endogenous TSH levels are high, so as to stimulate radionuclide uptake by metastatic neoplasms.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks.
2. Advise the patient that the imaging process may take several hours. If iodine allergies are suspected, observe the patient for possible reactions.
3. Refer to standard nuclear scan, pretest precautions on page 655.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Refer to standard nuclear scan posttest precautions on page 655.
2. Interpret test outcome and monitor appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Breast Imaging (Scintimammography); Lymph Node Imaging (Lymphoscintigraphy)

Although x-ray mammography is the preferred examination for routine breast screening, scintimammography is often used in cases of indeterminate mammography. Other indications for performing breast imaging include follow-up to surgery, biopsy, radiation therapy, or chemotherapy. The radiopharmaceutical used for scintimammography is $^{99m}$Tc miraluma. The test is more specific than x-ray mammography and may differentiate between benign and malignant lesions. The test is also used to detect any axillary lymph node involvement from breast cancer and decreases the number of unnecessary breast biopsies. Sentinel and satellite node identification and staging in early breast cancer using lymphoscintigraphy and intraoperative gamma node and tissue biopsy show micrometastasis more frequently than standard dissection. Lymphoscintigraphy assesses the lymphatic drainage of tumors.

Reference Values

**Normal Breast** Uniform distribution of radiopharmaceutical uptake in the breasts without focal points of concentration No focal uptake in lymphatic tissue

**Normal Lymph Node** No abnormal nodes (indicated by obstruction to tracer). (The first node the tracer goes to is identified.)

Procedure for Breast Imaging

1. Inject the radiopharmaceutical intravenously in the opposite arm from the breast of concern.
2. Have the patient lie prone on a special table with a cut out section that allows the breasts to hang through the table unobstructed.
3. Place the patient in the supine position with the arms raised for obtaining images of the axillary lymph nodes.
4. Be aware that the total patient time is approximately 45 to 60 minutes. The actual scan time is 25 to 30 minutes.
5. For sentinel node identification, see Chapter 11 for a complete discussion of the procedure.
6. Remember that an optional SPECT examination may be requested by the nuclear medicine physician. This examination may take an additional 30 to 40 minutes.
7. See Chapter 1 guidelines for safe, effective, informed intratest care.

Procedure for Lymph Node Imaging

1. Special positioning is required. (See sentinel node evaluation in Chapter 11.)
2. The tracer is injected by intradermal method for melanoma evaluation or subcutaneously for breast cancer evaluation. Massage after injection for at least 30 seconds, moving the breast to optimally clear the overlying soft tissue.
3. Images are obtained immediately and 2–4 hours after injection.
4. Sentinel lymph node identification may also be performed to evaluate metastatic spread of cancer to penis, vulva, uterus, head, and neck.

Clinical Implications of Breast Imaging
1. Abnormal increased focal uptake is found in cases of a fibroadenoma and adenocarcinoma.
2. Nonuniform increased diffuse uptake of activity is associated with fibrous dysplasia, which may be unilateral or bilateral.
3. Several areas of increased focal uptake are often seen in cases of multifocal breast cancer.
4. In patients with a breast prosthesis, a focal decrease in activity is observed in relation to the size and shape of the prosthesis.
5. Axillary metastasis is detected as focal areas of increased uptake in the axillary nodes.
6. This scan is used to evaluate radiation therapy and chemotherapy.

Clinical Implications of Lymph Node Imaging

1. Abnormal nodes show leaks into adjacent tissue, a blush around the affected node, and unusual collateral lymph drainage pathways.
2. The first lymph node to drain the tumor invariably contains the tumor.
3. It has been found that there are more than one lymphatic channels draining the tumor, and that there are one, two, or three sentinel lymph nodes (SLNs) as well as satellite nodes.
4. Micrometastasis of biopsied tissue is found more frequently than standard axillary node dissection.

Interfering Factors

1. There should not be any other detectable amount of radioactivity in the patient.
2. The patient should be lying supine for the injection of the radiopharmaceutical (for breast imaging) to prevent a “streaking” artifact found on the resulting image in the breast region, which corresponds to the arm that received the injection.
3. To eliminate a false-positive appearance, the patient should be injected on the side opposite of a known lymphatic lesion. If the patient is known to have bilateral breast cancer, a foot vein may be used for injection.
4. Extravasation of the radiopharmaceutical can result in hot spots of radioactivity in the location of the axillary lymph nodes.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of the nuclear scan. See Chapter 11 for more information on sentinel node biopsy.
2. Have the patient remove all clothing and jewelry from the waist up. The patient wears a hospital gown with the opening of the gown in the front. There are no dietary or medication restrictions. For lymph node imaging, position patient as she would be placed for surgical intervention.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.
4. See standard nuclear scan pretest precautions on page 655.

Posttest Patient Aftercare

1. Interpret test outcome, monitor, and counsel appropriately about need for further tests (eg, biopsy and possible immediate surgery).
2. Refer to standard nuclear scan posttest precautions on page 655.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

INFLAMMATORY PROCESS IMAGING

Leukocyte (WBC) Imaging (Indium- or Ceretec-Labeled WBCs)

The leukocyte imaging test, in which a sample of the patient's own white blood cells (WBCs) are isolated, labeled with indium oxine (111In) or 99mTc exametazime, and reinjected, is used for localization of acute abscess formation. The study is indicated in both adults and children with signs and symptoms of a septic process, fever of unknown origin, osteomyelitis, or suspected intraabdominal abscess. It is also helpful in determining the cause of complications of surgery, injury, or inflammation of the gastrointestinal tract and pelvis. The test results are based on the fact that any collection of labeled WBCs outside the liver, spleen, and functioning bone marrow indicates an abnormal area to which the cells localize. This procedure is 90% sensitive and 90% specific for acute inflammatory disease or acute abscess formation.

Reference Values

Normal Normal leukocyte concentration and radiopharmaceutical distribution in liver, spleen, and bone marrow No signs of leukocyte localization outside of the reticuloendothelial system

Procedure

1. Obtain a venous blood sample of 60 mL for the purpose of isolating and labeling the WBCs. The laboratory process takes about 2 hours to complete. The patient's WBC count needs to be at least 4.0 so that there are enough cells to label for this procedure.
2. Label the WBCs with radioactive 111In, oxine, or 99mTc exametazime and inject intravenously.
3. Have the patient return for imaging after 4 hours with Ceretec and after 24 or 48 hours with indium.
4. Be aware that imaging time is about 1 hour each session.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.
Clinical Implications Abnormal concentrations indicate:

1. Acute abscess formation
2. Acute osteomyelitis and infection of orthopedic prostheses
3. Active inflammatory bowel disease
4. Postsurgical abscess sites and wound infections

Interfering Factors

1. False-negative reactions are known to occur when the chemotactic function of the WBC has been altered, as in hemodialysis, hyperglycemia, hyperalimentation, steroid therapy, and long-term antibiotic therapy.
2. Gallium scans up to 1 month before the test can interfere.
3. False-positive scans occur in the presence of gastrointestinal bleeding and in upper respiratory infections and pneumonitis when patients swallow purulent sputum.

NOTE
See Clinical Considerations, Pretest Patient Care, and Posttest Patient Aftercare for nuclear scans on pages 654–655.

Clinical Alert
If the patient does not have an adequate number of WBCs, additional blood may have to be drawn. Gallium imaging may be necessary if too few WBCs are present, or donor cells can be used.

Part 2 Radionuclide Laboratory Procedures (Nonradioimmunoassay Studies)

OVERVIEW OF LABORATORY PROCEDURES

Very small amounts of radioactive substances may be administered to patients, and subsequently their body fluids and glands may be examined in the laboratory for concentrations of radioactivity. Minute quantities of radioactive materials may be detected in blood, feces, urine, other body fluids, and glands.

Some procedures (eg, Schilling test) check the ability of the body to absorb the administered radioactive compound. Others, such as blood volume determinations, test the ability of the body to localize or dilute the administered radioactive substance.

Part 2 of this chapter includes a sampling of tests that employ the use of radionuclides in the study of disease. Imaging may or may not be required as part of these procedures, which are all a form of tracer chemistry.

Schilling Test The Schilling test is a 24-hour urine test that is used to diagnose pernicious anemia (one form of macrocytic anemia) and malabsorption syndromes. It is an indirect test of intrinsic factor deficiency. This test evaluates the body's ability to absorb vitamin B\textsubscript{12} from the gastrointestinal tract and is based on the anticipated urinary excretion of radioactive vitamin B\textsubscript{12}. The procedure may be done in two stages: stage I, without intrinsic factor; and stage II, with intrinsic factor. The second stage is performed only when an abnormal first stage occurs. In stage I, the fasting patient is given an oral dose of vitamin B\textsubscript{12} tagged with radioactive cobalt (\textsuperscript{57}Co). An intramuscular injection of vitamin B\textsubscript{12} is given to saturate the liver and serum protein-binding sites, which allows radioactive vitamin B\textsubscript{12} to be excreted in the urine. A 24-hour urine specimen is then collected. The amount of the excreted radioactive B\textsubscript{12} is determined and expressed as a percentage of the given dose. Normal persons absorb (and therefore excrete) as much as 25% of the radioactive B\textsubscript{12}. Patients with pernicious anemia absorb little of the oral dose and therefore excrete little radioactive material in the urine.

Reference Values
Normal Excretion of 10% or more of the dose of cobalt-tagged vitamin B\textsubscript{12} in the urine within 24 hours Normal values assume near-normal kidney function

Procedure

1. Have the patient fast for 12 hours before the test. (Fasting is continued for 3 hours after the vitamin B\textsubscript{12} doses have been administered.)
2. Administer a tasteless capsule of radioactive B\textsubscript{12} labeled with \textsuperscript{57}Co orally by a nuclear medicine technologist.
3. Have a registered nurse or nuclear medicine technologist inject nonradioactive B\textsubscript{12} intramuscularly. This is done 2 hours later.
4. Collect a small sample of urine before the study begins. The patient also voids just before injection with nonradioactive B\textsubscript{12} (called a flushing dose). Collect all urine for 24 or 48 hours after the time the patient receives the injection of vitamin B\textsubscript{12}.
   a. Obtain a special 24-hour urine container from the laboratory. No preservative is required.
   b. Ensure that there is no contamination of the urine with stool.
   c. Continue collecting the urine for 24 hours (see Chapter 3).
4. In the presence of renal disease, a 48-hour urine collection may be necessary.
5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications
1. An abnormally low value (eg, <5%) or borderline (5.0%–9.9%) allows two interpretations:
   a. Absence of intrinsic factor
   b. Defective absorption in the ileum
2. When the absorption of radioactive vitamin B\textsubscript{12} is low from the first stage, the test must be repeated with intrinsic factor (stage II) to rule out intestinal malabsorption (confirmatory Schilling test).
   a. If urinary excretion then rises to normal levels, it indicates a lack of intrinsic factor, suggesting the diagnosis of pernicious anemia.
   b. If the urinary excretion does not rise, malabsorption is considered to be the cause of the patient’s anemia.

**NOTE**

A dual-radiouclide test is an alternative method in which both stages are performed at the same time.

**Interfering Factors**

1. Renal insufficiency may cause reduced excretion of radioactive vitamin B\textsubscript{12}. If renal insufficiency is suspected, a 48- to 72-hour urine collection is advised because eventually almost all of the absorbed material will be excreted, and urine specific gravity and volume are checked.
2. The patient should not undergo diagnostic procedures that interfere with B\textsubscript{12} absorption.
3. The single most common source of error in performing the test is incomplete collection of urine. Some laboratories may require a 48-hour collection to allow for a small margin of error.
4. Urinary excretion of B\textsubscript{12} is depressed in elderly patients, diabetic patients, patients with hypothyroidism, and patients with enteritis.
5. Fecal contamination in the urine leads to false results and invalidates the test.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose, procedure, benefits, and risks of the test.
2. Refer to general procedures, description of benefits, risks, clinical considerations, and standard nuclear scan pretest precautions on page 655.
3. Obtain a random urine sample before the vitamin B\textsubscript{12} doses are administered.
4. Give a written reminder to the patient about fasting and collection of a 24-hour urine specimen. Water is permitted during the fasting period.
5. Permit food and drink 3 hours after the doses of vitamin B\textsubscript{12} are given. Encourage the patient to drink as much as can be tolerated during the entire test.
6. Be certain the patient receives the nonradioactive B\textsubscript{12}. If the intramuscular dose of vitamin B\textsubscript{12} is not given, the radioactive vitamin B\textsubscript{12} will be found in the liver instead of the urine.
7. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Clinical Alert**

1. No laxatives are to be used during the test.
2. Bone marrow aspiration should be done before the Schilling test because the vitamin B\textsubscript{12} administered in the test destroys the diagnostic characteristics of the bone marrow.

**Posttest Patient Aftercare**

1. Assess for compliance with 24-hour urine collection protocols (see Chapter 3).
2. Refer to standard nuclear scan posttest precautions on page 655.
3. Interpret test outcome and monitor appropriately.
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Total Blood Volume; Plasma Volume; Erythrocyte (RBC) Volume** The purposes of the blood volume test are to determine circulating blood volume, to help evaluate the bleeding or debilitated patient, and to determine the origin of hypotension in the presence of anuria or oliguria when dehydration may be the cause. This determination is one way to monitor blood loss during surgery; it is used as a guide in replacement therapy after blood or body fluid loss and in the determination of whole-body hematocrit. The results are useful in choosing the most appropriate blood component for replacement therapy (whole blood, plasma, or packed RBCs). Total blood volume determinations are of value in the following situations:

1. To evaluate gastrointestinal and uterine bleeding
2. To aid in the diagnosis of hypovolemic shock
3. To aid in the diagnosis of polycythemia vera
4. To determine the required blood component for replacement therapy, as in persons undergoing surgery

These tests reveal an increased or decreased volume of RBC mass. A sample of the patient's blood is mixed with a radioactive substance, incubated at room temperature, and reinjected. Another blood sample is obtained 15 minutes later. The most commonly used tracers in blood volume determinations are serum albumin tagged with $^{131}$I or $^{125}$I and patient or donor RBCs tagged with chromium-51 ($^{51}$Cr). The combination of procedures (total blood volume) is the only true blood volume. Other volume studies are plasma volume and RBC volume, which may be done separately. The plasma volume is used to establish a vascular baseline, to determine changes in plasma volume before and after
surgery, and to evaluate fluid and blood replacement in patients with gastrointestinal bleeding, burns, or trauma. The 
\(^{51}\text{Cr} \) RBC volume study is done to see what percentage of the circulating blood is composed of RBCs. This procedure is
performed in connection with evaluation of RBC survival or gastrointestinal blood loss and in ferrokinetic studies. These tests can be done simultaneously.

**Reference Values**

**Normal**
- Total blood volume: 55–80 mL/kg or 0.055–0.080 L/kg
- Erythrocyte volume: 20–35 mL/kg or 0.020–0.035 L/kg (greater in men than in women)
- Plasma volume: 30–45 mL/kg or 0.030–0.045 L/kg

**NOTE**

Because adipose tissue has a sparser blood supply than lean tissue, the patient's body type can affect the proportion of blood volume to body weight; for this reason, test findings should always be reported in milliliters per kilogram of body weight.

**Procedure**

1. Record the patient's height and current weight.
2. Obtain venous blood samples, and mix one blood sample with a radionuclide.
3. Fifteen to 30 minutes later, reinject the blood radiopharmaceutical.
4. About 15 minutes later, obtain another venous blood sample and have it examined in the laboratory.
5. See Chapter 1 guidelines for **safe, effective, informed intratest care**.

**Clinical Implications**

1. A normal total blood volume with a decreased RBC content indicates the need for a transfusion of packed red cells.
2. Polycythemia vera may be differentiated from secondary polycythemia.
   a. Increased total blood volume due to an increased RBC mass suggests polycythemia vera. The plasma volume most often is normal.
   b. Normal or decreased total blood volume due to a decreased plasma volume suggests secondary polycythemia. The RBC most often is normal.

**Clinical Alert**

If intravenous blood component therapy is ordered for the same day, the blood volume determination should be done before the intravenous line is started.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose, procedure, benefits, and risks of the test. Blood samples and intravenous injection are part of this test. No imaging or scanning takes place.
2. Weigh the patient just before the test if possible.
3. Refer to standard nuclear scan **pretest precautions** on page 655.
4. See Chapter 1 guidelines for **safe, effective, informed pretest care**.

**Posttest Patient Aftercare**

1. Refer to standard nuclear scan **posttest precautions** on page 655.
2. Interpret test outcome and monitor appropriately.
3. Follow Chapter 1 guidelines for **safe, effective, informed posttest care**.

**Red Blood Cell (RBC) Survival Time Test**

The RBC survival time test has its greatest use in the evaluation of known or suspected hemolytic anemia and is also indicated when the cause for anemia is obscure (abnormal sequestration of RBCs in the spleen), to identify accessory spleens, and to determine abnormal RBC production or destruction. A sample of the patient's erythrocytes is mixed with a radioactive substance (\(^{51}\text{Cr}\)), incubated at room temperature, and reinjected. Blood specimens are drawn after 24 hours and at regular intervals for at least 3 weeks. After counting the specimens, the results are plotted, and the RBC survival time is calculated. Results are based on the fact that disappearance of radioactivity from the circulation corresponds to disappearance of the RBCs, thereby determining overall erythrocyte survival. Scanning of the spleen is often done as part of this test. The RBC survival test usually is ordered in conjunction with a blood volume determination and radionuclide iron uptake and clearance tests. When stool specimens are collected for 3 days, the test is often referred to as the "gastrointestinal blood loss test."

**Reference Values**

**Normal**
- Normal half-time for survival of \(^{51}\text{Cr}\)-labeled red blood cells is approximately 25 to 35 days.
- \(^{51}\text{Cr}\) in stool: <3 mL/24 hours

**Procedure**

1. Obtain a venous blood sample of 20 mL.
2. Ten to 30 minutes later, reinject the blood after being tagged with a radionuclide, \(^{51}\text{Cr}\).
3. Remember that blood samples are usually obtained on the first day; again after 24, 48, 72, and 96 hours; and then at weekly intervals for 3 weeks. Time may be shortened depending on the outcome of the test. As part of this procedure, a radioactive detector may be used over the spleen, sternum, and liver to assess the relative concentrations of radioactivity in these areas. This external counting helps to determine whether the spleen is taking part in excessive sequestration of RBCs as a causative factor in anemia.
4. Be aware that in some instances, a 72-hour stool collection may be ordered to detect gastrointestinal blood loss. Obtain special collection containers labeled for radiation hazard. At the end of each 24-hour collection period, the
total stool is to be collected by the department of nuclear medicine. This test can be completed in 3 days.

5. See Chapter 1 guidelines for safe, effective, informed intratest care.

Clinical Implications

1. Shortened RBC survival may result from blood loss, hemolysis, or removal of RBCs by the spleen, as in:
   a. Chronic granulocytic leukemia
   b. Hemolytic anemia
   c. Hemoglobin C disease
   d. Hereditary spherocytosis
   e. Pernicious anemia
   f. Megaloblastic anemia of pregnancy
   g. Sickle cell anemia
   h. Uremia
2. Prolonged RBC survival time may result from an abnormality in RBC production, as in thalassemia minor, and false-negative results when transfusion is given during the procedure.
3. If hemolytic anemia is diagnosed, further studies are needed to establish whether the RBCs have intrinsic abnormalities or whether anemia results from immunologic effects of the patient's plasma.
4. Results are normal in:
   a. Hemoglobin C trait
   b. Sickle cell trait
5. Half of the radioactivity in the plasma may not disappear for 7 to 8 hours.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test. Emphasize that this test requires a minimum of 2 weeks of the patient's time, with trips to the diagnostic facility for venipunctures.
2. If stool collection is required, advise the patient of the importance of saving all stool and that stool must be free of urine contamination.
3. Refer to standard nuclear scan pretest precautions on page 655.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert

1. The test usually is contraindicated in a patient who is actively bleeding.
2. Record and report signs of active bleeding.
3. Transfusions should not be given while the test is in progress. If it is necessary to do so, notify the nuclear medicine department to terminate the test.

Posttest Patient Aftercare

1. Refer to standard nuclear scan posttest precautions on page 655.
2. Interpret test outcome and monitor appropriately. Explain need for further testing and possible treatment (spleenectomy).
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Part 3 Positron Emission Tomography

Positron Emission Tomography (PET) Imaging

PET imaging involves the combined use of positron-emitting radionuclides and emission CT. PET technology generates high-resolution images of body function and metabolism. PET uses radiopharmaceuticals that are the basic elements of biologic substances. In this way, normal and abnormal biologic function of cells and organs can be determined. It produces images of molecular-level physiologic function, including glucose metabolism, oxygen utilization, blood flow, and tissue perfusion. The radiopharmaceutical dose is injected and emits radioactivity in the form of positrons, which are detected and transformed into a visual display by computer.

A broad spectrum of radiopharmaceuticals is currently used in PET imaging. A main advantage of PET derives from the positron-emitting isotopes themselves: carbon-11 (\(^{11}\)C), nitrogen-13 (\(^{13}\)N), and oxygen-15 (\(^{15}\)O), which are present in organic molecules, and fluorine-18 (\(^{18}\)F), which can be substituted for hydrogen. Typically, radionuclides used in PET imaging have very short half-lives (2 minutes to 2 hours).

\(^{18}\)F is used for several purposes. Its half-life is long enough to trace biochemical reactions. It can be used to label a glucose compound, permitting imaging of a variety of tissues. \(^{18}\)F is administered primarily in a glucose form called fluorodeoxyglucose (FDG). FDG is highly sensitive. Neoplastic cells are hypermetabolic and appear to have an FDG affinity that results in high contrast. FDG has greater than 90% specificity for myocardial viability, neoplastic processes, and infection. FDG is an outstanding tracer that can be used in many areas of the body. It is a glucose analogue and has a broad application because every cell uses glucose as fuel.

Indications for PET Clinical PET is a useful diagnostic tool aiding diagnosis of many disease states, primarily in oncology, neurology, and cardiology. However, the technique is applicable to all parts of the body for diagnosis, disease staging, and monitoring of therapy. Unlike MRI or CT, PET provides physiologic, anatomic, and biochemical data. Although PET is more sensitive than gamma SPECT, it is considerably more expensive. The use of FDG imaging with
specially equipped gamma cameras has been an alternative to exclusive PET imaging systems. The patient preparation for nuclear medicine use of FDG in gamma SPECT imaging is similar to that for PET imaging of FDG. Because of the physics of 18F, only multiheaded cameras can be used for gamma SPECT acquisitions. Currently, there are certain limitations with gamma SPECT imaging when compared with true PET imaging. In oncology, FDG-PET has proved useful in several areas, including the diagnosis of pulmonary nodules, the differentiation of pancreatic cancer from mass-forming pancreatitis, and the diagnosis of breast cancer in selected cases of mammography and biopsy failure. PET imaging is used for the initial preoperative staging of cancer involving the lung, liver, colon, breast, head, and neck as well as in melanomas and lymphomas. For example, in lung cancer, PET is useful in determining the degree of operability. With extensive metastasis in the mediastinum, surgery is contraindicated. Staging, detection of recurrence, and response to therapy also can be determined. In cardiology, PET has demonstrated excellent utility for measuring myocardial blood flow and perfusion and for detecting coronary artery disease. The high-energy photons of PET tracers produce high-quality images even in obese patients. In these cases, PET can provide important information for determining which patients will benefit from the more invasive procedures. In neurology, FDG-PET imaging is a noninvasive aid in predicting prognosis and for surgical planning in epilepsy. By revealing areas of increased and decreased glucose utilization, PET helps surgeons pinpoint the surgical site. PET is being used to diagnose a wide variety of dementias, including Alzheimer's disease, which shows a distinct pattern of glucose consumption in the temporal and parietal regions of the brain. Also, distinct brain patterns can be seen in the involuntary movement disorders, such as Parkinson's disease, Huntington's disease, and Tourette's syndrome.

Reference Values

**Normal** Normal patterns of tissue metabolism based on oxygen, glucose, and fatty acid utilization and protein synthesis

**Procedure**

1. Remember that the actual imaging time required for a single scan is 1 to 2 hours. The actual time involved with the patient may be several hours and occurs before and during radiopharmaceutical injection. Delayed imaging may produce different results than early imaging after injection (after 45 minutes for body tumor and 30 minutes for brain tumor).
2. Position the patient on a table, then within the scanner. Before administration of the radiopharmaceutical, perform a background transmission scan. In certain procedures, this preliminary scan is optional. A number of positions are assumed, 2–6 minutes at each position.
3. Administer the radioactive drug intravenously. The patient waits 30 to 45 minutes in the department, usually remaining on the table, and then the area of interest is scanned.
4. Be aware that patients undergoing PET procedures for colon cancer, suspected pelvic pathology, or kidney studies may require a urinary catheter.
5. Remember that cardiac patients do not require fasting, and glucose monitoring may be part of the patient preparation before the scan. Elevated glucose results in decreased FDG uptake in cancer cells. Hydrate patient before and after FDG injection to minimize bladder uptake.
6. Combined PET and CT scans result in more sensitive, improved images.
7. See Chapter 1 guidelines for safe, effective, informed intratest care.

**Interventions** Patient preparation for FDG-PET imaging varies among institutions. However, some generalizations can be made.

**Pretest Patient Care**

1. Explain test purpose and procedure. Fasting is required for all tests (except cardiac). Sometimes, fasting blood glucose levels are obtained. If blood glucose levels are too high, insulin may be ordered and administered by the physician. Caution must be taken if insulin is given because it suppresses glucose tissue uptake. Insulin also suppresses FDG tissue uptake, which affects the quality of the resulting scan.
2. Administer the FDG radiopharmaceutical intravenously. Blood pressure is monitored.

**Posttest Patient Aftercare**

1. Interpret test outcome and monitor appropriately for side effects.
2. Refer to standard nuclear scan posttest precautions on page 655.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Brain Imaging**

**Clinical Implications**

1. **Epilepsy.** Focal areas with increased metabolism have been seen during actual episodes of epilepsy, with decreased oxygen utilization and blood flow during interictal episodes. (PET becomes an alternative to depth electrode implants.)
2. **Stroke.** An extremely complex pathophysiologic picture is being revealed, including anaerobic glycolysis, depressed oxygen utilization, and decreased blood flow.
3. **Coronary artery disease.** Excellent images of decreased myocardial blood flow and perfusion are observed.
4. **Dementia.** Decreased glucose consumption (hypometabolic activity) is revealed by PET imaging. PET is used to differentiate Alzheimer's disease from other types of dementia, such as Huntington's disease and Parkinson's disease.
5. **Schizophrenia.** Some studies using labeled glucose indicate reduced metabolic activity in the frontal region. The PET scans can also distinguish the developmental stages of cranial tumors and give information about the operability of such tumors.
6. **Brain tumors.** Data have been collected concerning oxygen use and blood flow relations for these tumors. Gliomas have relatively good perfusion compared with their decreased oxygen utilization. The high uptake of
radiopharmaceutical in gliomas is reported to correlate with the tumor's histologic grade.

**Interfering Factors** Excessive anxiety can alter the test results when brain function is being tested. Tranquilizers cannot be given before the test because they alter glucose metabolism.

**Interventions**

**Pretest Patient Care**

1. Instruct the patient about the purpose, procedure, and special requirements of the PET scan (see page 700). Refer to [standard nuclear scan pretest precautions](#) on page 655.
2. Advise the patient that lying as still as possible during the scan is necessary. However, the patient is not to fall asleep nor count to pass the time.
3. Remember that during the scan, it is important to maintain a quiet environment.
4. See Chapter 1 guidelines for [safe, effective, informed pretest care](#).

**Posttest Patient Aftercare**

1. Interpret test outcome and monitor appropriately for side effects.
2. Refer to standard nuclear scan [posttest precautions](#) on page 655.
3. Follow Chapter 1 guidelines for [safe, effective, informed posttest care](#).

**Cardiac Imaging**

**Clinical Implications**

1. In cardiology, PET imaging provides measurements of blood flow, myocardial perfusion, and myocardial viability. These measurements are used to detect:
   a. Coronary artery disease, which is characterized by areas of decreased blood flow, decreased perfusion, or both.
   b. Transient ischemia (both stress and rest images are performed).
2. A high rate of glucose consumption is required to meet the energy needs of the heart. Low glucose metabolism in areas of decreased blood flow indicates nonviable myocardial tissue.

**Interventions**

**Pretest Patient Care**

1. Instruct the patient about the purpose, procedure, and special requirements of the PET scan (see page 700). Refer to [standard nuclear scan pretest precautions](#) on page 655.
2. Be aware that an intravenous line may be necessary. Cardiac patients do not require fasting and may be given glucose as part of patient preparation. Smoking and medication restrictions may be required before imaging. Consult with the referring physician or the nuclear imaging department.
3. It may be necessary to place ECG leads on the patient.
4. See Chapter 1 guidelines for [safe, effective, informed pretest care](#).

**Posttest Patient Aftercare**

1. Interpret test outcome and monitor appropriately for side effects.
2. Refer to standard nuclear scan [posttest precautions](#) on page 655.
3. Follow Chapter 1 guidelines for [safe, effective, informed posttest care](#).

**Tumor Imaging**

**Clinical Implications**

1. Measurements of glucose (FDG) metabolism are used to determine tumor growth. Because small amounts of FDG can be visualized, early tumor detection is possible before structural changes detectable by MRI or CT occur. Tumor grading can be assessed by the rate of increase in glucose metabolism. In cases of suspected tumor recurrence after therapy, PET differentiates any new growth from necrotic tissue.
2. PET is used to distinguish between recurrent, active tumor growth and necrotic masses in soft tissue; this differentiation is difficult to make by MRI or CT.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose, procedure, and special requirements of the PET scan (see page 700). Refer to [standard nuclear scan pretest precautions](#) on page 655.
2. Usually, no special preparation is needed. Sometimes, a urinary catheter may have to be inserted for colon or kidney tumor detection.
3. See Chapter 1 guidelines for [safe, effective, informed pretest care](#).

**Posttest Patient Aftercare**

1. Interpret test outcome and monitor appropriately for side effects.
2. Refer to standard nuclear scan [posttest precautions](#) on page 655.
3. Follow Chapter 1 guidelines for [safe, effective, informed posttest care](#).
BIBLIOGRAPHY


DePuey EG, Garcia EV, Berman DS: Cardiac SPECT Imaging, 2nd ed. Philadelphia, Lippincott Williams & Wilkins, 2001

Early PJ, Sodee DB: Principles and Practice of Nuclear Medicine, 2nd ed. St Louis, Mosby, 1995


OVERVIEW OF X-RAY STUDIES

USE OF CONTRAST AGENTS

ADVERSE REACTIONS TO CONTRAST AGENTS

Clinical Considerations When Iodine Contrast Agents Are Used

Clinical Alert

Clinical Considerations When Barium Contrast Is Used

Clinical Alert

Clinical Alert

Clinical Alert

RISKS OF RADIATION

Safety Measures

Precautions to Be Used with Pregnant Patients

PLAIN (CONVENTIONAL) X-RAYS/RADIOGRAPHY

Chest X-Ray

Mammography (Breast X-Ray)

Abdominal X-Ray: Plain Film or KUB (Kidney, Ureters, Bladder); Scout Film; Flat Plate, Abdominal Series

Dental X-Rays

CONTRAST X-RAYS/RADIOGRAPHY

Contrast X-Ray of the Stomach: Gastric X-Ray Including Upper Gastrointestinal Examination (Upper GI [UGI] Series, Barium Swallow, Esophagram)

Small Bowel X-Ray: Intestinal Radiography and Fluoroscopy

Colon X-Ray: Defecography (DEF); Barium Enema; Air Contrast Study (Evacuative Portography)

Education Alert

Bile Duct X-Ray (Cholangiography), T-Tube Cholangiogram, Operative Cholangiogram, Percutaneous Transhepatic Cholangiogram

Intravenous Urography (IVU); Excretory Urography or Intravenous Pyelography (IVP)

NOTE

Other Tests Used to Examine the Urinary System

Retrograde Pyelography and Other Tests to Examine the Urinary System

Urography (Urine X-Ray)

Myelography, Myelogram, Spinal Cord X-Ray

Hysterosalpingography (Uterine and Fallopian Tube X-Rays)

Angiography (Digital Subtraction Angiography [DSA], Transvenous Digital Subtraction, Vascular X-Ray)

Lymphangiography (X-Rays of Lymph Nodes and Vessels)

COMPUTED TOMOGRAPHY (CT)

Computed Tomography (CT) of the Head and Neck; Brain, Eyes, and Sinus Computerized Axial Tomography (CAT)

Computed Tomography (CT) of the Body, Computerized Axial Tomography (CAT); Body Scan, Chest, Spine, Extremities, Abdomen, and Pelvis Computed Tomography

CT Screening

BIBLIOGRAPHY

INTERNET SITES

OVERVIEW OF X-RAY STUDIES

X-ray studies, also known as radiographs or roentgenograms, are used to examine soft and bony tissues of the body. X-rays are short-wavelength electromagnetic vibrations produced when fast-moving electrons collide with substances in their pathways. X-rays travel in straight lines at the speed of light (186,000 miles/second). When an x-ray beam passes through matter, some of its intensity is absorbed; the more dense the matter, the greater the degree of x-ray absorption. The composite image produced represents these varying degrees of tissue density in shades of black, white, and gray. Images may be captured on photographic film, displayed on a video screen, or recorded on digital media. The basic principle of radiography is that differences in density among various body structures produce images of varying light or dark intensity, much like the negative print of a photograph. Dense structures appear white, whereas air-filled areas are black.

USE OF CONTRAST AGENTS

Many radiographic techniques use the natural contrasts and varying densities that exist in body tissues representing air, water (in soft tissue), fat, and bone. The lungs and gastrointestinal (GI) tract normally contain air or gases. Other body structures are encased in a fatty envelope. Bone contains naturally occurring mineral salts. However, diagnosis of certain pathologic conditions requires visualization of details that cannot be revealed through plain x-rays. In these cases, details can be highlighted by the presence of contrast media in the area. These contrast substances can be administered through oral, rectal, or injection administration.

The ideal contrast agent should be relatively harmless (low toxicity, nonantigenic, nonallergenic, and inert), should not interfere with any physiologic functions, and should allow high and repeated dosing at a moderate cost. A contrast medium may be classified as either radiopaque (not permitting the transmission of x-rays) or radiolucent (permitting partial transmission of x-rays). The adverse pharmacodynamics of contrast media causes death in an estimated 1 of every 20,000 to 40,000 administrations. The most commonly used contrast agents are water-soluble iodine agents for GI examinations and intravascular procedures. Ultimately, one must always be alert to the possibility of an adverse reaction to contrast media. Consequently, emergency supplies and equipment should be readily available when using these agents.
The following contrast agents are used routinely in x-ray studies:

1. Alimentary canal contrast agents
   a. Water-soluble agents (eg, Gastrografin, Gastroview, oral Hypaque)
   b. Water-insoluble agents (eg, BaSO$_4$, Polibar Plus, Esophotrast, Anatrast)
   c. Gases (CO$_2$ gas, gas-producing calcium citrate and magnesium citrate)

2. Injectable contrast agents
   a. Nonionic iodinated contrast (low osmolar agents, eg, Omnipaque, Isovue, Optiray)
   b. Ionic iodinated contrast (high osmolar agents, eg, Renavest, Hypaque, Conray)

3. Specific-use agents
   a. Bile agents (Cholebin, Bilivist)
   b. Iodized oil contrast agents

ADVERSE REACTIONS TO CONTRAST AGENTS

All contrast agents have the potential for causing allergic reactions that can range from mild (eg, nausea and vomiting) to severe anaphylaxis (eg, cardiovascular collapse and central nervous system depression leading to death if untreated). Table 10.1 lists the range of possible adverse reactions to iodine contrast media. Reactions happen quickly and usually occur within minutes of administration of the contrast agent. Such reactions can occur in anyone.

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<th>Cutaneous</th>
<th>Gastrointestinal</th>
<th>Neurologic</th>
<th>Genitourinary</th>
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<td>Erythema</td>
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<td>Anxiety</td>
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</tr>
<tr>
<td>Acute pulmonary edema</td>
<td>Cyanosis</td>
<td>Angioneurotic edema</td>
<td>Disorientation</td>
<td>Stupor</td>
<td>Urinary cramps</td>
</tr>
<tr>
<td>Shock</td>
<td>Laryngeal edema</td>
<td>Swelling of eyes</td>
<td>Disorientation</td>
<td>Coma</td>
<td>Angioneurotic edema</td>
</tr>
<tr>
<td>Congestive heart failure</td>
<td>Apnea</td>
<td></td>
<td></td>
<td>Coma</td>
<td></td>
</tr>
<tr>
<td>Cardiac arrest</td>
<td>Respiratory arrest</td>
<td></td>
<td></td>
<td>Convulsions</td>
<td></td>
</tr>
<tr>
<td>Dyspnea</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

All Iodine Contrast Reactions

| Minor reactions requiring no treatment: sensation of heat, nausea, vomiting, local urticaria, rash, dizziness, light-headedness, transient arrhythmia, pain at injection site, mild pallor, pruritus, facial swelling 1:20 (5)
| Intermediate reactions that require treatment but no hospitalization and are not life-threatening: vomiting, extensive urticaria, bronchospasm, faintness, dyspnea, mild chest pain, headache, chills, and fever 1:100 (1)
| Severe reactions that require hospitalization and are life-threatening: syncope, laryngeal and pulmonary edema, hypotension, convulsions, circulatory collapse, pulmonary edema, severe angina, myocardial infarction, cardiac arrhythmia, coma, respiratory arrest Cardiac arrest 1:2000 (0.05)
| Death 1:40,000 (0.0025) |

Clinical Considerations When Iodine Contrast Agents Are Used

1. Know the patient’s age and health status. Children and elderly people, especially those with medical problems, may be especially sensitive to contrast agents. This sensitivity may increase the chance for side effects.
2. The presence of other medical problems may increase the risk for side effects.
   a. Those with asthma or hay fever are at a greater risk for having an allergic reaction to the contrast agent.
   b. Those with diabetes have a greater risk for developing kidney problems.
   c. Those with severe hypertension may experience a dangerous rise in blood pressure and tachycardia.
   d. Those with kidney and liver disease may experience exacerbation of their disease.
   e. Those with multiple myeloma may develop severe kidney problems.
   f. Those with overactive thyroid may experience a sudden increase in symptoms or thyroid storm.
   g. Those with sickle cell disease may experience the formation of abnormal blood cells.
   h. Those using beta blockers may have a higher risk for developing anaphylactoid reactions.
   i. Those with chronic obstructive pulmonary disease (COPD) have an increased risk for postinjection dyspnea.
3. Patients who are allergic to iodine contrast media must have this information documented in their health care records. The risk for subsequent reactions increases three to four times after the first reaction; however, subsequent reactions will not necessarily be more severe than the first. The patient must be made aware of the implications of the situation. Assess for and document allergies to iodine-containing substances (eg, seafood, cabbage, kale, raw leafy vegetables, turnips, iodized salt). Also determine each person’s reactions to penicillin or to skin test for allergies because these patients have a greater chance of having a reaction.

4. Check the patient’s fasting status before the x-ray procedure has begun. Except in an extreme emergency, iodine contrast media should never be administered intravenously sooner than 90 minutes after the patient has eaten. In most instances, the patient should fast the night before undergoing any x-ray procedure using an iodine contrast agent.

5. Death from an allergic reaction can occur if severe symptoms go untreated. Staff in attendance must be qualified to administer cardiopulmonary resuscitation should it be necessary. Emergency equipment and supplies must be readily available.

6. Promptly administer antihistamines per physician’s order if mild to moderate reactions to iodine contrast substances occur (see Table 10.1).

7. When coordinating x-ray testing with a contrast agent, keep in mind that studies using iodine and those using barium should be scheduled at different times.

8. Some physiologic change can be expected when an iodine contrast substance is injected, as during an intravenous pyelogram (IVP). Physiologic responses to iodine given intravenously include hypotension, tachycardia, and arrhythmias. For this reason, always check blood pressure, pulse, and respiration before and after these tests are performed.

9. If appropriate for the patient, encourage intake of large amounts of oral fluids after the test to promote frequent urination. This flushes the iodine out of the body.

10. Possible contraindications to the administration of iodine contrast substances include the following conditions:
   a. Hypersensitivity to iodine
   b. Sickle cell anemia (use may increase sickling effect)
   c. Syphilis (use may lead to nephrotic syndrome)
   d. Long-term steroid therapy (iodine substances may render part of the drug inactive)
   e. Pheochromocytoma (iodine substances may produce sudden, potentially fatal rise in blood pressure)
   f. Hyperthyroidism
   g. COPD
   h. Multiple myeloma
   i. Acute asthma
   j. History of renal failure
   k. Pregnancy
   l. Diabetes mellitus
   m. Severe dehydration
   n. Congestive heart failure
   o. Drug therapy known to be nephrotoxic (eg, cisplatin)

11. Nonionic contrast agents tend to produce fewer side effects than do ionic materials.

12. Patients with renal failure may develop acidosis when iodine contrast is administered.
1. **Clinical Alert**
   
   Careful patient preparation considers patient safety, prevents complications, and can prevent repeat procedures. Assess for the following risk factors associated with a higher incidence of undesirable contrast agent reactions:
   
   - Allergy
   - Asthma
   - Previous reactions to contrast media
   - Repeat and high dosages administered
   - Diabetes mellitus
   - Renal failure (preexisting); many laboratories require preprocedural assessment of creatinine levels in older adults
   - Liver insufficiency
   - Multiple myeloma
   - Dehydration
   - Older adult (>65 yr)
   - Newborns
   - History of seizures
   - Pheochromocytoma

2. **Clinical Alert**
   
   No contrast agent is without risk for causing reactions. Benefit versus risk must be considered. For example, in a work-up to detect cancer, the benefits of early detection far outweigh the dangers of cumulative x-irradiation exposure. The patient must be informed of the risk-to-benefit ratio; the patient has a legal right to this knowledge. In instances in which contrast must be delivered to high-risk patients, prophylactic premedication with prednisone may be ordered. Consult the radiology department for further information.

3. **Clinical Alert**
   
   Never inject iodized oils or barium into the bloodstream.

4. **Clinical Alert**
   
   Contrast agent–induced acute renal insufficiency is a rare and dangerous complication that occurs 1 to 5 days following intravenous injection of a contrast medium. Dehydrated patients and those with serum creatinine levels >1.4 mg/dL (>123.8 µmol/L) are at greatest risk.

5. **Clinical Alert**
   
   Intravascular iodinated contrast may interact with certain IV medications. These interactions produce insoluble precipitates that may lead to embolism. For that reason, existing IV lines should be flushed with saline before using this line as the mechanism for delivering contrast.

6. **Clinical Alert**
   
   Special attention is necessary for diabetic patients because of their increased potential for renal failure and development of lactic acidosis. Diabetic persons taking oral hypoglycemic Glucophage/metformin should have this drug withheld the day of and 48 hours following the injection of iodinated contrast. In addition, advise the patient that his or her serum creatinine level be rechecked 24 to 48 hours after he or she has received parenteral contrast. Examinations requiring extremely small volumes of contrast (myelography, arthrography) may not require such stringent precautions. Check with the radiology department for specific instructions.

7. **Clinical Alert**
   
   Tests for thyroid function (serum tests as well as nuclear medicine studies) are adversely affected for several weeks to months following iodinated contrast injection.

8. **Clinical Alert**
   
   Late reactions (2–3 days after procedure) most often occur with the use of agents such as iotrolan and iodoxane for intravascular procedures such as angiography.

---

### Clinical Considerations When Barium Contrast Is Used

There is always some risk when introducing barium sulfate or a similar contrast agent into the GI tract.

1. Barium radiography may interfere with many other abdominal examinations. A number of studies, including other x-rays, tests using iodine, ultrasound procedures, radiisotope studies, tomograms, computed tomography (CT), and proctoscopy, must be scheduled before or several days following barium studies. Consult with the radiology department for the proper sequencing of studies.

2. Emphasize that a laxative should be taken after a barium sulfate procedure is completed. Increased consumption of fluids will help to clear the bowel of barium.

3. Elderly, inactive persons should be checked for stool impaction if they fail to defecate within reasonable length of time after a barium procedure. The first sign of impaction in an elderly person is fainting.

4. Observe and record findings regarding stool color and consistency for at least 2 days to determine whether barium has been evacuated. Stools will be light in color until all barium has been expelled. Outpatients should be given a written reminder to inspect their stools for at least 2 days following barium administration.

5. If possible, avoid giving narcotics, especially codeine, when barium x-rays are ordered because these drugs can cause decreased bowel motility that can compound possible barium-associated constipation.

---

### Clinical Alert

1. Rare instances of severe allergic reactions to barium sulfate have been reported. All patients should be questioned regarding their allergic history before administration of any type of contrast agent. A history of hay fever, asthma, and other allergies places the patient at higher risk for reactions to all types of contrast agents.

2. The risk for postprocedure constipation or blockage of the bowel is increased in patients with the following conditions:
   - Cystic fibrosis
   - Dehydration
   - Acute ulcerative colitis

3. Barium should **not** be used for intestinal study in the following circumstances:
   - When a bowel perforation is suspected
   - Following sigmoidoscopy or colonoscopy, especially if a biopsy was performed, because leakage of barium from the alimentary canal can cause peritonitis. Iodinated contrast should be used in these cases.
There are special clinical considerations for ostomy patients undergoing bowel preparation for GI studies; exam preparation and procedure should be tailored by the primary care provider and the radiology department to achieve the most optimal outcomes. In most cases, standard dietary and medication restrictions apply, but modifications involving mechanical bowel cleansing with enemas and physiologic cleansing with laxatives may be necessary.

**Clinical Alert for Patients with Ostomies**

1. Enemas and laxatives should not be given to a person with an ileostomy in preparation for x-rays or endoscopy (Chap. 12) because this puts the person at risk for dehydration and electrolyte imbalance. Conversely, a person with a sigmoid colostomy requires enemas before x-ray studies or endoscopy is performed. Consequently, it is important to identify the type of surgical procedure the patient has undergone. Moreover, not all colostomies need irrigation. For example, a person with an ascending right-sided colostomy will usually pass a liquid, pasty stool high in water content and digestive enzymes; such a patient may only require laxatives.

2. Notify the radiology department that the person has an ostomy.

3. Advise all patients to bring extra ostomy supplies and pouches for use after the procedure is completed.

**RISKS OF RADIATION**

Exposure of the human body to radiation carries certain risks. The biologic effects of ionizing radiation change the chemical makeup of cells, causing cell damage and mutation and promoting carcinogenesis. However, not all forms of radiation are equal in the potential for causing damage, and often no perceptible or long-lasting damage occurs. Generally speaking, the higher the dose (as determined by the "strength" of the radiation and the duration of the exposure), the greater the risk.

Deterministic effects (ie, early effects), such as erythema, acute radiation syndrome, and induced fertility, occur after the person has received massive doses of radiation. Stochastic or late effects of radiation (ie, those in which the risk for damage rises with increasing exposure levels and consequently are of most concern in diagnostic radiology) include radiocarcinogenesis and genetic effects. Because the most radiosensitive human is the embryo during the first trimester of pregnancy, special precautions must be taken to prevent or minimize radiation exposure to the pregnant uterus (Table 10.2, Table 10.3, Table 10.4 and Table 10.5).

**Table 10.2 Principal Early Effects of Radiation Exposure on Humans and Approximate Minimum Radiation Dose Necessary to Produce Them**

<table>
<thead>
<tr>
<th>Effect</th>
<th>Anatomic Site</th>
<th>Minimum Dose (Gray)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Death</td>
<td>Whole body</td>
<td>1</td>
</tr>
<tr>
<td>Hematologic depression</td>
<td>Whole body</td>
<td>0.25</td>
</tr>
<tr>
<td>Skin erythema</td>
<td>Small field</td>
<td>3</td>
</tr>
<tr>
<td>Epilation</td>
<td>Small field</td>
<td>3</td>
</tr>
<tr>
<td>Chromosome aberration</td>
<td>Whole body</td>
<td>0.05</td>
</tr>
<tr>
<td>Gonadal dysfunction</td>
<td>Local tissue</td>
<td>0.1</td>
</tr>
</tbody>
</table>


**Table 10.3 Relative Risk for Childhood Leukemia After Irradiation in Utero by Trimester**

<table>
<thead>
<tr>
<th>Time of X-Ray Examination</th>
<th>Relative Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>First trimester</td>
<td>8.3</td>
</tr>
<tr>
<td>Second trimester</td>
<td>1.5</td>
</tr>
<tr>
<td>Third trimester</td>
<td>1.4</td>
</tr>
<tr>
<td>Total</td>
<td>1.5</td>
</tr>
</tbody>
</table>


**Table 10.4 Summary of Effects After 10-rad in Exposure in Utero**

<table>
<thead>
<tr>
<th>Time of Exposure Type of Response</th>
<th>Natural Occurrence</th>
<th>Radiation Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–2 wk</td>
<td>25%</td>
<td>0.1%</td>
</tr>
<tr>
<td>2–10 wk</td>
<td>5%</td>
<td>1%</td>
</tr>
<tr>
<td>2–15 wk</td>
<td>6%</td>
<td>0.5%</td>
</tr>
<tr>
<td>0–9 mo</td>
<td>8/10,000</td>
<td>12/10,000</td>
</tr>
<tr>
<td>0–9 mo</td>
<td>Impaired growth and development 1%</td>
<td>Nil</td>
</tr>
<tr>
<td>0–9 mo</td>
<td>Genetic mutations 10%</td>
<td>Nil</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Examination</th>
<th>Technique (kVp/mAs)</th>
<th>Entrance Skin Exposure (mrad)</th>
<th>Mean Marrow Dose (mrad)</th>
<th>Gonad Dose (mrad)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skull</td>
<td>76/50</td>
<td>200</td>
<td>10</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Chest</td>
<td>110/3</td>
<td>10</td>
<td>2</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Cervical spine</td>
<td>70/40</td>
<td>150</td>
<td>10</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Lumbar spine</td>
<td>72/60</td>
<td>300</td>
<td>60</td>
<td>225</td>
</tr>
<tr>
<td>Abdomen</td>
<td>74/60</td>
<td>400</td>
<td>30</td>
<td>125</td>
</tr>
<tr>
<td>Pelvis</td>
<td>70/50</td>
<td>150</td>
<td>20</td>
<td>150</td>
</tr>
<tr>
<td>Extremity</td>
<td>60/5</td>
<td>50</td>
<td>2</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Head CT</td>
<td>125/300</td>
<td>3000</td>
<td>20</td>
<td>50</td>
</tr>
<tr>
<td>Pelvis CT</td>
<td>124/400</td>
<td>4000</td>
<td>100</td>
<td>3000</td>
</tr>
</tbody>
</table>


Safety Measures

Certain precautions must be taken to protect patients, visitors, and staff from unnecessary exposure to radiation.

General Precautions

1. The patient’s medical records should be reviewed for radiation therapy history and to minimize the potential for unwarranted repeat studies.
2. Fast film and high-resolution screens produce quality results. Filmless “computed radiography” may reduce radiation exposure and retakes.
3. The size or area irradiated must be carefully adjusted so that no more tissue than necessary is exposed to the x-irradiation. Collimators (shutters), cones, or lead diaphragms can assure proper sizing and x-ray exposure area.
4. Fluoroscopy yields a higher dose than static radiographs or CT studies. Significant dose reduction is achieved by employing pulsed digital fluoroscopy.
5. The gonads should be shielded in both female and male patients of childbearing age unless the examination involves the abdomen or gonad areas.
6. The primary x-ray beam should pass through layers of aluminum adequate to filter out excess radiation while still providing detailed images.
7. Staff in the radiology department should wear lead aprons (and gloves if indicated) when not within a shielded booth during x-ray exposures. Patients should be shielded appropriately insofar as the procedure allows.
8. The x-ray tube housing should be checked periodically to detect radiation leakage and to indicate when repairs or adjustments are necessary.

Precautions to Be Used with Pregnant Patients

1. Women of childbearing age who could possibly be in the first trimester of pregnancy should not have x-ray examinations involving the trunk or pelvic regions. A brief menstrual history should be obtained to determine whether a possible pregnancy exists. If pregnancy is possible, a pregnancy test should be done before proceeding with x-ray examination.
2. All pregnant patients, regardless of trimester, should avoid radiographic, fluoroscopic, and serial film studies of the pelvic region, lumbar spine, and abdomen if at all possible.
3. Should x-ray studies be necessary for obstetric regions, repeat films should be avoided.
4. If x-ray studies of nonreproductive tissues are necessary (eg, dental x-rays), the abdominal and pelvic region should be shielded with a lead apron.

Responsibilities in Ordering, Scheduling, and Sequencing X-Ray Examinations

Correct and complete information should be entered into the computer or on the x-ray requisition. An appropriate order will include the name of the exam, the ordering physician’s name, and the clinical indication for the exam. Explain to the patient the purpose and procedure of the x-ray examination. Written patient instructions may be helpful. When a complete genitourinary-gastrointestinal (GI/GU) workup is scheduled, the sequence of x-ray procedures should follow a definite order:

1. First day: IVP and barium enema
2. Second day (or subsequent day): upper GI [UGI] series

Barium studies should be scheduled after the following procedures:

1. Abdominal or pelvic ultrasound examination
2. Lumbar-sacral spine x-rays
3. Pelvic x-rays
4. Hysterosalpingogram
5. IVP

As a general rule, examinations that do not require contrast should precede examinations that do require contrast. All examinations that require iodine contrast should be completed before those that require barium contrast. In addition,
examinations that require iodine contrast must precede nuclear medicine examinations that require radioactive iodine
administrations (eg, thyroid scans). Other x-ray examinations that do not require preparation can be performed at any
time. Such examinations include the following:

1. X-rays of the head, spine, and extremities
2. Noncontrast abdominal x-rays (eg, kidney, ureters, bladder [KUB], abdomen series)
3. Mammograms

Clinical Alert for Nursing Home Patients

All nursing home patients should be accompanied by another adult to the x-ray testing site. If a nonfasting patient will
be in the x-ray department over lunchtime, the facility should send a bag lunch or money for lunch with the patient.

PLAIN (CONVENTIONAL) X-RAYS/RADIOGRAPHY

Chest X-Ray

The chest x-ray is the most frequently requested radiograph. It is used to diagnose cancer, tuberculosis and other
pulmonary diseases, and disorders of the mediastinum and bony thorax. The chest x-ray provides a record of the
sequential progress or development of a disease. It can also provide valuable information about the condition of the
heart, lungs, GI tract, and thyroid gland. A chest x-ray must be done after the insertion of chest tubes or subclavian
catheters to determine their anatomic position as well as to detect possible pneumothorax related to the insertion
procedure. A postbronchoscopy chest x-ray is done to ensure there is no pneumothorax following a biopsy. In addition,
the position of other devices such as nasogastric or enteric feeding tubes can be determined and adjusted if necessary.

Reference Values

**Normal** Normal-appearing and normally positioned chest, bony thorax (all bones present, aligned, symmetrical, and
normally shaped), soft tissues, mediastinum, lungs, pleura, heart, and aortic arch

Procedure

1. Remember that routine chest radiography consists of two images: a frontal view (posteroanterior [PA]) and a left
lateral view. Upright chest films are preferred and are of utmost importance because films taken in the supine
position do not demonstrate fluid levels. This observation is especially important when testing patients on bed rest.
2. Street clothing that is covering the chest is removed to the waist. Allow only cloth or paper hospital gowns free of
buttons and snaps to be worn during the x-ray. Remove jewelry on or adjacent to the chest.
3. Ensure that monitoring cables and patches do not obscure the chest area, if possible.
4. Instruct the patient to take a deep breath and to exhale; then to take another deep breath and to hold it while the
x-ray image is taken. After the x-ray is completed, the patient may breathe normally.
5. Be aware that the procedure takes only a few minutes.
6. Follow guidelines in Chapter 1 regarding [safe, effective, informed intratest care](#).

Clinical Implications

1. Abnormal chest x-ray results indicate the following lung conditions:
   a. Presence of foreign bodies
   b. Aplasia
   c. Hypoplasia
   d. Cysts
   e. Lobar pneumonia
   f. Bronchopneumonia
   g. Aspiration pneumonia
   h. Pulmonary brucellosis
   i. Viral pneumonia
   j. Lung abscess
   k. Middle lobe syndrome
   l. Pneumothorax
   m. Pleural effusion
   n. Atelectasis
   o. Pneumonitis
   p. Congenital pulmonary cysts
   q. Pulmonary tuberculosis
   r. Sarcoïdosis
   s. Pneumoconiosis (eg, asbestosis)
   t. Coccidioidomycosis
   u. Westermark's sign (indicates decreased pulmonary vascularity, sometimes thought to suggest pulmonary
   embolus)
2. Abnormal conditions of the bony thorax include the following:
   a. Scoliosis
   b. Hemivertebrae
   c. Kyphosis
   d. Trauma
   e. Bone destruction or degeneration
   f. Osteoarthritis
   g. Osteomyelitis
3. Cardiac enlargement

**Interfering Factors** An important consideration in interpreting chest radiographs is to ask whether the film was taken in full inspiration. Certain disease states do not allow the patient to inhale fully. The following conditions may alter the patient's ability to breathe properly and should be considered when evaluating radiographs:

1. Obesity
2. Severe pain
3. Congestive heart failure
4. Scarring of lung tissues

**Interventions**

**Pretest Patient Care**

1. No special preparation is required. However, the patient should be given a brief explanation of the purpose of and procedure for the test and assured that there will be no discomfort. Screen for pregnancy status of female patients. If positive, advise the radiology department.
2. Remove all jewelry and other ornamentation in the chest area before the x-ray.
3. Remind the patient of the need to remain motionless and to follow all breathing instructions during the procedure.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Clinical Alert**

A portable x-ray machine may be brought to the nursing unit if the patient cannot be transported. The nurse may need to assist x-ray personnel in positioning the patient and film. It is the x-ray technologist's responsibility to clear all unnecessary personnel from the radiation field before x-ray exposure.

**Posttest Patient Care**

1. Interpret test outcomes and monitor for pulmonary disease and chest disorders. Explain changes in therapy based on chest x-ray results (e.g., diuretics for pulmonary edema, endotracheal tube repositioning, starting or stopping mechanical ventilation, further testing to determine new chest infiltrates).
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Mammography (Breast X-Ray)**

Soft tissue mammography visualizes the breast to detect small abnormalities that could warn of cancer. Its primary use is to screen for and discover cancers that escape detection by other means such as palpation. Typically, cancers <1 cm cannot be detected by routine clinical or self-examinations. Because the average breast cancer has probably been present for some time before it reaches the clinically palpable 1-cm size, the prognosis for cure is excellent if detected in this preclinical or presymptomatic phase.

The low-energy x-ray beam used for this procedure is applied to a tightly restricted area and consequently does not produce significant radiation exposure to other areas of the body. Therefore, it is quite acceptable from a radiation safety standpoint to recommend routine screenings. Diagnosis by mammography is based on the radiographic appearance of gross anatomic structures. Benign lesions tend to push breast tissue aside as they expand, whereas malignant lesions may invade surrounding breast tissue. Although false-negative and false-positive readings can occur, mammography is highly accurate.

Most breast lumps are not malignant; many are benign cysts. For women >40 years of age, the benefits of using low-dose mammography to find early, curable cancers outweigh possible risks from radiation exposure (Table 10.6).

**Table 10.6 Likelihood of Breast Cancer**

<table>
<thead>
<tr>
<th>Age (yr)</th>
<th>Odds</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>1:19,608</td>
</tr>
<tr>
<td>30</td>
<td>1:2525</td>
</tr>
<tr>
<td>35</td>
<td>1:622</td>
</tr>
<tr>
<td>40</td>
<td>1:217</td>
</tr>
<tr>
<td>45</td>
<td>1:93</td>
</tr>
<tr>
<td>50</td>
<td>1:50</td>
</tr>
<tr>
<td>55</td>
<td>1:33</td>
</tr>
<tr>
<td>60</td>
<td>1:24</td>
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<tr>
<td>65</td>
<td>1:17</td>
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<tr>
<td>70</td>
<td>1:14</td>
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<tr>
<td>75</td>
<td>1:11</td>
</tr>
<tr>
<td>80</td>
<td>1:10</td>
</tr>
<tr>
<td>85</td>
<td>1:9</td>
</tr>
<tr>
<td>&gt;95</td>
<td>1:8</td>
</tr>
</tbody>
</table>

The American College of Radiology (ACR) accredits mammography machines, and the Food and Drug Administration (FDA) certifies mammographic facilities. To earn accreditation, mammograms must be performed by specially trained and credentialed radiographers, and the resulting images must be interpreted by radiologists who meet criteria for continuing education in mammography. Additionally, the ACR has stringent standards for equipment, image quality, and radiation dose. Health insurers, including Medicare, require mammographic services to be performed at an accredited institution. Recently, the FDA has approved certain “digital” systems; these systems record breast anatomy on a computer rather than on film.

**Indications for Mammography**

1. To detect clinically nonpalpable breast cancers in women >40 years of age, younger women at high risk, or those having a history of breast cancer
2. When signs and symptoms of breast cancer are present
   a. Skin changes (eg, “orange peel” skin associated with inflammatory type cancer)
   b. Nipple or skin retraction
   c. Nipple discharge or erosion
3. Breast pain
4. “Lumpy” breast; multiple masses or nodules
5. Pendulous breasts that are difficult to examine
6. Survey of opposite breast after mastectomy
7. Patients at risk for having breast cancer (eg, family history of breast cancer)
8. Adenocarcinoma of undetermined origin
9. Previous breast biopsy
10. Tissue samples removed from the breast may be radiographed using detailed mammography techniques.
11. Follow-up studies for questionable mammographic images

**NOTE**

The American Cancer Society recommends a baseline mammogram for all women at 40 years of age, an annual or biannual mammogram for those 40 to 49 years of age, and a yearly mammogram for those >50 years of age.

**Reference Values**

**Normal** Essentially normal breast tissue: calcification, if present, should be evenly distributed; normal ducts with gradual narrowing ductal system branches

**Procedure**

1. Mammogram
   a. Perform mammograms with the person in an upright position, preferably standing. Make accommodations for patients using wheelchairs.
   b. Expose the breast and lift onto a film holder or digital plate. Adjust the breast tissue by hand, smoothing out all skin folds and wrinkles. Lower a movable paddle onto the breast, rigorously compressing the breast tissue.
   c. Make an x-ray exposure quickly, and immediately lift the compression.
   d. Typically, take two views (craniocaudal and mediolateral) of each breast.
   e. Be aware that before or after the x-ray examination, the technologist visually observes and manually palpates the breasts.
   f. Tell patient that the complete examination takes about 30 minutes.
   g. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.
2. X-ray–guided biopsy (stereotactic technique)
   a. Administer a local anesthetic and a sedative.
   b. Have the patient lie on her abdomen, allowing her breast to protrude through an opening in a special table.
   c. Take two stereoview mammograms, allowing precise positioning of hollow-core needle.
   d. Insert the needle into the breast at precise locations using sterile lacerations. Take multiple core tissue samples because tumors have both benign and malignant areas.
   e. Cleanse the breast, and apply a sterile dressing.

**Comparison of Core Needle vs. Vacuum-Assisted Biopsy**

<table>
<thead>
<tr>
<th>Core Needle Biopsy</th>
<th>Technique</th>
<th>Disadvantages</th>
<th>Advantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABB1 Site-select Centricca</td>
<td>Automated gun</td>
<td>Requires multiple passes into tissue</td>
<td>Excellent for dense lesions</td>
</tr>
<tr>
<td>M1BB Mammatome ATEC</td>
<td>Dual lumen needle/probe with rotating cutter</td>
<td>Increased potential for postprocedural bleeding</td>
<td>Single pass into tissue yields multiple samples</td>
</tr>
<tr>
<td><strong>Vacuum-Assisted Biopsy (VAB)</strong></td>
<td></td>
<td>Expensive equipment</td>
<td>Larger tissue sample</td>
</tr>
</tbody>
</table>

3. Needle x-ray localization and surgical biopsy
   a. Administer a local anesthetic and sedative. In some instances, general anesthesia is used.
   b. Insert a needle that holds a fine wire, clip, or biodegradable marker into the breast tissue, using breast x-ray films as a guide. When the needle point is at the tip of the x-ray–defined abnormality, the device is released. It
stays there until the surgeon, guided by the wire, removes a specimen of the abnormal tissue.

**NOTE**

Rigorous compression is a brief and uncomfortable but critical step in ensuring a high-quality mammogram. It lowers dose and improves image quality.

**Clinical Alert**

1. Computer-assisted diagnosis (CAD) recently became a reimbursable procedure for Medicare patients. Computer software scans the image and notes suspicious areas that a radiologist could miss, thus acting as a second opinion.
2. Many radiologists double-read all mammograms.
3. Comparison with old mammograms is very important. Consequently, patients are advised to have all mammograms performed at the same facility or retrieve old mammograms and bring them along when having a new study performed.
4. Mammographic examination of augmented breasts requires additional views that add to procedure time. The presence of implants should be communicated to the radiology department when scheduling the procedure.

**Clinical Implications**

Abnormal mammogram findings reveal the following conditions:

1. **Breast mass**
   a. Benign breast masses (eg, cysts, fibroadenomas) are usually round and well demarcated.
   b. Malignant breast masses are often irregularly shaped with extensions into adjacent tissue, generally with an increased number of blood vessels (Fig. 10.1).

2. Calcifications present in the malignant mass (duct carcinoma) or in adjacent tissue (lobular carcinoma) are described as innumerable punctate calcifications resembling fine grains of salt or rod-like calcifications that appear thin, branching, and curvilinear. Macrocalcifications (large mineral deposits) generally represent benign degenerative processes. Microcalcifications (<1/50 inch) are of more concern and require close examination.
3. The likelihood of malignancy increases with a greater number of calcifications in a cluster. However, a cluster with as few as three calcifications, particularly if they are irregular in shape or size, can occur in cancer.

4. Typical parenchymal patterns are as follows:
   a. N1: normal
   b. P1: mild duct prominence on less than one fourth of the breast
   c. P2: marked duct prominence

*FIGURE 10.1* Half of all breast cancers develop in the upper outer section. (Source: Department of Health and Human Services, 1994.)

*FIGURE 10.2* Examples of (A) whole breast compression and (B) spot compression.
d. DY: dysplasia (some diagnosticians believe that the person who exhibits dysplasia is 22 times more likely to develop breast cancer than the person with normal results)

5. Findings of breast cancer when contrast is injected are associated with extravasation of contrast, filling defects, obstruction or irregular narrowing of ducts (Chart 10.1) (see Fig. 10.1 and Fig. 10.2).

<table>
<thead>
<tr>
<th>Chart 10.1 Clinical Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Contrast mammography (ductogram, galactogram) is a valuable aid for diagnosing intraductal papillomas. Mammary duct injection is used when cytologic examination of breast fluid or discharge is abnormal. In contrast mammography, after careful cannulation of a discharging duct, about 1 mL of a radiopaque substance (eg, 50% sodium diatrizoate) is injected into the breast duct with a blunt, 25-gauge needle. 2. Ductal lavage is a new technique in which the milk ducts are cannulized. Saline is injected and, when withdrawn, will &quot;wash out&quot; ductal cells. These cells are examined in the laboratory, in much the same way as a Pap smear is reviewed. (See Chap. 11 for additional information.)</td>
</tr>
</tbody>
</table>

Interventions
Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of mammograms. Mammography is the single best method for detecting breast cancer while it is still in a curable stage (Fig. 10.3). Some discomfort is to be expected when the breast is compressed.

2. Assess pregnancy status of female patients. If positive, advise radiology department.
3. Instruct the patient not to apply deodorant, perfume, powders, or ointment to the underarm area on the day of the examination. Residue from these preparations can obscure optimal visualization.
4. Advise the patient to wear separates rather than a dress because clothing must be removed from the upper body.
5. Suggest that patients who have painful breasts refrain from caffeinated foods and beverages (eg, coffee, tea, cola, chocolate) for 5 to 7 days before testing.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

NOTE

Patients in the reproductive age group are advised to have mammograms performed in the 2 weeks that follow their last menstrual period.

Posttest Patient Care

1. Interpret test outcomes and counsel appropriately. If a biopsy is necessary, see procedures for biopsy using x-ray technology.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Clinical Alert

1. A mammogram detects abnormalities that could warn of cancer. The actual diagnosis of cancer is made by biopsy. Only one in five biopsies test positive for cancer.
2. Several methods can be used to provide a breast tissue sample necessary for cancer diagnosis. These include core-needle biopsy, surgical biopsy, and vacuum-assisted biopsy. Any of these methods can utilize either x-ray mammography or ultrasound for image guidance.

Orthopedic X-Ray: Bones, Joints, and Supporting Structures

Orthopedic radiography examines a particular bone, group of bones, or joint. The bony or osseous system presents five functions of radiologic significance: structure support of the body, locomotion, red marrow storage, calcium storage, and protection of underlying soft tissue and organ structures. Orthopedic radiography is performed on the following structures:

1. The extremities (eg, hand, wrist, shoulder, foot, knee, hip)
2. The bony thorax (eg, ribs, sternum, clavicle)
3. The spine (eg, cervical, thoracic, lumbar, sacrum, coccyx)
4. The head and skull (eg, facial bones, mastoids, sinuses)

Optimal results from orthopedic x-ray examinations depend on proper immobilization of the area being studied. To
produce a thorough image of the body part, at least two and sometimes more projections are required. These are usually taken at angles of 90 degrees to one another (e.g., anteroposterior and lateral views).

To examine more complex structures such as the spine and skull, or to examine a structure in greater detail, several projections from various angles may be required.

Reference Values

Normal Normal osseous (bone) and supporting tissue structures

Procedure

1. Inform the patient that dietary restrictions are not necessary.
2. Have the patient assume the positions most favorable to capturing the best images. However, the degree of patient mobility and physical condition may also need to be considered. Typically, the anatomic structures being studied are examined from several angles and positions. This may require the examiner to manipulate the body area physically into a position that will allow optimal visualization.
3. Be aware that jewelry, zippers, snaps, monitoring cables, and so forth interfere with proper visualization. These objects must be removed from the visual field if possible. Skull x-rays require removal of dentures and partials.
4. Remove surgical-type hardware used to stabilize a traumatized area. This should be done only under the direction of the attending physician.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications Abnormal orthopedic x-ray results may reveal the following conditions:

1. Fractures
2. Dislocations
3. Arthritis
4. Osteoporosis
5. Osteomyelitis
6. Degenerative joint disease
7. Hydrocephalus
8. Sarcoma
9. Abscess and aseptic necrosis
10. Paget's disease
11. Gout
12. Acromegaly
13. Metastatic processes
14. Myeloma
15. Osteochondrosis, for example, a. Legg-Calvé-Perthes disease b. Osgood-Schlatter disease
16. Bone infarcts
17. Histiocytosis X
18. Bone tumors (benign and malignant)
19. Foreign bodies

Interfering Factors Radiography of the lumbosacral spine, coccyx, or pelvis must be completed before barium studies because residual barium may interfere with proper visualization. Jewelry and accessories, heavy clothing, metallic objects, zippers, buttons, snaps, cables, and other monitoring equipment and supplies can interfere with optimal views and need to be removed before the examination.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test. No preparation or dietary restrictions are necessary. Screen for pregnancy status of female patients. If positive, advise the radiology department.
2. Assure the patient that the procedure in and of itself causes no pain. However, necessary manipulation of the body may cause discomfort. If appropriate, pain medication may be administered before the procedure.
3. Advise the patient that all dentures, partials, jewelry, and other ornamentation worn in the anatomic area being examined must be removed before the study. If possible, simple clothing should be worn, and the previously mentioned items should be left at home or in the patient's room.
4. Emphasize the importance of not moving during the procedure unless specifically instructed otherwise. Movement distorts or "blurs" the image and often requires repeat exposures.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Care

1. Interpret test outcomes and monitor for fractures, dislocations, and other orthopedic disorders. Counsel about need for follow-up procedures and treatment.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.
Clinical Alert

1. Orthopedic radiography also can provide information about soft tissue structures, such as swelling or calcifications. However, radiography alone cannot provide data about the condition of cartilage, tendons, or ligaments.
2. Portable x-ray machines can be taken to the nursing unit if the patient cannot be transported to the radiology department. Nursing personnel may need to assist in the process. The x-ray technologist is responsible for clearing all unnecessary personnel from the immediate radiation field before activating the exposure.

Abdominal X-Ray: Plain Film or KUB (Kidney, Ureters, Bladder); Scout Film; Flat Plate, Abdominal Series

This radiographic study does not use contrast media. It is done to aid in the diagnosis of intraabdominal diseases such as nephrolithiasis, intestinal obstruction, soft tissue mass, or ruptured viscus. It may be the preliminary step in evaluating the GI tract, the gallbladder, or the urinary tract, and it is done before IVP or other renal studies. Abdominal films may provide information on the size, shape, and position of the liver, spleen, and kidneys.

Reference Values

Normal Normal abdominal structures

Procedure

1. Have the patient wear a hospital gown. All metallic objects must be removed from the abdominal area.
2. Have the patient lie in a supine position on the x-ray table.
3. Take multiple images (including upright and left decubitus) for an abdominal series to assess air-fluid levels.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications Abnormal abdominal x-ray results reveal the following conditions:

1. Calcium deposits in blood vessels and lymph nodes; cysts, tumors, or stones
2. Ureters are not clearly defined, although calculi may be visualized within the ureters.
3. The urinary bladder can often be identified by the shadow it casts, especially in the presence of urine with high specific gravity
4. Abnormal kidney size, shape, and position
5. Appendicolithiasis
6. Foreign bodies
7. Abnormal fluid; ascites
8. Large tumors and masses, (eg, bladder, ovarian, or uterine), if they displace normal bowel configurations
9. Abnormal gas distribution associated with bowel perforation or obstruction
10. Fusion anomalies
11. Horseshoe-shaped kidneys

Interfering Factors

1. Barium may interfere with optimal visualization. Therefore, this examination should be done before barium studies.
2. A “flat plate” of the abdomen does not detect free air.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test. Normal diet is allowed unless contraindicated. Assure the patient that the procedure in itself is not painful.
2. Remove belts, zippers, jewelry, and other ornamentation from the abdominal area.
3. Instruct the patient to remain still and to follow breathing instructions.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Clinical Alert

1. Abdominal plain films are not diagnostic for certain conditions, such as esophageal varices or bleeding peptic ulcer.
2. A portable x-ray machine may be brought to the nursing unit if the patient cannot be moved. Assist with positioning as necessary. The x-ray technologist is responsible for clearing all unnecessary personnel from the radiation field before the x-ray is taken.

Posttest Patient Care

1. Interpret test outcomes and monitor for intraabdominal disease.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Dental X-Rays

Dental x-rays screen and diagnose causes of pain and other symptoms related to the teeth, jaws, and temporomandibular joints and are also used as follow-up for dental therapy. Many different types of dental radiographs are available because of the complex tissue density found within the human masticatory system. The x-rays are categorized by the location at which the film is placed during the procedure (intraoral versus extraoral). The most common x-rays taken are the bite
wing and the periapical, both of which are intraoral. The various types of dental x-rays include the following:

1. Intraoral (film positioned inside the mouth)
   a. Bite wing: shows coronal portion of the tooth; also done for caries detection; shows bite correlation between upper and lower teeth
   b. Peripheral: shows x-ray of the whole tooth and immediate surrounding area
   c. Occlusal: shows chewing surfaces and curve of mandibular molar teeth
2. Extraoral (film positioned outside the mouth)
   a. Shows various projections of the skull, maxilla, sinuses, or temporomandibular joints
   b. Panorex (full-mouth x-ray)
   c. CT
   d. Arthrography of the temporomandibular joint

Reference Values

**Normal**
Normal mandible, maxilla, temporomandibular joints, maxillary sinuses, and primary or permanent dentition

**Procedure**

1. Have the patient sit upright and place the film and holder in the mouth for intraoral studies. The patient may bite on the holder or may anchor it with a finger to keep it in place. Drape a lead apron with a cervical collar over the patient's torso and neck area.
2. Remember that different designs of film holders facilitate proper alignment for correct x-ray tube orientation. There are also many different types of extraoral films that can be taken, each with their own procedures. For example, with the lateral skull projection, the patient sits upright, and the film packet is placed on one side of the head while the x-ray source is placed on the opposite side. In other instances, such as Panorex imaging, rotate the x-ray machine around the face.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

**Clinical Alert**

Previous extensive radiation therapy or a current state of pregnancy may present contraindications to dental x-rays. Consult the patient's physician if in doubt.

**Clinical Implications**

1. Abnormal dental x-ray results reveal the following conditions:
   a. Dentition
      1. Changes in number of teeth
      2. Changes in shape of teeth
      3. Changes in pulp canal
      4. Miscellaneous other tooth lesions
   b. Radiolucent lesions of the jaw and teeth
      1. Lesions at the tooth apex or its midline
      2. Lesions in place of a missing tooth
      3. Lesions around the crown of an impacted tooth
      4. Bubble-like radiolucencies
      5. Other multiple but different radiolucent lesions
      6. Lesions that destroy the cortical plate of the tooth
   c. Mixed lesions (radiopaque and radiolucent)
   d. Salivary gland lesions
   e. Soft tissue lesions
   f. Temporomandibular joint abnormalities

**Interfering Factors**
The following factors can interfere with proper visualization:

1. Braces and retainers
2. Partials and dentures
3. Restorations
4. Jewelry (e.g., earrings)
5. Bony growths on the inside of the mandible and the midline of the hard palate (torus) or excess deposits of bone

**Interventions**

**Pretest Patient Care**

1. Explain purpose, procedure, benefits, and risks (minimum radiation exposure). Stress the importance of holding still and breathing through the nose to lessen the gag reflex.
2. Assist the patient to rinse his or her mouth before the procedure.
3. Assess for contraindications and interfering factors.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Care**

1. Evaluate x-ray films and explain abnormalities. Comparison with a normal x-ray film may be helpful.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.
CONTRAST X-RAYS/RADIOGRAPHY

To visualize hollow internal viscera, contrast media is administered to highlight the structure. Refer to pages 706–711 for special care when using contrast media. Careful sequencing of multiple examinations is necessary. As a general rule, the following instructions for sequencing should be followed:

1. Perform abdominal pelvic plain film or CT, ultrasound, and nuclear medicine studies before contrast studies of the intestines.
2. Perform examinations of the lower intestine (barium enema) 1 or 2 days before UGI examinations.
3. Perform examinations requiring an injection of iodinated contrast, such as an IVP, before any barium studies (eg, barium enema, UGI).
4. Consult the radiology department for specific sequencing information.
5. Take special caution when administering contrast agents to diabetic persons and persons with kidney problems (see page 710).
6. Use fluoroscopy for imaging of diagnostic (moving) structures such as those of the alimentary canal. Use fluoroscopy to localize tumors for biopsy and drainages, guide catheter, or filter stent placement and monitor vascular filling for both diagnostic and therapeutic purposes (angioplasty).

Fluoroscopic radiation dose is higher than conventional x-rays. Dose is directly related to time of exposure. The use of digital fluorography tends to reduce dose by pulsing the x-ray beam.

Contrast X-Ray of the Stomach: Gastric X-Ray Including Upper Gastrointestinal Examination (Upper GI [UGI] Series, Barium Swallow, Esophagram)

Gastric radiography visualizes the form, position, mucosal folds, peristaltic activity, and motility of the stomach and upper GI tract. A UGI series includes the esophagus, duodenum, and upper portion of the jejunum.

Preliminary films without the use of a contrast medium are useful in detecting perforation, presence of radiopaque foreign substances, gastric wall thickening, and displacement of the gastric air bubble, which may indicate a mass external to the stomach.

Oral contrast substances, such as barium sulfate or diatrizoate meglumine (Gastrografin), highlight conditions such as hiatal hernia, pyloric stenosis, gastric diverticulitis, presence of undigested food, gastritis, congenital anomalies (eg, dextroposition, duplication), or diseases of the stomach (eg, gastric ulcer, cancer, stomach polyps).

Reference Values

<table>
<thead>
<tr>
<th>Normal</th>
<th>Normal stomach size, contour, motility, and peristaltic activity Normal esophagus</th>
</tr>
</thead>
</table>

NOTE

A video-esophagram is typically performed to evaluate swallowing disorders, particularly in poststroke patients, and after head and neck surgery with plastic repair. This examination generally includes evaluation by a speech pathologist.

Procedure

1. Have patient change from street clothing into a hospital gown. Neck and torso jewelry and other ornamentation must be removed.
2. Instruct the patient to swallow the barium after the patient is properly positioned in front of the fluoroscopy machine. Some changes in position may be necessary during the procedure. A motorized tabletop shifts the patient from an upright to a supine position when appropriate. Fluoroscopy allows visualization and filming of actual activity taking place in real time.
3. Take several conventional x-ray films following fluoroscopic examination. The patient will need to hold his or her breath during each exposure.
4. Tell patient that examination time may be 20 to 45 minutes.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications Abnormal UGI x-ray results reveal the following conditions:

1. Congenital anomalies
2. Gastric ulcer
3. Carcinoma of stomach
4. Gastric polyps
5. Gastritis
6. Foreign bodies
7. Gastric diverticula
8. Pyloric stenosis
9. Reflux and hiatal hernia
10. Volvulus of the stomach
Normal contours may be deformed by intrinsic tumors or consistent filling defects as well as by stenosis in conjunction with dilation.

Interfering Factors

1. If the patient is debilitated, proper examination may be difficult; it may be impossible to visualize the stomach adequately.
2. Retained food and fluids interfere with optimal film clarity.

Interventions

Pretest Patient Care

1. Explain purpose and procedure (consult barium contrast test precautions on pages 710–711). Written instructions on pretest preparation are helpful for the patient. Screen female patients for pregnancy status. If positive, inform the radiology department.
2. Inform patient that complete fasting from food and fluids is required for a minimum of 8 hours before the procedure. Necessary oral medications (other than Glucophage/metformin) may be taken with a tiny sip of water. Inform radiology department because pills may be visualized during the study.
3. Instruct the patient to hold still and follow breathing instructions during the procedure.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Clinical Alert

1. If the patient has diabetes, alert the radiology department and schedule examination for early morning. If diabetic patient is taking Glucophage/metformin, special considerations may be necessary. Consult with radiology department to determine whether this medication regimen must be suspended for the day of and several days after the study.
2. Determine whether the patient is allergic to barium. Although rare, presence of this allergy must be communicated to the radiology department so that alternate contrast can be used.
3. All female patients of reproductive age must be screened for pregnancy before performing this study.

Posttest Patient Care

1. Pretest diet and activity may be resumed. Provide food and ample fluids.
2. Administer laxatives as ordered. If barium sulfate or diatrizoate meglumine has been administered, a laxative should be taken.
3. Observe and record stools for color and consistency. Monitor evacuation of barium. Counsel that follow-up procedures may be necessary.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Small Bowel X-Ray; Intestinal Radiography and Fluoroscopy

These small intestine studies, usually scheduled in conjunction with UGI series, are done to diagnose small bowel diseases (eg, ulcerative colitis, tumors, active bleeding, obstruction). A contrast material such as barium sulfate or meglumine diatrizoate highlights Meckel's diverticulum, congenital atresia, obstruction, filling defects, regional enteritis, lymphoid hyperplasia, tuberculosis of small intestine (malabsorption syndrome), sprue, Whipple's disease, intussusception, and edema.

The mesenteric small intestine begins at the duodenojejunal valve and ends at the ileocecal valve. The mesenteric small intestine is not routinely included as part of a UGI study.

Reference Values

Normal Normal small intestine contour, position, and motility

Procedure

1. Have the patient change into a hospital gown after removing street clothes and accessories. Perform a preliminary plain-film study with the patient on the examining table.
2. Have the patient swallow the prescribed amount of contrast media while the patient is standing in front of the fluoroscopy machine.
3. Take timed films after contrast material is swallowed, usually every 30 minutes.
4. Remember that the examination is not complete until the ileocecal valve has filled with contrast material. This may take several minutes (for those patients with a bypass) to several hours.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications Abnormal small bowel x-ray results indicate the following conditions:

1. Anomalies of small intestine
2. Errors of rotation
3. Meckel's diverticulum
4. Atresia
5. Neoplasms
6. Regional enteritis (Crohn's disease)
7. Tuberculosis
8. Malabsorption syndrome
9. Intussusception
10. Roundworms (ascariasis)
11. Intraabdominal hernias

Interfering Factors

1. Delays in small intestine motility can be due to the following circumstances:
   a. Morphine use
   b. Severe or poorly controlled diabetes
2. Increases in motility in the small intestine can be due to the following circumstances:
   a. Fear or anxiety
   b. Excitement
   c. Nausea
   d. Pathogens
   e. Viruses
   f. Diet (eg, very high fiber)

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test. Refer to barium contrast test precautions (see pages 710–711). Written reminders for pretest instructions are helpful, especially for diet limitations. Screen female patients for pregnancy status. If positive, advise the radiology department.
   2. Maintain total fast from midnight until the examination is completed.
   3. Do not administer laxatives or enemas to a patient with an ileostomy.
   4. Instruct the patient regarding the need to hold still and to follow breathing instructions during the procedure.
   5. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Clinical Alert

1. If the patient has diabetes, alert the radiology department and schedule examination for early morning. If the diabetic patient is taking Glucophage/metformin, special considerations may be necessary. Consult with the radiology department to determine whether this medication regimen must be suspended during and for several days after study.
2. Determine whether the patient is hypersensitive to barium. Although rare, presence of this allergy must be communicated to the radiology department so alternate contrast can be used.
3. All female patients of reproductive age must be screened for pregnancy before performing this study.

Posttest Patient Care

1. Resume pretest diet and activity. Assist patient if necessary.
2. Administer laxatives if ordered. If a barium sulfate swallow has been done, a laxative should be taken. However, do not give laxatives to a patient with an ileostomy unless specifically ordered.
3. Monitor stools for color and consistency.
4. Counsel patient about motility disorders and other small intestine abnormalities. Follow-up procedures may be necessary.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Colon X-Ray: Defecography (DEF); Barium Enema; Air Contrast Study (Evacuative Portography)

This fluoroscopic and filmed examination of the large intestine (colon) allows visualization of the position, filling, and movement of contrast medium through the colon. It can reveal the presence or absence of diseases such as diverticulitis, mass lesions, polyps, colitis, obstruction, or active bleeding. Barium or diatrizoate meglumine (Hypaque) is instilled into the large intestine through a rectal tube inserted into the colon. The radiologist, with the aid of a fluoroscope, observes the barium as it flows through the large intestine. X-ray films are taken concurrently.

Education Alert

A pretest preparation is vital for this exam. For a satisfactory examination, the colon must be thoroughly cleansed of fecal matter. This is most important. Accurate identification of small polyps is possible only in a clean bowel. The presence of stool can also make the search for bleeding sources much more difficult.

If polyps are suspected, an air-contrast colon examination may be done. The procedure is basically the same as that for the barium enema; however, more complex radiographs need to be taken with the patient in several different positions. A double-contrast mixture of air and barium is instilled into the colon under fluoroscopic visualization.

Reference Values

Normal Normal colon position, contour, filling, movement time, and patency

Procedure

1. Have the patient lie on his or her back while a preliminary x-ray film is made; this step may be omitted at some institutions.
2. Have the patient then lie on his or her side while barium is administered by rectal enema (ie, through the rectum
and up through the sigmoid, descending, transverse, and ascending colon to the ileocecal valve).

3. Take conventional x-ray films following fluoroscopy, which includes several spot films. After these are completed, the patient is free to expel the barium. After evacuation, another film is made.

4. Be aware that defecography and evacuative portography are contrast-enhanced studies of the anus and rectum function during evacuation. Often used in young patients to evaluate rectoceles, rectal prolapse, or rectal intussusception, this examination requires the patient to evacuate into a specially designed commode while being evaluated fluoroscopically.

5. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications

1. Abnormal colon x-ray results indicate the following conditions:
   a. Lesions or tumors (benign)
   b. Obstructions
   c. Megacolon
   d. Fistulas
   e. Inflammatory changes
   f. Diverticula
   g. Chronic ulcerative colitis
   h. Stenosis
   i. Right-sided colitis
   j. Hernias
   k. Polyps
   l. Intussusception
   m. Carcinoma

2. Appendix size, position, and motility can also be evaluated; however, a diagnosis of acute or chronic appendicitis cannot be made from x-ray findings. Instead, typical signs and symptoms of appendicitis provide the most accurate data for this diagnosis.

Interfering Factors A poorly cleansed bowel is the most common interfering factor. Fecal matter interferes with accurate and complete visualization. Therefore, it is imperative that proper bowel cleansing be conscientiously carried out, or the procedure may need to be repeated.

Interventions

Pretest Patient Care Preparation involves a three-step process over a 1- to 2-day period and includes diet restrictions, physiologic cleansing of the large bowel by means of oral laxatives, and mechanical cleansing with enemas. Twelve- to 18-hour protocols are common. Follow institutional protocols.

1. Explain the purpose and procedure of the test. Patients may be apprehensive or embarrassed. Include a family member in this process if it appears likely that the patient will need assistance with preparation. Explain the need to cooperate to expedite the procedure. Emphasize that the actual time frame when the colon is full is quite brief. Screen female patients for pregnancy status. If positive, advise the radiology department.

2. A written reminder about the following may be helpful to the patient:
   a. Only a clear liquid diet should be taken before testing (according to protocols).
   b. Stool softeners, laxatives, and enemas need to be taken to ensure bowel cleanliness necessary for optimal visualization. Agents such as X-Prep, citrate of magnesia, and bisacodyl assist in emptying the ascending and right to midtransverse colon (proximal large bowel). Enemas cleanse the left transverse, descending, and sigmoid colon and the rectum. Suppositories also empty the rectum.
   c. Fasting from food and fluids is prescribed before the test. Nothing should be eaten or drunk from midnight until the test is completed. Oral medications should also be temporarily discontinued unless specifically ordered otherwise. Check with the clinician who orders the test.

3. Refer to barium contrast test precautions on pages 710–711.

4. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Care

1. Resume pretest activity and diet. Assist the patient if necessary. This bowel examination can be very exhausting. Patients may be weak, thirsty, hungry, and tired. Provide a calm, restful environment to promote return to normal status.

2. Administer laxatives for at least 2 days after these studies or until stools return to normal. Instruct the patient to assess stools during this time. Stools will be light colored until all barium has been expelled. Outpatients should be given a written reminder to inspect stools for 2 days.

3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.
Clinical Alert

1. Multiple enemas given before the procedure, especially to a person at risk for electrolyte imbalances, could induce a rather rapid hypokalemia. Enema fluid, if not expelled within a reasonable time, can be absorbed through the bowel wall and deposited into the intestinal spaces and eventually within extracellular spaces.

2. Caution should dictate administration of cathartics or enemas in the presence of acute abdominal pain, active bleeding, ulcerative colitis, or obstruction. Consult with the physician or radiology department and consider the following points:
   a. Introducing large quantities of water into the bowel of a patient with megacolon should be avoided because of the potential danger of water intoxication. In general, patients with toxic megacolon should not receive enemas.
   b. In the presence of colon obstruction, large volumes of water from enemas may be reabsorbed, and impaction may occur.
   c. Rectal obstruction makes it difficult or impossible to give cleansing enemas because the solution will not be able to enter the colon. Consult the physician or radiology department.

3. Strong cathartics administered in the presence of obstructive lesions or acute ulcerative colitis can present hazardous or life-threatening situations.

4. Be aware of complications that can occur when barium sulfate or other contrast media are introduced into the GI tract. For example, barium may aggravate acute ulcerative colitis or cause a progression from partial to complete obstruction. Barium also should not be given as contrast for intestinal studies when a bowel perforation is suspected because leakage of barium through the perforation may cause peritonitis. Iodinated contrast substances should be used if perforation is suspected.

5. Determine whether the patient is hypersensitive to barium. Although rare, the presence of this allergy must be communicated to the radiology department so that alternate contrast media can be used.

6. Fasting orders include oral medications except when specified otherwise.

7. If patient has diabetes, alert the radiology department and schedule examination for early morning. If diabetic patient is taking Glucophage/metformin, special consideration may be necessary. Consult with the radiology department to determine whether this medication regimen must be suspended the day of and several days after the study.

8. Determine whether patient is allergic to latex. Latex products are typically used to administer the contrast agent; alternate materials must be used if the patient is hypersensitive. Inform the radiology department of any known or suspected latex allergies.

9. Inform the radiology department if this procedure is to follow a sigmoidoscopy or colonoscopy, particularly if a biopsy was performed. In the case of biopsy, an iodinated contrast agent, rather than barium, is used.

Clinical Alert

Colonic Transit Time

This examination is performed on patients with suspected colonic motility disorder. The patient must not take any laxatives, enemas, or suppositories before beginning this test or during the 4–7 days it takes to perform this test. The procedure is quite simple:

1. The patient receives several pills that contain radiopaque markers (sitz markers).
2. A KUB or series of KUBs are performed at fixed times several days later.
3. The passage of or retention of these markers is noted and recorded.
4. Retention of a significant portion of markers 5 days after administration is considered abnormal and is evidence of dysmotility or an outlet obstruction.

Special Considerations

1. Children or elderly patients receiving barium enemas
   a. Because a successful examination of the large intestine depends on the ability of the bowel to retain contrast medium during visualization and filming, special techniques are used for infants and young children and the infirm or uncooperative adult patient.
   b. After inserting a small enema tip into the rectum, the infant's buttocks are gently taped together to prevent leakage of contrast material during the study.
   c. For the older patient, a special retention enema tip may by used. This device resembles a regular enema tip, but it can be inflated, much like an indwelling urinary catheter, after insertion into the rectum. When the examination is done, the retention balloon is deflated and the tip removed.

2. Barium enema in the presence of a colostomy
   a. See page 711 for assessment criteria.
   b. Laxatives can be taken.
   c. Suppositories are of no value.
   d. Follow physician's diet orders.
   e. If irrigation is necessary, a preassembled colostomy irrigation kit or a soft, no. 28, standard-tip Foley catheter attached to a disposable enema bag may be used.
   f. Advise the patient that a Foley catheter is used to introduce the barium into the stoma.
   g. The patient should bring additional colostomy supplies to the radiology department for posttest use.

3. Patients with stomas
   a. Patients with descending or sigmoid colostomies may need a normal saline or tap-water irrigation to wash out the barium.
   b. Advise those who normally irrigate their colostomy to wear a disposable pouch for several days until all the barium has passed.

Bile Duct X-Ray (Cholangiography), T-Tube Cholangiogram, Operative Cholangiogram, Percutaneous
Transhepatic Cholangiogram

A cholangiogram visualizes the bile ducts by enhancing them with an iodinated contrast agent. Often performed on the postcholecystectomy patient, the cholangiogram is used to identify intraductal mass lesions and calculi. A number of approaches may be used to opacify and image the bile ducts:

1. **T-tube cholangiogram**: Following cholecystectomy, a self-retaining T-shaped drainage tube may be surgically inserted into the common bile duct. Before removal, patency is verified by injecting iodinated contrast into the T-tube to fill the biliary tree.

2. **Cholangiogram with stone removal**: This study combines diagnostic visualization of the bile ducts with therapeutic capture and removal of ductal calculi.

3. **Intravenous cholangiography**: This study allows radiographic visualization of the large hepatic ducts and the common ducts by means of intravenous injection of a contrast medium. It is rarely performed.

4. **Operative cholangiography**: Cannulation and injection of contrast medium into the exposed cystic duct or common bile duct is performed during surgery.

5. **Percutaneous transhepatic cholangiography**: A needle or small-diameter catheter is percutaneously introduced into the liver and the bile duct. Following injection of the contrast agent, the hepatic and common ducts should be visualized. The dilated biliary tree can be shown up to the point of obstruction, which is usually in the common duct. This procedure is frequently done for jaundiced patients whose liver cells are unable to transport oral or intravenous contrast agents properly.

6. **Intravenous cholecystography**: Radiographic visualization of the gallbladder is performed after intravenous injection of a contrast agent. It is rarely performed.

7. **Oral cholecystography**: Radiographic visualization of the gallbladder is performed after oral administration of an opaque medium. This test is often combined with or replaced by gallbladder sonography.

8. **Endoscopic retrograde cholangiopancreatography (ERCP)**: This endoscopic procedure uses an injection of a contrast agent to evaluate the patency of pancreatic and common bile ducts, the duodenal papilla, and the normalcy of the gallbladder (see Chap. 12). Often, the ERCP is performed therapeutically, involving stone extraction, stent placement, or other treatments.

Reference Values

**Normal** Patent bile ducts

**Procedure for T-Tube Cholangiogram**

1. Have the patient lie on the x-ray table as an iodine contrast medium is injected into the T-tube.
2. Be aware that no pain or discomfort should be felt; however, some persons may feel pressure during the injection.
3. Unclamp the T-tube after the procedure and allow it to drain freely unless otherwise ordered. This minimizes prolonged, irritating contact of residual contrast in the bile duct.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications

Abnormal duct and gallbladder x-ray results reveal stenosis obstruction or choledocholithiasis (bile duct calculi of the common bile duct).

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the test. Assure the patient that the procedure is not painful, but some discomfort or pressure may be felt when the contrast is injected. If the patient is diabetic, special precautions may be necessary (see page 710).
2. Instruct patient to remove street clothing and accessories such as jewelry before the study. Provide a gown for patient use.
3. Stress the importance of remaining still and following breathing instructions during the procedure.
4. Refer to iodine test precautions. Assess female patients for pregnancy status. If positive, advise the radiology department.
5. Omit food and fluid before the examination. Check institutional protocols for specific dietary and fluid restrictions. A laxative may be ordered the evening before the examination.
6. Inform the patient and family that a cholangiogram can be a lengthy procedure lasting > 2 hours.
7. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Care**

1. Be aware that posttest nausea, vomiting, and transient elevated temperature may occur as a reaction to the iodine contrast.
2. Document observations and notify physician if necessary.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.
Clinical Alert

1. Persistent fever, especially if associated with chills, may indicate bile duct inflammation.
2. If the patient has diabetes, assess whether he or she is taking Glucophage/metformin. Due to an increased risk for renal failure, this medication regimen must be discontinued the day of and several days after administration of contrast media. Consult the radiology department for specific instructions.
3. Assess patient for allergies to all substances, specifically latex, and inform the radiology department of any known or suspected sensitivities before study.
4. Assess whether patient is allergic to iodine. If iodine contrast sensitivities are known or suspected, inform the radiology department before study.
5. Monitor for hemorrhage, pneumothorax, or peritonitis after percutaneous transhepatic cholangiography. Unusual pain or tenderness, difficulty breathing, or change in vital signs may signal these complications. If these side effects occur, take immediate action to treat.

Intravenous Urography (IVU); Excretory Urography or Intravenous Pyelography (IVP)

IVU is one of the most frequently ordered tests in cases of suspected renal disease or urinary tract dysfunction.

NOTE

IVU is indicated during the initial investigation of any suspected urologic problem, especially to diagnose kidney and ureter lesions and impaired renal function.

An intravenous radiopaque iodine contrast substance is injected and concentrates in the urine. Following this injection, a series of x-ray films are made at predetermined intervals over the next 20 to 30 minutes. A final postvoid film is taken after the patient empties the bladder.

These films demonstrate the size, shape, and structure of the kidneys, ureters, and bladder and the degree to which the bladder can empty. Renal function is reflected by the length of time it takes the contrast material first to appear and then to be excreted by each kidney. Kidney disease, ureteral and bladder stones, and tumors can be detected with IVU.

CT also may be done in conjunction with IVU to obtain better visualization of renal lesions. This increases examination time. If kidney tomography or nephrotomograms are ordered separately, the procedure and preparation are the same as for IVU.

Reference Values

Normal

1. Normal size, shape, and position of the kidneys, ureters, and bladder. Normal kidneys are approximately as long in dimension as three and one half vertebral bodies. Therefore, kidney size is estimated in relation to this rule of thumb.
2. Normal renal function
   a. Two to 5 minutes after the injection of contrast material, the kidney outline appears on an x-ray film. Threadlike strands of contrast material appear in the calyces.
   b. When the second film is taken several minutes after contrast injection, the entire renal pelvis can be visualized.
   c. Later films show the ureters and bladder as the contrast material makes its way into the lower urinary tract.
   d. No evidence of residual urine should be found on the postvoid film.

Procedure

1. Take a preliminary x-ray (KUB) with the patient in a supine position to ensure that the bowel is empty and the kidney location can be visualized.
2. Inject the intravenous contrast material, usually into the antecubital vein.
3. Alert the patient that during and following the intravenous contrast injection they may experience warmth, flushing of the face, salty taste, and nausea.
   a. Instruct the patient to take slow, deep breaths should these sensations occur. Have an emesis basin and tissue wipes available. Use standard precautions when handling secretions.
   b. Assess for other untoward signs, such as respiratory difficulty, diaphoresis, numbness, palpitations, or urticaria. Be prepared to respond with emergency drugs, equipment, and supplies. These items should be readily available whenever this procedure is performed.
4. Take at least three x-ray films at predetermined intervals following injection of the contrast material.
5. After these three films are taken, instruct the patient to void before the final film is made to determine the ability of the bladder to empty.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications

1. Abnormal IVU findings may reveal the following conditions:
   a. Altered size, form, and position of the kidneys, ureters, and bladder
   b. Duplication of the pelvis or ureter
   c. Presence of only one kidney
   d. Hydronephrosis
   e. Supernumerary kidney
f. Renal or ureteral calculi (stones)
g. Tuberculosis of the urinary tract
h. Cystic disease
i. Tumors
j. Degree of renal injury subsequent to trauma
k. Prostatic enlargement in males
l. Enlarged kidneys suggesting obstruction or polycystic disease kidney
m. Evidence of renal failure in the presence of normal-sized kidneys suggesting an acute rather than chronic disease process
n. Irregular scarring of the renal outlines, suggesting chronic pyelonephritis

2. A time delay in radiopaque contrast visualization is indicative of renal dysfunction. No contrast visualization may indicate very poor or no renal function.

Interfering Factors

1. Feces or intestinal gas will obscure urinary tract visualization.
2. Retained barium can obscure optimal views of the kidneys. For this reason, barium tests should be scheduled after IVU when possible.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test. A written reminder may be helpful to the patient. Screen patients for pregnancy status. If positive, advise the radiology department. If patient has diabetes, special precautions may be necessary (see page 710).
2. Observe iodine contrast test precautions. Assess for all allergies and determine prior allergic reaction to contrast substances. Many radiology departments require a recent creatinine level for all patients >40 years of age before performing this procedure in order to ensure the absence of renal insufficiency.
3. Because a relative state of dehydration is necessary for contrast material to concentrate in the urinary tract, instruct the patient to abstain from all food, liquid, and medication (if possible) for 12 hours before examination. Fasting after the evening meal the day before the test will meet this criterion.
4. Instruct the patient to take a laxative the evening before the examination, and alert the patient that he or she may receive an enema the morning of the test.
   a. Patients with intestinal disorders such as ulcerative colitis should be given a cathartic only when specified by the physician.
   b. Elderly patients may need assistance to the bathroom. Be alert for signs of weakness and stress.
5. Do not give children <7 years of age pretest cathartics or enemas. Should the preliminary x-ray film show intestinal gas obscuring the kidneys, a few ounces of infant formula or carbonated beverage may relieve the concentration of gas at that particular location.
6. Evaluate stool and check for abdominal distention to evaluate for possible barium retention if it has been used in previous studies. Additional bowel preparation may be necessary.
7. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Care

1. Resume prescribed diet and activity after the examination.
2. Teach and encourage the patient to drink sufficient fluids to replace those lost during the pretest phase.
3. Encourage rest, as needed, following the examination. Instruct patients to “let their body tell them” about rest needs.
4. Observe and document mild reactions to the iodine material, which may include hives, skin rashes, nausea, or swelling of the parotid glands (iodism). Notify the physician if the signs and symptoms persist. Oral antihistamines may relieve more severe symptoms.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

NOTE

Elderly or debilitated patients with poor renal reserves may not tolerate these de-hydration protocols (fasting, laxatives, enemas). In such instances, consult with the radiologist or the patient’s physician to ascertain the proper procedure. For infants and small children, fasting time usually varies from 6 to 8 hours pretest. If in doubt, verify protocols with the radiologist or attending physician.
Contraindications to an IVU or IVP include the following conditions:

- Sedation and analgesia may precede insertion of a local anesthetic into the urethra (see Drip infusion pyelography).
- Patients receiving drug therapy for chronic bronchitis, emphysema, or asthma.
- Multiple myeloma, unless the patient can be adequately hydrated during and after the study.
- Advanced pulmonary tuberculosis.
- Follow guidelines in Chapter 1 regarding Combined renal and hepatic disease.
- Diabetes, especially diabetes mellitus.

If a patient has diabetes, assess whether he or she is taking Glucophage/metformin. Due to an increased risk for renal failure and lactic acidosis, this medication regimen must be discontinued the day of and several days after administration of contrast media. Consult the radiology department for specific instructions.

Some physiologic changes can be expected after radiopaque iodine injections. Hypertension, hypotension, tachycardia, arrhythmias, or other electrocardiographic (ECG) changes may occur.

An iodine-based contrast medium is given with caution in the presence of hyperthyroidism, asthma, hay fever, or other allergies.

Observe for anaphylaxis or severe reactions to iodine, as evidenced by shock, respiratory distress, precipitous hypotension, fainting, convulsions, or actual cardiopulmonary arrest. Resuscitation supplies and equipment should be readily available.

In all cases except emergencies, a contrast medium should not be injected sooner than 90 minutes after eating.

Intravenous iodine can be highly irritating to the intimal layer of the veins and may cause painful vascular spasm. If this occurs, a 1% procaine intravenous injection may relieve vascular spasm and pain. Sometimes local vascular irritation is severe enough to induce thrombophlebitis. Warm or cold compresses to the area may relieve pain; however, these do not prevent sloughing. The attending physician should be notified. Anticoagulant therapy may need to be instituted.

Local reactions to iodine may be evidenced by extensive redness, swelling, and pain at the injection site. Even a small amount of iodine contrast entering subcutaneous tissues can cause tissue sloughing, which may require skin grafting. Radiographic evidence of iodine contrast leakage within soft tissues surrounding the injection site confirms extravasation. Treatment may include a local infiltration of hyaluronidase.

Assess for latex allergy and inform the radiology department of any known or suspected sensitivities before study.

Other Tests Used to Examine the Urinary System

1. Excretion urography or intravenous pyelography (IVP): After injection of an intravenous contrast agent, the collecting system (ie, calyces, pelvis, and ureter) of each kidney is progressively opacified. Radiographs are made at 5- to 15-minute intervals until the urinary bladder is visualized.

2. Drip infusion pyelography: This is a modification of conventional pyelography. An increased volume of contrast agent is administered by continuous intravenous infusion.

3. Cystography: The urinary bladder is opacified by means of a contrast agent instilled through a urethral catheter. After the patient voids, air may be introduced into the bladder to obtain a double-contrast study.

4. Retrograde cystourethrogram: After catheterization, the bladder is filled to capacity with a contrast agent, and radiography is used to visualize the bladder and urethra.

5. Voiding cystourethrogram: After contrast material has been instilled into the urinary bladder, films are made of the bladder and urethra during the process of voiding.

Retrograde Pyelography and Other Tests to Examine the Urinary System

Retrograde pyelography generally confirms IVU findings and is indicated when IVU yields insufficient results because of kidney nonvisualization (congenital kidney absence), decreased renal blood flow that impairs renal function, obstruction, kidney dysfunction, presence of calculi, or patient allergy to intravenous contrast material. This x-ray examination of the upper urinary tract begins with cystoscopy to introduce ureteral catheters up to the level of the renal pelvis. Following this, iodine contrast is injected into the ureteral catheter, and x-ray films are then taken. The chief advantage of retrograde pyelography lies in the fact that the contrast substance can be indirectly injected under controlled pressure so that optimal visualization is achieved. Renal function impairment does not influence the degree of visualization.

Reference Values

Normal

Normal contour and size of ureters and kidneys

Procedure

1. This examination is usually done in the surgical department in conjunction with cystoscopy (see Chap. 12).
2. Sedation and analgesia may precede insertion of a local anesthetic into the urethra (see Cystoscopy in Chap. 12). General anesthesia may be required if the patient is not able to cooperate fully with the procedure.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications

Urinary system x-ray results may reveal the following conditions:
1. Intrinsic abnormality of ureters and kidney pelvis (eg, congenital defects)
2. Extrinsic abnormality of the ureters (eg, obstructive tumor or stones)

**Interfering Factors** Because barium may interfere with test results, these studies must be done before barium x-rays are performed.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the test. Screen female patients for pregnancy status. If positive, advise the radiology department.
2. Be aware that the patient or other authorized person must sign and have witnessed a legal consent form before examination in the operating room.
3. Follow iodine contrast test precautions. A recent creatinine level may be required by the radiology department to evaluate the kidney's ability to clear the contrast.
4. Have the patient fast from food and fluids after midnight before the test.
5. Administer cathartics, suppositories, or enemas as ordered.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Care**

1. Observe patient for signs of allergic reaction to iodine contrast.
2. Check vital signs frequently for the first 24 hours following the test. Follow institutional protocols if general anesthetics were administered.
3. Record accurate urine output and appearance for 24 hours following the procedure. Hematuria or dysuria may be common after the examination. If hematuria does not clear and dysuria persists or worsens, notify the physician. Instruct the patient to do the same.
4. Administer analgesics as necessary. Discomfort may be present immediately following the examination and may require a prescriptive analgesic (eg, codeine).
5. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Clinical Alert**

1. Renal function tests of blood and urine must be completed before this examination is done.
2. Assess whether the patient is allergic to iodine. If iodine contrast sensitivities are known or suspected, inform the radiology department before study.
3. Refer to Clinical Alerts posted in the text for Cystoscopy in Chapter 12.

**Arthrography (Joint X-Ray)**

Arthrography involves multiple x-ray examinations of encapsulated joint structures following injection of contrast agents into the joint capsular space. Arthrography is done in cases of persistent, unexplained joint discomfort. Although the knee is the most frequently studied joint, the shoulder, hip, elbow, wrist, temporomandibular joint, and other joints may also be examined. Local anesthetics are used, and aseptic conditions are observed.

**Reference Values**

**Normal** Normal filling of encapsulated joint structures, joint space, bursae, menisci, ligaments, and articular cartilage

**Procedure**

1. Position the patient on the examining table.
2. Surgically prepare and drape the skin around the joint.
3. Inject a local anesthetic into tissues around the joint. It is usually unnecessary to anesthetize the actual joint space.
4. Aspirate any effusion fluids present in the joint. Inject the contrast agents (eg, gas, water, soluble iodine). Remove the needle, and manipulate the joint to ensure even distribution of the contrast material. In some cases, ask the patient to walk or exercise the joint for a few minutes.
5. Remember that during the examination, several positions are assumed to obtain various x-ray views of the joint.
6. Be aware that a special frame may be attached to the extremity to widen the joint space for a better view. Pillows and sandbags also may be used to position the joint properly.
7. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

**Clinical Implications** Abnormal joint x-ray results reveal the following conditions:

1. Arthritis
2. Dislocation
3. Ligament tears
4. Rotator cuff rupture
5. Synovial abnormalities
6. Narrowing of joint space
7. Cysts

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the test. Advise the patient that some discomfort is normal during contrast
injection and joint manipulation.
2. Remember that in most instances, a properly signed and witnessed consent form is required.
3. Refer to iodine test precautions on pages 707–710. Check for known allergies to iodine, other contrast substances, and latex.
4. Advise patient to bring any prior x-ray films of the joint in question to the arthrogram appointment.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Care

1. The joint should be rested for 12 hours.
2. An elastic bandage may be applied to the knee joint for several days after the examination.
3. Ice can be applied to the area if swelling occurs. Pain can usually be controlled with a mild analgesic.
4. Cracking or clicking noises in the joint may be heard for 1 or 2 days following the test. This is normal. Notify the physician if crepitant noises persist or if increased pain, swelling, or restlessness occurs.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Clinical Alert

If diabetic patient is taking Glucophage/metformin, special considerations may be necessary. Consult with the radiology department to determine whether this medication regimen must be discontinued the day of and several days after the study.

Myelography, Myelogram, Spinal Cord X-Ray

Myelography is a radiographic study of the spinal subarachnoid space in which iodine contrast material is introduced into that space so that the spinal cord and nerve roots are outlined and dura mater distortions can be detected.

This study is done to detect neoplasms, ruptured intravertebral disks, or extraspinal lesions such as arthritic stenosis or ankylosing spondylloses. This examination is also indicated when compression of the spinal or posterior fossa neural structure or nerve roots is suspected. The test is frequently done before surgical treatment for a ruptured vertebral disk or release of stenosis. Symptoms may include unrelieved back pain, pain radiating down the leg, absent or abnormal ankle and knee reflexes, claudication of neurospinal origin, or past history of cancer with loss of mobility or bladder control.

Myelograms fall into three categories: positive contrasts using water-soluble iodine, iodized oil contrast, and negative air contrast. Water-soluble iodine contrast is the most commonly used medium for myelograms and is often followed by CT scanning to improve visualization. In low-dose myelograms, injection of a very small amount of water-soluble contrast is immediately followed by scanning.

Reference Values

Normal lumbar, cervical, or thoracic myelogram

Procedure

1. The test is usually done in the radiography department with the patient positioned on his or her abdomen during the procedure.
2. Prepare and drape the puncture area.
3. The procedure is the same as that for lumbar puncture (see Chap. 5), except for the injection of the contrast substance and fluoroscopic x-ray films. With the use of water-soluble contrast, a narrow-bore needle (22-gauge) may be used. A lumbar puncture is done when a lumbar defect is suspected; a cervical puncture is done for a suspected cervical lesion. In children, the level at which the lumbar puncture is performed is much lower than the level in adults to avoid puncturing the spinal cord. Depending on the contrast substance used, it may be removed (oil) or left to be absorbed (water or air).
4. Tilt the table during the procedure to achieve optimal visualization. Use shoulder and foot braces to maintain correct position.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications Abnormal myelogram results reveal distorted outlines of the subarachnoid space that indicate the following conditions:

1. Ruptured intervertebral disk
2. Compression and stenosis of spinal cord
3. The exact level of intravertebral tumors
4. Spinal canal obstruction
5. Avulsion of nerve roots

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of the test. Explain that some discomfort may be felt during the procedure. Disadvantages of water and air contrast include poor visualization and painful headache (air contrast) because of the difficulty in controlling the gas introduced into the area. Oil contrast substances can cause tissue irritation or can be poorly absorbed from the subarachnoid space. Oil may remain visible on x-ray examination for up to 1 year following the original examination. For these reasons, oil and air contrast are rarely used. Refer to iodine contrast test precautions if iodine is used (see pages 707–710).
2. Be aware that a legal consent form must be properly signed and witnessed before the test.
Assess pregnancy status of female patients. Advise the radiology department if positive.

Explain that the examination table may be tilted during the test but that the patient will be securely fastened and will not fall off the table.

Most diagnostic departments require the patient to refrain from eating for about 4 hours before testing. Clear liquids may be permitted and even encouraged to lower the incidence of headaches after the test. Check with the radiology department and physician for specific orders.

Inform the patient that a myelogram usually produces some discomfort. If the patient has trouble moving, a pain reliever may be necessary to allow easier positioning and movement during the test.

Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Posttest Patient Care

1. Bed rest is necessary for 4 to 24 hours after testing. If a water-soluble contrast is used, the head of the bed should be elevated at 45 degrees for 8 to 24 hours after the procedure. The patient is also advised to lie quietly. This position reduces upward dispersion of the contrast medium and keeps it out of the head, where it may cause headache. If oil contrast dye is used, the patient usually must lie prone for 2 to 4 hours and then remain on his or her back for another 2 to 4 hours. If the entire amount of oil contrast is not withdrawn at the end of the procedure, the head must be elevated to prevent the oil from flowing into the brain.
2. Encourage fluid intake to hasten absorption of residual contrast material, to replace cerebrospinal fluid, and to reduce risk for headache and unusual or metallic taste.
3. Check for bladder distention and adequate voiding, especially if metrizamide has been used.
4. Check vital signs frequently (at least every 4 hours) for the first 24 hours after the examination.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Clinical Alert

1. Observe the patient for possible complications such as continued nausea and vomiting, headache, fever, seizure, paralysis of one side of the body or both arms or legs (rare), arachnoiditis (inflammation of the spinal cord coverings), change in level of consciousness, hallucinations, drowsiness, stupor, neck stiffness, and sterile meningitis reaction (severe headache, symptoms of arachnoiditis, slow-wave patterns on electroencephalogram).
2. Alteration of cerebrospinal fluid pressure may cause an acute exacerbation of symptoms that may require immediate surgical intervention. Lumbar punctures should not be done unless absolutely necessary.
3. This test is to be avoided unless there is a reason to suspect a lesion. Multiple sclerosis, for example, may be worsened by this procedure.
4. Determine whether water-soluble, oil, or air contrast was used for the procedure because posttest interventions differ.
5. If nausea or vomiting occurs after the procedure and a water-soluble contrast has been used, do not administer phenothiazine antiemetics such as prochlorperazine (Compazine).
6. Assess whether the patient is allergic to latex or iodine and inform the radiology department of any known or suspected sensitivities before study.
7. If patient has diabetes, assess whether patient is taking Glucophage/metformin. Because of an increased risk for renal failure and lactic acidosis, this medication regimen may need to be discontinued the day of and several days after administration of contrast media. Consult the radiology department for specific instructions.
8. Many radiology departments require the discontinuation of warfarin sodium (Coumadin) therapy for several days before performance of a myelogram. Often, a prothrombin time is required before beginning the examination.

Hysterosalpingography (Uterine and Fallopian Tube X-Rays)

Hysterosalpingography involves radiographic visualization of the uterine cavity and the fallopian tubes to detect abnormalities that may be the cause of infertility or other problems. Normally, a contrast agent introduced into the uterine cavity will travel through the fallopian tubes and “spill” into the peritoneal cavity, where it will be naturally reabsorbed.

Reference Values

Normal Normal intrauterine cavity Patent fallopian tubes

Procedure

1. Have the patient remove all clothing and put on a hospital gown. The bladder should be emptied before the study begins.
2. Have the patient lie supine on the x-ray table in a lithotomy position. Preliminary pelvic x-ray films may be taken.
3. The radiologist or gynecologist introduces a speculum into the patient's vagina and inserts a cannula through the cervical canal. Administer the iodinated contrast agent into the uterus through this cannula.
4. Remove the speculum (unless it is radiolucent), and perform both fluoroscopic and conventional films.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications Abnormal uterine and fallopian tube x-ray findings include the following conditions:

1. Bicornuate uterus or other uterine cavity anomalies
2. Tubal tortuosity
3. Tubal obstruction evidenced by failure of the contrast dye to spill into the peritoneal cavity on one or both sides (bilateral tubal obstruction causes infertility).

Interventions

Pretest Patient Care
1. Explain test purpose and procedure. Some institutions require a properly signed and witnessed informed consent.
2. Refer to iodine contrast test precautions on pages 707–710.
3. Verify date of last menstrual period to ensure that the patient is not pregnant.
4. Advise patient that some discomfort may be experienced but subsides quickly.
5. Suggest that the patient bring along sanitary napkins to wear because some spotting and contrast dye discharge may occur.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Clinical Alert**

1. Pregnancy, active vaginal bleeding, and active pelvic inflammatory disease are contraindications to hysterosalpingography. It is best to perform this test 7–10 days after the onset of menses.
2. If patient has diabetes and is taking Glucophage/metformin, special considerations may be necessary. Consult with the radiology department to determine whether this medication regimen must be discontinued the day of and several days after the study.
3. Assess whether patient is allergic to latex, and inform the radiology department of any known or suspected sensitivities before study.
4. Assess whether patient is allergic to iodine. If iodine contrast sensitivities are known or suspected, inform the radiology department prior to study.

**Posttest Patient Care**

1. Monitor patient for discomfort and administer analgesics as ordered.
2. Instruct the patient to report heavy vaginal bleeding, abnormal discharge, unusual pain, or fever to the referring physician.
3. Interpret test outcomes and counsel about infertility problems.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Angiography (Digital Subtraction Angiography [DSA], Transvenous Digital Subtraction, Vascular X-Ray)**

Digital angiography is a computer-based imaging method of performing vascular studies that require catheterization of certain venous or arterial vessels. Vasculature studies include the carotid vessels; intracranial vessels; those vessels originating from the aortic arch; abdominal vessels, including the celiac, renal, and mesenteric branches; and other peripheral vessels. Digital subtraction angiography began as an intravenous technique, but because of its limitations, other methods of iodine contrast administration may be employed. Although carrying a greater complication risk, intraarterial injection can be used for detailed visceral studies. The presence of the contrast material blocks the path of x-rays and makes blood vessels visible. An image taken just before contrast injection is subtracted from that taken when the contrast material is actually within the vascular system. The resulting image shows only the distribution of the contrast substance. Digital subtraction is used to isolate a clinically relevant subset of information and is particularly useful in preoperative and postoperative evaluations for vascular and tumor surgery.

Visualization of the carotid and vertebral vasculature is possible in patients with a history of stroke, transient ischemic attacks, bruit, or subarachnoid hemorrhage. The procedure may be used as an adjunct to CT or magnetic resonance scanning and may be performed just before these studies in persons who have evidence of an aneurysm, vascular malformation, or hypervascular tumor. Often, a “bi-plane” imaging device is used, producing simultaneous images 90 degrees apart.

The study names are derived from the vascular structure studied and the study method used. Arteriography refers to contrast agent studies of arterial vessels. Venous structures may also be visualized as these procedures progress. Venography is the contrast agent study of peripheral or central veins. Lymphography studies lymph vessels and nodes. Angiocardiography investigates the interior of the heart and adjacent great vessels such as the pulmonary arteries. Aortography refers to a contrast study of aortic segments such as the thoracic aorta (thoracic aortography), the abdominal aorta (abdominal aortography), or the lumbar aorta (lumbar aortography).

Angiographic examinations also can be named for the route used to inject the contrast substance. For example, renal arteriography is performed by inserting a catheter into the abdominal aorta and then directing it into the renal artery. During peripheral arteriography, the contrast is injected directly into the vessel being studied (eg, femoral artery). If done through the venous route, a large bolus of contrast medium is directly injected into a peripheral vein (eg, venous aortography). X-ray films are taken to track the flow of contrast through the right side of the heart, the lungs, and the left side of the heart.

**Reference Values**

**Normal** Normal carotid arteries, vertebral arteries, abdominal aorta and its branches, renal arteries, and peripheral arteries

**Procedure**

1. Cleanse, prepare, and inject the vascular access area with a local anesthetic, using the sterile technique. Depending on the type of study and patient factors, this is commonly the groin or the antecubital area of the arm. Follow standard precautions.
2. Advance the catheter containing a guide wire into the desired vessel or right atrium of the heart. Remove the guide wire, and connect the catheter to a power injector that administers iodine under pressure in defined quantities and at prescribed intervals. Take x-ray images and store on digital or film media. Therapeutic procedures such as angioplasty, stents, and stent placement may be done in concert with this examination.
3. Remove the catheter after the procedure is terminated.
4. Place a dressing over the insertion site, and apply manual pressure to the puncture site for about 5 minutes or until bleeding stops. Tape a more permanent pressure dressing in place; this usually can be removed in 24 hours.
5. Monitor the patient frequently for hemorrhage or hematoma formation.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications Abnormal digital subtraction angiography results reveal the following conditions:

1. Arterial stenosis
2. Large aneurysms
3. Intravascular or extravascular tumors or other masses
4. Total occlusion of arteries
5. Thoracic outlet syndrome
6. Large or central pulmonary emboli
7. Ulcerative plaque
8. Tumor circulation

Interfering Factors
1. Because this examination is very sensitive to physical movement, motion artifact will produce poor images. Consequently, uncooperative or agitated patients cannot be studied. Even the act of swallowing results in unsatisfactory images. Measures to reduce swallowing, such as breath holding, using a bite block, or exhaling through a straw, do not always yield satisfactory results.
2. Vessel overlap of external and internal carotid arteries makes it almost impossible to obtain a select view of a specific carotid artery because contrast fills both arteries almost simultaneously.

Clinical Alert

1. These tests should be used cautiously in patients with renal insufficiency or unstable cardiac disease. Assess for contraindications to iodinated contrast drugs listed on page 709.
2. In the presence of diabetes, assess whether the patient is taking Glucophage/metformin. Due to an increased risk for renal failure and lactic acidosis, this medication regimen must be discontinued the day of and several days after administration of contrast media. Consult the radiology department for specific instructions.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure and document instructions given. Reinforce explanation of test benefits and risks.
2. Ensure that the patient is coherent and cooperative and able to hold his or her breath and remain absolutely still when so instructed.
3. Be aware that a legal consent form must be properly signed and witnessed.
4. Refer to iodine contrast test precautions (see pages 707–710).
5. Determine whether the patient has any known allergies, especially those to iodine, contrast media, or latex. See pages 706–711 for additional assessment criteria.
6. Assess pregnancy status of female patients. If positive, advise the radiology department.
7. Ensure that preprocedure laboratory work is performed in accordance with departmental standards. This generally will include the following tests:
   a. Prothrombin time drawn on day of procedure for any patients on anticoagulation therapy (eg, warfarin sodium [Coumadin])
   b. Creatinine levels for all patients
   c. Recent prothrombin time and partial thromboplastin time (PT/PTT) and platelet count (generally within 30 days)
8. In many instances, administer glucagon intravenously just before abdominal examinations. This serves to reduce motion artifacts by stopping peristalsis.
9. Remember that the few risks include venous thrombosis and infection. When contrast is administered through the venous route, the arteries—which are normally under higher pressure than the veins—can clear the contrast agent through the process of normal circulation. For the same reason, there is less risk for loosening plaques.
10. Advise patient that no food or fluids should be taken within 2 hours before the study to minimize vomiting if an iodine contrast reaction occurs.
11. Be aware that all arteries in a specific area can be visualized during one series of exposures. This overview gives the advantage of being able to evaluate the entire blood supply to a given area at one time. In contrast, during routine angiography, only one specific artery at a time can be visualized.
12. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Care

2. Observe the catheter insertion site for signs of infection, hemorrhage, or hematoma. Use sterile aseptic technique at all times. Monitor neurovascular status of the extremity. Report problems to the physician promptly.
3. Observe for allergic reactions to iodine. Mild side effects include nausea, vomiting, dizziness, and urticaria. Also watch for other complications such as abdominal pain, hypertension, congestive heart failure, angina, myocardial infarction, and anaphylaxis. In susceptible persons, renal failure may occur because higher doses of contrast materials are given compared with conventional arteriograms. Resuscitation equipment and emergency supplies should be readily available. Immediately report these conditions to the physician.
4. Instruct the patient to increase fluid intake to at least 2000 mL during the 24 hours following the procedure to facilitate excretion of the iodine contrast substance.
5. Interpret test outcomes and monitor appropriately.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

### Clinical Alert

1. The catheter puncture site must be observed frequently and closely for hemorrhage, pseudoaneurysm, or hematoma formation. These can be serious complications and require immediate attention should they occur. Many such patients received anticoagulants before the procedure.
2. Vital sign assessment, puncture site assessment, and neurovascular assessments may need to be done as frequently as every 15 minutes for the first few hours after the procedure. Neurovascular assessments include evaluation of color, motion, sensation, capillary refill time, pulse quality, and temperature (warm or cool) of the affected extremity. Compare the affected extremity with the nonaffected extremity.
3. Review the chart or question the patient or physician regarding deficits that were present before the procedure to establish baseline levels of circulatory function. Report postprocedure changes immediately.
4. If an arterial puncture was performed, the affected extremity must not be bent for several hours, and the patient must lie flat other than a pillow under the head. Do not raise the head of the bed or cart because this can put a strain on a femoral puncture site. The patient may turn if the affected extremity is maintained in a straight position without putting strain on the femoral puncture site. If needed, a fracture bedpan can lessen strain on a groin puncture site.
5. If bleeding or hematoma occurs, apply pressure to the site. Sometimes “sandbags” may be applied to the puncture site as a routine part of postprocedure protocols.
6. Maintain a functional intravenous access site. Usually, the patient will return to the nursing unit with an IV in place.
7. A Doppler device may reveal audible pulse sounds if pulses are nonpalpable.
8. Sudden onset of pain, numbness or tingling, greater degree of coolness, decreased or absent pulses, and blanching of extremities are always cues to notify the physician immediately. These signs can indicate arterial occlusion, which may require rapid surgical intervention.

### Lymphangiography (X-Rays of Lymph Nodes and Vessels)

Lymphangiography examines the lymphatic channels and lymph nodes by means of radiopaque iodine contrast injected into the small lymphatics of the foot. This test is commonly ordered for patients with Hodgkin's disease or cancer of the prostate to check for nodal involvement. Lymphangiography is also indicated to evaluate edema of an extremity without known cause, to determine the extent of adenopathy, to stage lymphomas, and to localize affected nodes as part of surgical or radiotherapeutic treatment.

#### Reference Values

**Normal**

Normal lymphatic vessels and nodes

#### Procedure

1. Place the patient in the supine position on the x-ray table.
2. Inject a blue contrast intradermally between each of the first three toes of each foot to stain the lymphatic vessels.
3. Make a 1- to 2-inch incision on the dorsum of each foot after the site is infiltrated with local anesthetic.
4. Identify and cannulate the lymphatic vessel to facilitate extremely low-pressure injection of the iodine contrast medium.
5. Discontinue the injection when the contrast medium reaches the level of the third and fourth lumbar vertebrae as seen on fluoroscopy.
6. Observe that abdominal, pelvic, and upper body films demonstrate the lymphatic vessels filling.
7. Obtain a second set of films 12 to 24 hours later to demonstrate filling of the lymph nodes.
8. View the nodes in the inguinal, external iliac, common iliac, and periarterial areas, as well as the thoracic duct and supraclavicular nodes, using this procedure.
9. Be aware that when a lymphatic of the hand is injected, the axillary and supraclavicular nodes should be visible.
10. Because the contrast dye remains present in the nodes for 6 months to 1 year after lymph-angiography, repeat studies can be done to track disease activity and to monitor treatment without the need to repeat contrast injection. The patient may need to have additional films taken.
11. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

#### Clinical Implications

Abnormal lymph node and vessel x-ray results indicate the following conditions:

1. Retroperitoneal lymphomas associated with Hodgkin's disease
2. Metastasis to lymph nodes
3. Abnormal lymphatic vessels

#### Interventions

**Pretest Patient Care**

1. Explain test purpose and procedure. Obtain a signed, witnessed consent form.
2. See iodine contrast test precautions on pages 707–710.
3. Assess pregnancy status of female patients. If positive, advise the radiology department.
4. Tell patient that no fasting is necessary. Usual medications can be taken.
5. Instruct the patient that he or she may feel some discomfort when the local anesthetic is injected into the feet.
6. Administer oral antihistamines per physician orders if allergy to the iodized contrast agents is suspected.
7. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.
Posttest Patient Care

1. Check and record the patient's temperature every 4 hours for 48 hours after the examination.
2. Provide a restful environment.
3. If ordered, elevate the legs to prevent swelling.
4. Watch for complications such as delayed wound healing, infection, extremity edema, allergic dermatitis, headache, sore mouth and throat, skin rashes, transient fever, lymphangitis, and oil embolism.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Clinical Alert

1. Lymphangiography is usually contraindicated in the following conditions:
   a. Known iodine hypersensitivity
   b. Severe pulmonary insufficiency
   c. Cardiac disease
   d. Advanced renal or hepatic disease
2. The major complication of this procedure relates to contrast media embolization into the lungs. This will diminish pulmonary function temporarily and, in some patients, may produce lipid pneumonia. The patient may require aggressive respiratory management if this complication is life-threatening.
3. If patient has diabetes and is taking Glucophage/metformin, special considerations may be necessary. Consult with the radiology department to determine whether this medication regimen must be discontinued the day of and several days after the study.
4. Assess whether patient is allergic to latex, and inform radiology department of any known or suspected sensitivities before study.
5. Assess whether patient is allergic to iodine. If iodine contrast sensitivities are known or suspected, inform the radiology department before study.

COMPUTED TOMOGRAPHY (CT)

Computed tomography (CT), also called CT scanning, computerized tomography, or computerized axial tomography (CAT), produces x-rays similar to those used in conventional radiography; however, CT scans are taken with a special scanner system. Conventional x-rays pass through the body and produce an image of bone, soft tissues, and air. With CT scans, a computer provides rapid complex calculations that determine the extent to which tissues absorb multiple x-ray beams. CT is unique because it can produce cross-sectional images (ie, “slices”) of anatomic structures without superimposing tissues on each other. Additionally, CT can discern the different characteristics of tissue structures within solid organs. Agents may be used for delineation of blood vessels, the opacification of certain tissue (eg, kidneys), demonstration of bowel, and blood flow patterns.

The patient lies on a motorized table positioned inside a doughnut-shaped frame called the gantry. The gantry contains the x-ray tubes, which rotate around the patient during the scan. By rotating the narrow-beamed x-ray source around the patient's body, multiple attenuation readings are gathered and processed by the computer. The display, similar to a conventional radiograph, demonstrates varying densities that correspond to the absorption of x-rays by the patient's anatomy. As with traditional x-ray techniques, bones appear white, and gas and fat appear black. However, with CT, discrete differences in attenuation can be quantified. This means a CT scan can demonstrate minor differences in density and composition in shades of gray. A CT scan can differentiate tumors from soft tissues, air space from cerebrospinal fluid, and normal blood from clotted blood.

By interpreting the scan, structures are identified by appearance, shape, size, symmetry, and position. Usually, space-occupying lesions show characteristic displacement of surrounding viscera. Scans can be performed at different levels and planes and in different slice thicknesses to isolate small lesions. Often, hollow viscera (eg, intestines) and blood vessels need to be accentuated with the use of contrast media.

Spiral CT scanners, also known as helical CT scanners, are a modification of the conventional CT technique. A spiral scan employs a continuous, “corkscrew” scan pattern that produces a three-dimensional raw data set. This allows for three-dimensional reconstruction and CT angiography. Multi-row scanners are capable of producing up to 16 image slices simultaneously.

Following image acquisition on a multi-row or spiral CT scanner, several postprocessing techniques can be applied to the data sets. This computer manipulation allows for:

1. CT angiography—allowing the vascular system to be viewed in three dimensions without the visualization of overlying structures. Considered a complement to true angiography, the CT technique does have the advantage of requiring only an intravenous stick rather than an arterial puncture.
2. Shaded surface display—a computer-generated surface rendering. The resultant images have the perception of depth, which may be of particular value to surgeons, especially during reconstruction (eg, posttrauma) procedures.

CT scans can be performed on virtually any body part and can isolate virtually any abdominal organ. Typical CT applications include the following studies:

1. Abdomen: to include liver, pancreas, gallbladder, kidneys, adrenals, spleen, retroperitoneum, and abdominal blood vessels
2. Pelvis: to include urinary bladder, uterus, ovaries, distal colon, and prostate
3. Spine
4. Head, sinuses, orbits, mastoids, internal auditory canals, facial bones, neck
5. Chest: to include lungs, mediastinum, and heart
6. Joints and specific bones
7. CT-guided biopsy
8. Fee-for-service screening test may be available to evaluate the heart, lungs, colon, or the entire body.

Computed Tomography (CT) of the Head and Neck; Brain, Eyes, and Sinus Computerized Axial Tomography (CAT)

CT of the head is a relatively simple x-ray examination done by means of a special scanning machine to evaluate for suspected intracranial lesions (see pages 753–754 for CT explanation). The results form a cross-sectional picture of the anatomic structure of the head that includes the internal cranial structure, brain tissue, and surrounding cerebrospinal fluid. This axial image of the head is similar to a view looking down through the top of the head.

Reference Values
Normal: No evidence of tumor, other pathology, or fracture. Typically, low-density tissue areas appear black, whereas higher-density tissues appear as shades of gray. The lighter the shading, the higher the density of the tissue or structure.

Procedure
1. During the test, have the patient lie perfectly still on a motorized table with his or her head comfortably immobilized. The table is moved into a doughnut-shaped frame called a gantry. X-ray tubes situated within this gantry move around the patient in a circular fashion.
2. Inject an iodinated radiopaque contrast substance if tissue density enhancement is desired because a questionable area needs further clarification. Some patients experience nausea and vomiting after receiving this contrast agent.
3. Take additional images during contrast injection.
4. Be aware that during and after the intravenous injection, the patient may experience warmth, flushing of the face, salty taste, or nausea. Encourage the patient to breathe deeply. An emesis basin should be readily available.
5. Watch for other untoward signs such as respiratory difficulty, diaphoresis, numbness, or palpitations.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications
Abnormal CT head and neck scan results reveal the following conditions:
1. Bony and soft tissue tumor masses such as meningiomas, astrocytomas, angiomas, and cysts
2. Intracranial bleeding or hematoma
3. Aneurysm
4. Infarction
5. Infection
6. Sinusitis
7. Foreign bodies

Interfering Factors
1. A false-negative CT scan can occur in the presence of hemorrhage. As hematomas age, their appearance on CT scans changes from high-intensity to low-intensity levels, partly because older hematomas become more transparent to x-rays.
2. Patient movements negatively affect image quality and accuracy.

Interventions
Pretest Patient Care
1. Explain test purpose and procedure. Provide written instructions. Reinforce knowledge regarding possible adverse effects such as radiation exposure or allergy to iodine contrast media. The amount of x-ray exposure for this examination is about the same as that received during a routine skull x-ray.
2. Assess pregnancy status of female patients. If positive, advise the radiology department.
3. Refer to iodine contrast test precautions on pages 706–710. A creatinine level may be required before the study.
4. Generally, the patient should fast 2 to 3 hours before the test if a contrast study is planned. In most cases, prescribed medications can be taken before CT studies.
5. Reassure the patient that scanning produces no greater radiation than conventional x-ray studies.
7. Reassure the patient who is prone to claustrophobia that claustrophobic fear of the scanner is common. Pictures of the scanner or introduction to the scanner may alleviate these fears.
8. Administer analgesics and sedatives, especially to minimize pain and unnecessary movement.
9. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Care
1. Determine whether an iodine contrast substance was used. If used, observe and record information about reactions if they occur. Reactions may include hives, skin rashes, nausea, swelling of parotid glands (iodism), or, most serious of all, anaphylaxis.
2. Notify the physician immediately if allergic reactions occur. Antihistamines may be necessary to treat symptoms.
3. Documentation should include assessment of information needs, instructions given, time examination was completed, patient response to the procedure, and any allergic reactions.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.
Clinical Alert

1. If patient has diabetes and is taking Glucophage/metformin, special considerations may be necessary. Consult with the radiology department to determine whether this medication regimen must be discontinued the day of and several days after the study.
2. Assess whether patient is allergic to iodine or latex. If iodine contrast or latex sensitivities are known or suspected, inform the radiology department before study.

Computed Tomography (CT) of the Body; Computerized Axial Tomography (CAT) Body Scan; Chest, Spine, Extremities, Abdomen, and Pelvis Computed Tomography

Body CT imaging provides detailed cross-sectional images of the chest, abdomen, pelvis, spine, and extremities. When used to evaluate neoplastic and inflammatory disease, CT data acquisition can be rapidly sequenced to evaluate blood flow and to determine vascularity of a mass. This technique, known as dynamic CT scanning, requires the administration of intravenous contrast. In addition, CT can be used to detect intervertebral disk disease, herniation, and soft tissue damage to ligaments within joint spaces.

Conventional x-ray machines produce a “flat” picture, with organs in the front of the body appearing to be superimposed over organs toward the back of the body. The result is a two-dimensional image of the three-dimensional body part. CT imaging produces many cross-sectional anatomic views without superimposing structures. Spiral scanners allow CT angiography and three-dimensional reconstruction techniques.

Reference Values

Normal No apparent tumor or pathology On CT scans, air appears black, bone appears white, and soft tissue appears in various shades of gray. Shade patterns and their correlation to different tissue densities, together with the added dimensions of depth, allow identification of normal body structures and organs.

Procedure

1. Have the patient drink a special contrast preparation several minutes before the CT abdominal examination. This contrast material outlines the bowel so that it can be more readily differentiated from other structures.
2. Have the patient lie supine on a motorized couch that moves into a doughnut-shaped frame called a gantry. X-ray tubes within the gantry move around the patient as the pictures are taken. These films are concurrently projected onto a monitor screen.
3. Have the patient lie without moving, and give breathing instructions.
4. Inject iodine contrast substance and take more pictures if a questionable area requires further clarification. Patients having pelvic CT scans are given a barium contrast enema. Furthermore, female patients undergoing pelvic CT scans may require insertion of a contrast enhanced vaginal tampon to delineate the vaginal wall. Another indication for contrast is blood vessel delineation, the opacification of well-vascularized tissue, and evaluation of blood flow patterns (as for differential diagnosis of hemangioma).
5. Be aware that the patient may experience warmth, flushing of the face, salty taste, and nausea with intravenous injection of the contrast material. Slow, deep breaths may alleviate these symptoms. Have an emesis basin readily available. Watch for other untoward signs such as respiratory difficulty, heavy sweating, numbness, palpitations, or progression to an anaphylactic reaction. Resuscitation equipment and drugs should be readily available. Notify the physician immediately should any of these side effects occur.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

Clinical Implications Abnormal body CT scan findings reveal the following conditions:

1. Tumors, nodules, and cysts
2. Ascites
3. Fatty liver
4. Aneurysm of abdominal aorta
5. Lymphoma
6. Enlarged lymph nodes
7. Pleural effusion
8. Cancer of pancreas
9. Retropertoneal lymphadenopathy
10. Abnormal collection of blood, fluid, or fat
11. Skeletal bone metastasis
12. Cirrhosis of liver
13. Fractures
14. Soft tissue or ligament damage
15. Abscess

Interfering Factors

1. Retained barium can obscure organs in the upper and lower abdomen. Barium tests should be scheduled after CT scans when possible.
2. Inability of the patient to lie quietly produces less-than-optimal pictures.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure. Written explanations may be helpful. Benefits and risks of the test should be
explained to the patient before the procedure.
2. Assess pregnancy status of female patients. If positive, advise the radiology department.
3. Refer to iodine and barium contrast test precautions on pages 706–711.
4. In most cases, allow patient to take usual prescribed medications before CT studies.
5. Inform the patient that an iodine contrast substance may be administered before and during the examination. Determine whether the patient is allergic to iodine. See pages 706–710 for additional assessment criteria. Pelvic CT examinations usually require both intravenous and rectal administration of contrast material. A creatinine level may be required before the study.
6. Be aware that abdominal cramping and diarrhea may occur; therefore, drugs such as glucagon, Lipomul, or Donnatal may be ordered to decrease these side effects.
7. Inform the patient that solid foods are usually withheld on the day of the examination until after test completion. Clear liquids may be taken up to 2 hours before examination. If in doubt, check with the diagnostic department for specific protocols. A patient with diabetes may need to adjust his or her insulin dose and diet before the test (see Clinical Alert on next page).
8. Instruct the patient that he or she may experience warmth, flushing of the face, a salty metallic taste, and nausea or vomiting if intravenous iodine is administered.
9. Claustrophobic sensations while in the CT scanner are common. Show the patient a picture of the scanner before the procedure to alleviate anxiety.
10. Remember that sedation and analgesics may help the patient lie quietly during the test to achieve optimal results.
11. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Care

1. Observe and document reactions to iodine contrast material such as hives, skin rashes, nausea, swelling of parotid glands (iodism), or anaphylactic reaction.
2. Notify the physician immediately if symptoms are serious.
3. Administer antihistamines to relieve the more severe symptoms.
4. Document preparation and instructions given to the patient or significant others, the time the procedure was completed, patient's response to the procedure, any allergic reactions, and subsequent treatment.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

CT Screening

Typically performed as a fee-for-service exam, CT screening is receiving much media attention. Practitioners are divided as to the efficacy and value of these screens. Generally not covered by medical insurance, these procedures are available without a prescription to those who are able to pay. Further controversy exists as to the type of scanner used for the screening studies. No definitive data exist to show superiority of one technique over another. Scanners used for this purpose may be a spiral, multi-row CT scanner or an electron-beam scanner (EBT). The most common screening CT exams are:

1. Calcium scoring (cardiac scoring, heart scan)—used to detect and measure the amount of calcium plaque within the coronary arteries. Considered predictive for the potential for coronary events.
2. Lung screening—used to detect the presence of lung masses for patients with a significant risk (smoking history, toxin exposure).
3. Whole-body screening—a head-to-toe scan to detect the presence of lesions.
4. CT colonography—a new technique that may potentially serve as a substitute to colonoscopy.

In CT colonography, a small amount of CO₂ gas is administered through a rectal catheter. Rapid scans are taken with the patient in a prone and supine position. Using complex postprocessing software, images of the entire colon are laid out, presented in a “filet” format, or they can be “flown through” with computer animation. Although the preparation for CT colonography is the same as for traditional colonoscopy, patient discomfort is lessened, and acceptance of the process appears to be enhanced.

BIBLIOGRAPHY


Cochran ST: Determination of serum creatinine levels prior to administration of radiographic contrast media. JAMA 277(7): 517–518, 1997

INTERNET SITES

intelhealth.com
diabetesmonitor.com
epa.gov/radiation
postgradmed.com
imaginis.com
acr.org
breastcancer.org
gastronews.com
emedicine.com
auntminnie.com
CYTOLOGIC AND HISTOLOGIC STUDIES

Overview of Cytologic (Cells) Studies

Exfoliated cells in body tissues and fluid are studied to determine the types of cells present and to diagnose malignant and premalignant conditions. The staining technique developed by Dr. George N. Papanicolaou has been especially useful in diagnosis of malignancy and is now used routinely in the cytologic study of the female genital tract as well as in many types of nongynecologic specimens.

Some cytologic (cells) specimens (e.g., smears of the mouth, genital tract, nipple discharge) are relatively easy to obtain for study. Other samples (e.g., amniotic fluid, pleural effusions, cerebrospinal fluid [CSF]) are from less accessible sources, and special techniques, such as fine-needle aspiration, are required for collection. Tissue (histologic) samples may be obtained by biopsy during surgery or during outpatient diagnostic procedures such as endoscopy. In all studies, the source of the sample and its method of collection must be noted so that the evaluation can be based on complete information.

Specimens for cytologic and histologic study usually consist of many different cells. Some are normally present, whereas others indicate pathologic conditions. Cells normally observed in one sample may, under certain conditions, be indicative of an abnormal state when observed elsewhere. All specimens are examined for the number of cells, cell distribution, surface modifications, size, shape, appearance and staining properties, functional adaptations, and inclusions. The cell nucleus is also examined. Any increases or decreases from normal values are noted.

Gynecologic specimens may be smeared and fixed in 95% alcohol. Some types of spray fixative are also available. (Gynecologic specimens collected using the liquid-based technique are collected in special [i.e., PreservCyt] solution.) Nongynecologic specimens are generally collected without preservative. They may be placed in saline, and they must be handled carefully to prevent drying or degeneration. Check with your individual laboratory for collection requirements. It is important that all cytology specimens be sent to the laboratory as soon as they are obtained to prevent disintegration of cells or any other process that could cause alteration of the material for study.

Clinical Alert

1. The test is only as good as the specimen received.
2. Specimens collected from patients in isolation should be clearly labeled on the specimen container and requisition form with appropriate warning stickers. The specimen container should then be placed inside two sealed, protective biohazard bags before it is transported to the laboratory.
3. The U.S. Occupational Safety and Health Administration (OSHA) requires that all specimens be placed in a secondary container before transportation to the laboratory. Most laboratories prefer plastic biohazard bags. Requisitions should be kept on the outside of the bag or in a separate compartment in the biohazard bag, if available.

In practice, results of cytologic studies are commonly reported as:

1. Inflammatory
2. Benign
3. Atypical
4. Suspicious for malignancy
5. Positive for malignancy (in situ versus invasive)

Overview of Histologic (Tissue) Studies

Material submitted for tissue examination may be classified according to its histologic or cellular characteristics. A basic method for classifying cancers according to the histologic or cellular characteristics of the tumor is Broder's classification of malignancy:

1. Grade I: tumors showing a marked tendency to differentiate; 75% or more of cells differentiated
2. Grade II: 75% to 50% of cells differentiated, slight to moderate dysplasia and metaplasia
3. Grade III: 50% to 25% of cells differentiated, marked dysplasia, marked atypical features, and cancer in situ
4. Grade IV: 25% to 0% of cells differentiated

The tumor-node-metastasis (TNM) system is a method of identifying tumor stage according to spread of the disease. This system evolved from the work of the International Union Against Cancer and the American Joint Committee on Cancer. In addition, the TNM system further defines each specific type of cancer, (eg, breast, head, neck). This staging system (Chart 11.1) is employed for previously untreated and treated cancers and classifies the primary site of cancer and its extent and involvement, such as lymphatic and venous invasion.

<table>
<thead>
<tr>
<th>Chart 11.1 TNM System</th>
</tr>
</thead>
<tbody>
<tr>
<td>Three capital letters are used to describe the extent of the cancer:</td>
</tr>
<tr>
<td>T: Primary tumor</td>
</tr>
<tr>
<td>N: Regional lymph nodes</td>
</tr>
<tr>
<td>M: Distant metastasis</td>
</tr>
<tr>
<td>Lower-case letters are used to indicate the chronology of classification:</td>
</tr>
<tr>
<td>c: Clinical—diagnostic</td>
</tr>
<tr>
<td>p: Postsurgical treatment—pathologic</td>
</tr>
<tr>
<td>r: Retreatment</td>
</tr>
<tr>
<td>a: Autopsy</td>
</tr>
<tr>
<td>This classification is extended by the following designations:</td>
</tr>
<tr>
<td>T Subclasses (Extent of Primary Tumor)</td>
</tr>
<tr>
<td>TX: Tumor cannot be adequately assessed</td>
</tr>
<tr>
<td>T0: No evidence of primary tumor</td>
</tr>
<tr>
<td>Tis: Carcinoma in situ</td>
</tr>
<tr>
<td>T1, T2, T3, T4: Progressive increase in tumor size and involvement</td>
</tr>
<tr>
<td>N Subclasses (Involvement of Regional Lymph Nodes)</td>
</tr>
<tr>
<td>NX: Regional lymph nodes cannot be assessed clinically</td>
</tr>
<tr>
<td>N0: Regional lymph node metastasis</td>
</tr>
<tr>
<td>N1, N2, N3, N4: Increasing degrees of demonstrable abnormality of regional lymph nodes</td>
</tr>
<tr>
<td>HISTOPATHOLOGY GX: Grade cannot be assessed</td>
</tr>
<tr>
<td>G1: Well-differentiated grade</td>
</tr>
<tr>
<td>G2: Moderately well-differentiated grade</td>
</tr>
<tr>
<td>G3: Poorly differentiated grade</td>
</tr>
<tr>
<td>G4: Undifferentiated</td>
</tr>
<tr>
<td>Metastasis</td>
</tr>
<tr>
<td>MX: The minimum requirements to assess the presence of distant metastasis cannot be met</td>
</tr>
<tr>
<td>M0: No evidence of distant metastasis</td>
</tr>
<tr>
<td>M1: Distant metastasis present (specify sites of metastasis)</td>
</tr>
<tr>
<td>The category M1 may be subdivided according to the following notations:</td>
</tr>
<tr>
<td>Pulmonary: PUL</td>
</tr>
<tr>
<td>Hepatic: HEP</td>
</tr>
<tr>
<td>Osseous: OSS</td>
</tr>
<tr>
<td>Brain: BRA</td>
</tr>
<tr>
<td>Lymph nodes: LYM</td>
</tr>
<tr>
<td>Skin: SKI</td>
</tr>
<tr>
<td>Bone marrow: MAR</td>
</tr>
<tr>
<td>Peritoneum: PER</td>
</tr>
<tr>
<td>Pleura: PLE</td>
</tr>
<tr>
<td>Other: OTH</td>
</tr>
</tbody>
</table>

In certain sites, further information regarding the primary tumor may be recorded under the following headings:
In certain sites, further information regarding the primary tumor may be recorded under the following headings:

<table>
<thead>
<tr>
<th>LYMPHATIC INVASION (L)</th>
<th>LX: Lymphatic invasion cannot be assessed</th>
</tr>
</thead>
<tbody>
<tr>
<td>L0: No evidence of lymphatic invasion</td>
<td></td>
</tr>
<tr>
<td>L1: Lymphatic invasion</td>
<td></td>
</tr>
<tr>
<td>VENOUS INVASION (V)</td>
<td>VX: Venous invasion cannot be assessed</td>
</tr>
<tr>
<td>V0: No venous invasion</td>
<td></td>
</tr>
<tr>
<td>V1: Microscopic venous invasion</td>
<td></td>
</tr>
<tr>
<td>V2: Macroscopic venous invasion</td>
<td></td>
</tr>
</tbody>
</table>

Information on residual tumor does not enter into establishing the stage of the tumor but should be recorded for use in considering additive therapy. When the cancer is treated by definitive surgical procedures, residual cancer, if any, is recorded.

**RESIDUAL TUMOR (R)**

RX: Residual tumor at primary site cannot be assessed
R0: No residual tumor
R1: Microscopic residual tumor
R2: Macroscopic residual tumor

(M): Symbol in parentheses indicates multiple tumors
Y: Symbol—Y prefix indicates classification occurring with intense multimodality therapy
Z: Symbol—Z prefix indicates recurrent tumors after a disease-free interval


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**Fine-Needle Aspirates: Cell (Cytologic) and Tissue (Histologic) Study**

Fine-needle aspiration is a method of obtaining diagnostic material for cytologic (cell) and histologic (tissue) study that causes a minimal amount of trauma to the patient. Aspirates may be obtained from all parts of the body, including the mouth, breast, liver, genital tract, respiratory tract, urine, cerebrospinal fluid, and thyroid. Bacteriologic studies may also be done on material obtained during fine-needle aspiration. Unfixed material, left in the syringe or on a needle rinsed in sterile saline, may be taken to the microbiology department for study.

**Reference Values**

Normal: Benign or negative: no abnormal cells or abnormal tissue present
No pathogenic organisms

**Procedure**

1. Use local anesthesia in most cases. Aspirate superficial or palpable lesions without radiologic aid, but aspirate nonpalpable lesions using radiographic imaging as an aid for needle placement. Use sterile technique.
2. Position the needle properly, then retract the plunger of the syringe to create negative pressure. Move the needle up and down, and sometimes at several different angles. Release the plunger of the syringe and remove the needle.
3. Express material obtained onto glass slides, which must either be fixed immediately in 95% alcohol, spray fixed, or air dried, depending on the staining procedure used by the laboratory. The remaining material may be placed in a preservative solution, such as 50% alcohol. Check with your laboratory for recommended fixation requirements. Material may also be sent to the laboratory in the syringe.
4. Record the source of the sample and method of collection so that evaluation can be based on complete information.
5. Clearly label specimens collected from patients in isolation on the specimen container and on the requisition form with an appropriate warning sticker. Place the specimen container inside two sealed, protective biohazard bags before transport.
6. See Chapter 1 guidelines for intratest care.

**Clinical Implications**

Abnormal results reported as atypical, suspicious for malignancy, and positive for malignancy (in situ versus invasive) are helpful in identifying:

1. Infectious processes. The infectious agent may be seen, or characteristic cellular changes may indicate the infectious agent that is present.
2. Benign conditions. Some characteristic cellular changes may be present, indicating the presence of a benign process.
3. Malignant conditions, either primary or metastatic. If the disease is metastatic, the findings may be reported as consistent with the primary malignancy.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure, benefits, and risks of the test. Even though a local anesthetiic is used, the procedure causes some discomfort, and this should not be minimized. If the approach involves passing near a rib, the pain may be greater because of the sensitivity of the bone; this is not a cause for alarm. Unexpected pain may induce a vasovagal or other undesirable response. Other risks include infection and hematoma or hemorrhage, depending on the site aspirated.
2. See guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Monitor for signs of inflammation and use site care infection control measures. Treat pain, which may be common in sensitive areas such as the breast, nipple, prostate, and scrotum. Monitor for specific problems, which vary depending on the site aspirated (eg, hemoptysis after a lung aspiration).
2. Counsel about follow-up procedures for infections and malignant conditions.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Sentinel Node Location Before Biopsy (Breast, Melanoma): Special Prebiopsy Study**

The concept of identifying and localizing the sentinel node or nodes before biopsy is that these nodes receive initial lymphatic drainage and are the first filter to remove metastatic cells; thus, if this sentinel node is free of disease, then the rest of the nodes in the patient will...
be free of disease, also. Three methods (along with marking of the skin) are used: (1) lymphoscintigraphy (preoperative), (2) nuclear probe localization (intraoperative), and (3) blue dye injection (intraoperative). Often, all three techniques are used together, the lymph nuclear scan being the most common (see Procedures). These special prebiopsy procedures are done before biopsies to diagnose cancer of breast or melanoma. Indications for lymph nuclear scan lymphoscintigraphy include detection of metastasis, mapping of all sentinel nodes, and staging and monitoring cancers, such as melanoma, breast, head, neck, and skin. Indications for using the nuclear gamma-radiation probe include detecting the most sentinel nodes and providing auditory confirmation. Also, see Chapter 9 for more information on nuclear scans. Indications for blue dye staining include to provide visual confirmation of nodes and to map tumor route (urine will turn blue and skin will stain).

### Clinical Alert

Only when the sentinel node is positive is a complete nodal dissection performed.

### Reference Values

**Normal** No evidence of tumor activity No blocked lymphatic drainage

**Procedure**

1. **Lymph nuclear scan (lymphoscintigraphy)**
   a. For the breast, inject the radiopharmaceutical (large volume) subcutaneously into the breast and adjacent to suspected breast tumor; and for lymphedema, into webs of fingers and toes.
   b. For melanoma, make four to six intradermal injections around the tumor or excision site, avoiding scar tissue.
   c. Perform immediate imaging with the patient in the position expected during surgery.
2. **Nuclear radiation (gamma) probe (which produces sound)**
   a. Remember that a previously administered radiopharmaceutical and the sound radiation detector permit node detection and localization to determine where the initial operative incisions can be made.
   b. Use the sound-radiation gamma probe to locate the area of radioactivity, not associated with the injection sites. Of the three procedures, the probe is the most sensitive.
3. **Blue dye (not externally visible)**
   a. In order to identify the nodes to undergo biopsy, inject the feet in the web between the toes and the hands between the second and third fingers (allergic reaction to the dye may occur).
   b. Remember that an operative biopsy procedure follows.

### Clinical Implications

1. Abnormal findings reveal metastatic nodes and routes of spread.
2. Asymmetry may indicate lymph flow obstruction.

### Interventions

#### Pretest Patient Care

1. Explain purpose of sentinel node identification procedures.
2. Inform the patient that if the results are positive, surgery usually follows soon after.

#### Intratest Patient Care

1. Be aware that usually sedation or analgesia is not ordered.
2. Mark the site of lymph nodes with indelible pen.
3. Provide support, assist with positioning, and assure the patient that testing is proceeding as expected.

#### Posttest Patient Aftercare

1. Monitor injection site (breast, toes, fingers, or around tumor excision site). Check for signs of inflammation or bleeding.
2. If surgery is planned, prepare according to established protocol. Also, see Biopsies Overview in alphabetical listing.
3. When surgery is scheduled, take images just before transfer to the operating room (2 hours later or next morning).
4. Counsel about outcomes and possible need for further testing or treatment.

### Tissue (Histologic) Biopsy Studies: Overview; Prognostic and Predictive Markers

Tissue biopsies from many body sites (ie, breast, liver, kidney, lymph nodes, skin, bone, muscle, lung, bladder, prostate, thyroid, cervix) may be examined for the presence of benign, toxic, or malignant cells and conditions. The amount of tissue obtained and submitted to the laboratory depends on the specimen site and disease process (eg, in liver biopsy, at least two to three liver cores > 2 cm in length). These procedures may be performed in outpatient or inpatient settings. Some specimens should be collected early in the day. For ultrasound-guided prostate specimens (ie, transrectal ultrasound [TRUS]), 6 to 12 threadlike sections of tissue are obtained, ranging from 0.5 to 1.5 cm in length. Pain and bloody urine are common after. Depending on the body site sampled, anesthetic (ie, local or general) or conscious sedation and analgesia (see Appendix C) is necessary. Tissue obtained for routine histologic (tissue study) and pathologic examination requires special handling (eg, place in 10% formalin or send fresh and intact). Tissue needed for frozen-section examination must be delivered to the laboratory immediately with no fixative added. Tissue needed for special studies (eg, special stains for microorganisms, hormonal studies, DNA ploidy, bone biopsies) may need special handling. A frozen section is done upon the pathologist's recommendation. Tissue freezing (frozen section) may actually be contraindicated and not in the patient's best interest. Contact your individual laboratory for specific instructions. After the biopsy specimen is sent to the laboratory, various tests are done to identify the unique characteristics of the patient's tumor cells and to select correct chemotherapy based on resistance to specific drugs. Multiple and complex genetic changes result from loss of control over normal cell growth, and these alterations may influence the tumor's response to chemotherapy. To measure these changes, four major testing groups are used and include the following:
1. **Extreme drug resistance (EDR) assay** tests of solid tumors and malignant fluids (blood, bone marrow effusions), which determine the probability of a tumor’s resistance to specific chemotherapeutic drugs (eg, 5-fluorouracil [5-FU] for colon cancer). If the tumor cells grow in the presence of extreme exposures to a specific drug, this indicates the presence of significant drug resistance and, by identifying inactive agents, avoids exposing patients to the toxicity of drugs that are likely to be ineffective, saves valuable treatment time, and decreases the possibility of cross-resistance to other effective therapies.

2. **Differential staining and cytotoxicity (DISC) assay** uses special stains and techniques to detect drug resistance in leukemia, lymphoma, blood, and bone marrow specimens.

3. **Prognostic markers** measure the tumor’s growth potential or ability to invade other tissues (metastasis). Tumor cells release proteases and angiogenic factors to break down basement membranes and induce new vascularization of the tumor, which delivers oxygen and nutrients to the tumor and allows micrometastasis to distant sites.

4. **Predictive markers** identify specific mechanisms of drug resistance and provide information on how effective clinically indicated chemotherapy agents will be in treating the patient's tumor cells. Prognostic and predictive markers use molecular probes to determine the genetic characteristics, amount of protein, proliferation index, resistance mechanisms, receptor status, and other defining factors of the patient's malignant tumor. To obtain the most comprehensive analysis of the patient's unique tumor biology, drug resistance testing is done in combination with oncoprofiles and prognostic and predictive markers for the specific cancer type. A **radiation resistance assay** can also be done before the treatment actually begins.

These combined studies identify cervical cancer resistive to internal and external radiation plus chemotherapy (the standard treatment is prognostic indicators of progression-free survival). Also included are p53, thrombospondin-1 (Tsp-1), CD31, and angiogenesis index (AI). Prognostic and predictive markers are as follows:

1. **Androgen receptor.** This receptor predicts prostate cancer's response to hormone therapy.

2. **Angiogenesis index** (p53, Tsp-1, CD31). The AI defines a patient's risk for occult metastatic disease and is composed of factors that characterize the capacity for new blood vessel formation: p53, Tsp-1, and CD31 (vessel count). The p53 gene contributes to tumor growth suppression by slowing cell cycle progression and promoting apoptosis in damaged tumor cells. It also suppresses tumor angiogenesis. Tsp-1 levels have been found to decrease after the tumor sustains mutations in p53. CD31 is expressed on the membrane of endothelial cells, allowing for microvessel count in the tumor.

3. **BAX.** Increased levels of BAX, a 21-kd protein and amino acid, indicate accelerated programmed cell death induced by apoptotic stimulus.

4. **Proto-oncoprotein bc12 (apoptosis regulator).** The translocation of the bc12 gene, occurring in follicular lymphomas, is brought under control of the immunoglobulin gene promoter, resulting in increased intracellular levels of bc12 protein. This protein suppresses programmed cell death (apoptosis). Induction of cell death is an important mechanism for many chemotherapeutic agents. An abnormal expression of bc12 protein can render tumor cells resistant to chemotherapeutic agents.

5. **Cathepsin D (invasion potential).** Cathepsin D, a lysosomal acid protease, has been associated with metastatic potential. Elevated levels of cathepsin D are predictors of early recurrence and death in node-negative cancer and breast cancer.

6. **CD31 (component of tumor angiogenesis index).** CD31 stains microvessels, allowing for counting, and helps to predict more aggressive disease, metastases, poor survival, and new vascularization of the tumor mass.

7. **DNA ploidy and S phase (flow cytometry).** DNA ploidy and proliferative index are independent indicators of prognosis. Patients with aneuploid tumors or high S-phase fractions have poor disease-free survival compared with patients with diploid or low S-phase fraction tumors. DNA ploidy (image analysis) (Feulgen stain) is an indicator of prognosis in selected tumor types in fresh specimens.

8. **Epidemial growth factor receptor (EGF-R).** This growth factor receptor is a glycoprotein tyrosine kinase, either EGF or transforming growth factor-a (TGF-a). When high levels occur in breast, prostate, ovarian, lung, and squamous cell carcinomas, there is an association with poorer prognosis and poor disease-free survival.

9. **Endoglin (CD105).** Endoglin normally occurs in vascular endothelial cells of capillaries, arterioles, small arteries, and venules. Increased levels are found in tumor vessels and proliferating endothelial cells. Endoglin has been found in non-Thor-B and pre-B acute lymphoblastic leukemia (ALL) and acute myelocytic and myelomonocytic leukemia cells.

10. **Estrogen receptor (ER) and progesterone receptor (PR).** ER and PR positivity is associated with a 70% response rate to antihormonal therapy. In contrast, the response rate is less than 10% among patients whose tumors are ER and PR negative. Patients whose tumors are ER and PR positive generally achieve superior disease-free survival.

11. **Glutathione S-transferase (GST): alkylator resistance.** GST is an enzyme that inactivates certain anticancer agents by linking glutathione to the drug. Increased GST levels are associated with drug resistance to chlorambucil and melphalan.

12. **HER2/neu c-erbB2 oncoproteins.** The presence of HER2/neu, a protein that functions as an oncogene, is associated with poorer prognosis. HER2/neu detection also provides information on the potential treatment response to Herceptin.

13. **Ki-67 (proliferative index).** This is a staining technique. Monoclonal antibody Ki-67 is associated with increased cell proliferative activity in tumors and is associated with more aggressive tumors and poor disease-free survival.

14. **MDR-1 (P170 glycoprotein: multidrug resistance).** The presence of MDR-1 cancer cells is associated with resistance to naturally produced chemotherapeutic agents such as paclitaxel (Taxol), doxorubicin, and etoposide and plays a critical role in the selection of a treatment regimen.

15. **O6-methylguanine-DNA methyltransferase (MGMT) (nitrosourea resistance).** MGMT, a repair protein, occurring after DNA damage caused by nitrosoureas, such as BCNU. Brain cancer patients with high levels of the MGMT gene and alkyltransferase (AT) have shorter disease-free and overall survival.

16. **Multidrug resistance protein (MRP).** This protein is similar to, but distinct from, MDR-1, and is strongly associated with resistance to cisplatin drugs in ovarian cancer.

17. **p21.** A protein-like tumor suppressor like p53, p21 controls when and how the cell replicates. Low levels of p21 are associated with increased risk for tumor occurrence, and no p21 contributes to aggressive growth in some tumors.

18. **p53 (cell cycle and Tsp-1 regulator).** The tumor suppressor gene p53 regulates cell cycle progression, cellular...
proliferation, DNA repair, apoptosis (cell death), and angiogenesis. Increased levels of mutated p53 protein in tumor cell nuclei are associated with tumor progression and a poorer prognosis.

19. **PCNA (proliferative index).** Presence of PCNA protein is associated with cell proliferation and with increased levels occurring with more aggressive tumors and poor disease-free survival.

20. **Thymidylate synthase (TS; 5-FU resistance).** Drug resistance tests of thymidylate synthase (TS), a cellular enzyme essential for DNA biosynthesis and cell proliferation that is a target for 5-FU, is an important component of some breast cancer and colon cancer treatment regimens. Increased TS expression correlates with poorer response rates to 5-FU and with shorter survival in breast and colon cancer.

21. **Thrombospondin-1 (Tsp-1).** This extracellular matrix protein is involved in wound healing. Low value is associated with increased tumor neovascularity and mutant p53 expression.

22. **Thymidylate synthase (TS; 5-FU resistance).** Drug resistance tests of thymidylate synthase (TS), a cellular enzyme essential for DNA biosynthesis and cell proliferation that is a target for 5-FU, is an important component of some breast cancer and colon cancer treatment regimens. Increased TS expression correlates with poorer response rates to 5-FU and with shorter survival in breast and colon cancer.

23. **Vascular endothelial growth factor (VEGF).** Vascular endothelial growth factor, or vascular permeability factor (VPF), plays an important role in angiogenesis, which promotes tumor progression and metastasis.

Oncoprofiles provide the maximum useful information from a single biopsy specimen. These disease-specific marker studies include tests that have been associated with clinical outcomes for each cancer type. Oncoprofiles identify relative risk for relapse and assist in planning therapy to each patient's specific tumor. Table 11.1 shows an example of oncoprofiles offered by Oncotech, Inc., of Irvine, California.

**Table 11.1 Useful Information From a Single Biopsy Specimen**

<table>
<thead>
<tr>
<th>Oncoprofile</th>
<th>Basic Profile</th>
<th>Comprehensive Profile</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bladder cancer</td>
<td>DNA, p53, HER2/neu</td>
<td>DNA, p53, HER2/neu, CD31</td>
</tr>
<tr>
<td>Brain cancer</td>
<td>DNA, p53, HER2/neu</td>
<td>DNA, p53, HER2/neu, CD31</td>
</tr>
<tr>
<td>Breast cancer</td>
<td>DNA, ER/PR, HER2/neu</td>
<td>DNA, ER/PR, HER2/neu, p53, CD31</td>
</tr>
<tr>
<td>Colon cancer</td>
<td>DNA, p53</td>
<td>DNA, p53, TS, MDR-1, CD31</td>
</tr>
<tr>
<td>Endometrial cancer</td>
<td>DNA, ER/PR, Ki-67</td>
<td>DNA, ER/PR, Ki-67, CD31, MDR-1, p53</td>
</tr>
<tr>
<td>Kidney cancer</td>
<td>DNA, MDR-1</td>
<td>DNA, MDR-1, p53, CD31</td>
</tr>
<tr>
<td>Leukemia/non-Hodgkin's lymphoma</td>
<td>DNA, Ki-67</td>
<td>DNA, Ki-67, bc12, p53, MDR-1</td>
</tr>
<tr>
<td>Lung cancer</td>
<td>DNA, p53</td>
<td>DNA, p53, MDR-1, bc12</td>
</tr>
<tr>
<td>Melanoma</td>
<td>DNA, MDR-1</td>
<td>DNA, MDR-1, p53, CD31</td>
</tr>
<tr>
<td>Ovarian cancer</td>
<td>DNA, ER/PR, HER2/neu, EGF-R</td>
<td>DNA, ER/PR, HER2/neu, EGF-R, p53, MDR-1</td>
</tr>
<tr>
<td>Prostate cancer</td>
<td>DNA, AR</td>
<td>DNA, AR, p53, CD31</td>
</tr>
<tr>
<td>Sarcoma</td>
<td>DNA, p53</td>
<td>DNA, p53, MDR-1</td>
</tr>
<tr>
<td>Unknown primary site</td>
<td>DNA, p53, HER2/neu</td>
<td>DNA, p53, HER2/neu, MDR-1</td>
</tr>
</tbody>
</table>

*The laboratory report from these tumor studies should provide answers to questions such as “Is tumor malignant?”, “Is type of cancer identified?”, “How aggressive is the cancer?”, “Is the cancer likely to recur?”, and “To which drugs is the tumor resistant?”

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and biopsy procedure, and obtain a signed, witnessed consent form.
2. Remember that patient preparation depends on the predetermined biopsy site. Complete blood count (CBC), prothrombin time (PT), and other bleeding time determinants may be required. Obtain a pertinent history (eg, prior radiation therapy, other cancer, current medications, pregnancy).
3. Follow Chapter 1 guidelines for safe, effective, and informed pretest care.

**Clinical Alert**

Contraindications to tissue biopsy depend on the body site sampled: bleeding diathesis (uncontrolled), anticoagulant therapy, highly vascular lesions, sepsis, seriously impaired lung function, an uncooperative patient, or local infection near the biopsy site.

**Posttest Patient Aftercare**

1. Monitor for signs of bleeding, inflammation, infection, laceration of tissue and organs, and perforation. Treat pain, which may be experienced to various degrees depending on the body site sampled.
2. Counsel the patient about follow-up procedures and treatment for infections and malignant conditions.
3. Follow the guidelines in Chapter 1 for safe, effective, and informed posttest care.

**Breast Biopsy: Cell (Cytologic) and Tissue (Histologic) Study and Prognostic Markers** Breast biopsies are among the most common type of biopsy done. The cells and tissue obtained by breast biopsy establish the presence of breast disease, diagnose histopathology, and classify the process. They also confirm and characterize calcifications noted in prebiopsy mammograms. The breast tissue is examined to determine surgical margins, presence or absence of vesicular invasion, tumor type, staging, and grading. Secondary studies relevant to survival may include imaging procedures, along with the following prognostic markers. (Also see Tumor Markers in Chapter 8 for more information.)
1. ER and PR. These hormone receptors are indicators of prognosis and are used to manage hormonal therapy in breast and endometrial cancer. Immunohistochemical (IHC) staining aids recognition of metastatic breast cancer.

2. DNA ploidy. This test measures cell turnover or replication; it is used to predict prognosis and shorter survival times by the presence of aneuploid (rapidly replicating cells) for certain tumor types, breast, prostate, and colon; less clear for ovarian, lung, kidney, and bladder (ureter) (66% of breast cancers are aneuploid).

3. S-phase fraction (SPF)—to predict survival and reduced chance of relapse. Low levels of SPF appear to have longer survival and reduced chance of relapse. SPF is the DNA synthesis phase obtained by a statistical method.

4. Cathepsin D—done to determine prognosis. The presence of this lysosomal protease is estrogen related and may promote tumor spread. Prognostic significance remains ambiguous.

5. EGF reception—done to predict survival time. Presence is correlated with ER negativity, aneuploidy, increased S-phase factors, and lymph node metastases. Increased EGF reception may be associated with worse relapse free and survival time.

6. p53 gene—used to predict prognosis. This tumor suppression gene regulates cell cycles. Some clinicians believe that the p53 gene’s prognostic value is second only to lymph node status.

7. c-erbB2 (HER-2) oncogene—determines which patients are most likely to benefit from high doses of chemotherapy. High levels of this oncogene receptor are associated with poor response to conventional chemotherapy and may be a marker for patients likely to benefit from high doses of chemotherapy. HER-2/neu levels may also be determined in a blood specimen.

Reference Values

Normal No atypical or normal cells Gail Index of breast cancer risk = odds ratio = 1.7. For more information, see Gail, Brinton, Byar, et al., 1989.

Procedure

1. See Chapter 10 (mammography) for image guided tumor localization study before biopsy.

2. Be aware that breast tissue specimens may be obtained by open surgical technique by x-ray–guided core biopsy or by needle biopsy. Place these specimens in a biohazard bag, take directly to the laboratory, and give to the pathologist or histotechnologist. The breast tissue is examined and the extent of the tumor determined. Reaction margins and the grade and stage of disease are identified.

3. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. After breast tissue is examined, the extent of the tumor is determined. Resection margins are evaluated, and grade and stage of disease are identified. The further dedifferentiated a tumor becomes, the further it deviates from the normal diploid state. This may be expressed as a tetraploid or aneuploid state according to the amount of DNA on the stained tissue = DNA index of between 1.0 and 2.0. The more cells in the S or DNA phase, the more aggressive the tumor.

2. Favorable prognostic indicators include tumor size of less than 1 cm, a low histologic grade, negative axillary lymph nodes, and positive ER and PR.

3. Fibroplasia and fibroadenophasia are benign conditions.

Interventions

Pretest Patient Care

1. Explain biopsy purpose and procedure. Obtain and record relevant family or personal history of prior biopsy, trauma, recent or current pregnancy, nipple discharge, location of lump, and how lesion was detected. Obtain informed consent.

2. Be aware that open breast biopsies are performed under local or general anesthesia. Sedation may be used with the presence of an anesthetic. NPO is required when general anesthesia is used (see Appendix C).

3. Provide information and support, recognizing the fear the patient experiences about the procedure.

4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. If general anesthesia is used, follow the recovery protocols. See analgesia precautions in Appendix C for conscious sedation.

2. Interpret biopsy outcome and counsel appropriately about possible further testing and treatment (surgery, radiation, and medication [chemotherapy]).

3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Ductal Lavage of Breast Cells (Cytologic) Study; Gail Index of Breast Cancer Risk

Ductal lavage collects cells from the milk ducts of the nipple where most breast cancers begin. If cytologic study shows abnormal cells, this is an indication of increased risk for breast cancer development. Ductal lavage is used to assess breast cancer risk and for ongoing surveillance. A statistical model computes a Gail Index Score in a woman of a given age and with the presence of certain risk factors that indicates risk for developing breast cancer over a specified interval. The Gail Index Score is based on risk factors (eg, late age at menarche, late age at first live birth, number of previous biopsies, and number of first-degree relatives with breast cancer).

Reference Values

Normal No atypical or abnormal cells Gail Index of breast cancer risk = odds ratio = 1.7. For more information, see Gail, Brinton, Byar, et al., 1989.

Procedure
1. Apply a local anesthetic cream to the nipple area using a special kit; use a suction device to draw tiny amounts of fluid droplets from the milk ducts to the nipple surface. These droplets locate the milk ducts' natural opening on the surface of the nipple.

2. Insert a very fine (hair-thin) catheter (Fig. 11.1) into the periareolar duct. Administer local anesthetic into the duct. Use a saline wash to separate the cells. Place the specimen in a special collector vial and send for examination in a biohazard bag.

**FIGURE 11.1** A ductal lavage microcatheter. (Source: CAP Today [College of American Pathologists]; 16 (2), February 2002)

**Clinical Implications**

1. Abnormal findings include atypical hyperplasia and evidence of proliferative breast disease. The presence of atypical cells increases the risk for breast cancer by 4 to 5 times as compared with women who do not have atypical cells.
2. Relative risk is increased even further in presence of a family history of breast cancer (mother, daughter, sister, or two or more close relatives with history of breast cancer), specific genetic change (BRCA1 and BRCA2 mutations), and a Gail Index Score of at least 1.7.
3. The age-specific composite evidence rate of the Gail Model increases rapidly with age, although the conversion model changes little with age.
4. Later relative risk (%) or estimate of developing breast cancer within 10, 20, or 30 years of follow-up is based on projected probability.

**Interventions**

**Pretest Patient Care**

1. Explain the lavage purpose, procedure, benefits, and risks.
2. Be aware that relative high-risk women of any age may be good candidates for ductal lavage. Obtain appropriate history of risk.
3. Describe sensations that might be felt: feelings of fullness, pinching and gently tugging on the breast, uncomfortable, but not usually painful.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results and counsel appropriately about chance of breast cancer development, follow-up, close monitoring (yearly examinations), and preventive drug treatment (eg, tamoxifen) or surgery (oophorectomy or bilateral mastectomy).
2. Remember that test outcomes are interpreted in conjunction with mammogram and physical examination findings.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Liver Biopsy: Cell (Cytologic) and Tissue (Histologic) Study** Liver needle biopsy is an invasive procedure and is done to confirm diagnosis of chronic hepatitis and liver cirrhosis, evaluate disease severity, and establish etiology. Cellular material from the liver may be useful in evaluating the status of the liver in diffuse disorders of the parenchyma and in the diagnosis of space-occupying lesions. Liver biopsy is especially useful when the clinical findings and laboratory test results are not diagnostic (eg, an aspartate aminotransferase [AST] level 10 to 20 times less than the upper defined limit with an alkaline phosphatase [ALP] level less than 3 times the limit) and when the diagnosis or cause cannot be established by other means (enlarged liver of unknown cause or systemic disease affecting the liver, such as miliary tuberculosis). Other indications for liver biopsy include evaluation of chronic hepatitis, portal hypertension, and fever of unknown origin (tuberculosis and brucellosis) and to confirm alcoholic liver disease.

**Reference Values**

**Normal** Negative for malignant or other abnormal cells and abnormal tissue No evidence of local or diffuse liver disease No evidence of toxic reaction to drugs or inflammatory reactions No pathogenic organisms present

**Procedure**

1. See Fine-Needle Aspirates: Cell (Cytologic) and Tissue (Histologic) Study.
2. Be aware that in most cases the patient is hospitalized overnight.
3. Remember that the test may be done at the bedside in a special area, usually under local anesthesia. Obtain
specimens with ultrasound or computed tomography (CT) x-ray guidance and a tissue core biopsy needle, such as the Menghini needle, that provides histologic and cytologic material; or use a fine-needle aspiration needle, which obtains cytologic material only and is useful for cancer diagnosis but not diagnosis of other liver diseases.

4. Place tissue specimens in 10% formalin for fixation. Do not place specimens for culture in a fixative. Check with your laboratory for specific instructions for handling special cases (eg, liver biopsies for copper levels).

5. Express cytology specimens on glass slides and fix immediately in 95% alcohol. Needle rinses may provide helpful diagnostic material as well.

6. See Chapter 1 for safe, effective, informed intratest care. See Chapter 12 on endoscopic examination and liver biopsy.

Clinical Implications Abnormalities in test results of liver biopsies may be helpful in detecting the following liver diseases:

1. Benign disorders, such as those causing liver cirrhosis, and presence of pathogenic organisms in liver abscess
2. Metabolic disorders:
   a. Fatty metamorphosis
   b. Hemosiderosis
   c. Accumulation of bile (hepatitis, obstructive jaundice, malignancy)
   d. Diabetic pathology and Wilson's disease (tissue copper is elevated per dry tissue weight)
   e. Hepatic cysts (congenital or hydatid)
   f. Malignant processes such as end-stage of lymphomas

Interfering Factors The reported effectiveness of liver aspirates or biopsies varies in the limited published information. Because a very small fragment of tissue, often partially destroyed, is taken in a random manner from a large organ, localized disease is easily missed.

1. False-negative results may be caused by:
   a. Sampling error. Detection rate of liver metastases is approximately 50% to 70% with blind biopsy and about 85% (range, 67%–96%) with the use of ultrasound guidance. Also, many diseases produce nonspecific changes that may be spotty, healing, or minimal.
   b. Degeneration or distortion caused by faulty preparation of specimen.

2. False-positive results may be caused by misinterpretation of markedly reactive hepatocytes.

Interventions

Pretest Patient Care

1. Explain the purpose, procedure, benefits, and risks of the test. Obtain properly signed, informed consent. The procedure usually causes minimal discomfort, but only for a short while. Explain that a local anesthetic will be injected into the skin. Remember to ask whether the patient has ever had a reaction to any numbing medicines. Discontinue all aspirin and NSAIDs for at least 7 days before the procedure. PT, partial thromboplastin time (PTT), blood urea nitrogen (BUN), bleeding time, and type L screen cross-match for possible transfusion are usually ordered before biopsy.

2. Ensure that the patient takes nothing by mouth (NPO) for 4 to 6 hours before the procedure. Ask the patient to lie supine with the right arm above the head. During the biopsy, the patient should take a deep breath in, blow the air out, and then hold the breath.

3. Be aware that risks include a small but definite risk for intraabdominal bleeding and bile peritonitis. Percutaneous liver biopsy results in complications in only about 1% of cases.

4. See guidelines in Chapter 1 for safe, effective, informed pretest care.

Clinical Alert

Contraindications include:

1. Bleeding diathesis—anticoagulant therapy
2. Highly vascular lesions
3. Uncooperative patient
4. A prothrombin time in the anticoagulant range, PTT more than 20 seconds over control
5. Severe anemia (Hb < 9.5 g/dL or < 95 g/L) or marked prolonged bleeding time
6. Infection
7. A platelet count of >50,000/mm$^3$ (50 × 10$^9$/L)
8. Marked or tense ascites (risk for leakage)
9. Septic cholangitis

Posttest Patient Aftercare

1. Remember that strict bed rest for at least 6 hours is usually ordered, with observation for 24 hours.
2. Monitor in a recovery area. Assess pulse, blood pressure, and respiration every 15 minutes for the first hour, every 30 minutes for the next 2 hours, once in each of the next 4 hours, and then every 4 hours until the patient's condition is stable.
3. Notify the surgeon if the blood pressure differs markedly from baseline or if the patient is in severe pain.
4. Maintain NPO status for 2 hours; previous diet can then be resumed. Take action immediately if a bleeding episode occurs. Assess for pain and treat as ordered.
5. After 6 hours, a blood specimen for hematocrit testing is usually ordered to rule out internal bleeding. A small number of patients need transfusion for intraperitoneal bleeding.
6. Warn the patient not to cough hard or strain for 2 to 4 hours after the procedure. Heavy lifting and strenuous
activities should be avoided for about 1 week.

7. Follow the guidelines in Chapter 1 for **safe, effective, informed posttest care**.

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**Clinical Alert**

The most common complications include uncontrolled pain, hemorrhage (cause of death from liver biopsy), peritonitis, bile leakage, lacerations of other organs, sepsis, and bacteremia.

**Kidney Biopsy: Cell (Cytologic) and Tissue (Histologic) Study**

Kidney biopsy is used to establish a diagnosis in the presence or renal dysfunction, evaluate severity and extent of disease, guide therapy, and identify candidates for kidney transplantation.

**Reference Values**

**Normal**
- No patterns of abnormality or abnormal glomeruli
- No evidence of drug toxicity, infection, or inflammation

**Procedure**

1. Obtain a specimen of kidney tissue (containing 8–10 glomeruli) by needle biopsy or open surgical technique using x-ray or ultrasound as a guide.
2. Place in normal saline until frozen or place in a fixative or saline and send immediately to the laboratory. Check with your laboratory for specific handling instructions. Proper handling is critical to ensure that the specimen is properly preserved for necessary testing.
3. See section on of **Fine-Needle Aspirates: Cytologic Study** for information regarding obtaining kidney material for cytologic study.

**Clinical Implications**

Abnormal patterns reveal interstitial fibroses and scleroses, diabetic nephrotic pathology syndrome, chronic renal failure, kidney transplant reactions, rejection or failure, past infections, glomerulonephritis, and renal pathology in systemic diseases.

**Interventions**

**Pretest Patient Care**

1. Explain purpose and procedure, benefits, and risks of kidney biopsy.
2. Use sedation and local or general anesthesia if necessary (see Appendix C).
3. Obtain signed, witnessed consent.
4. Be aware that contraindications include uncontrolled bleeding, cancer, large cysts, abscess, pregnancy, acute pyelonephritis, aneurysm, and renal artery.

**Posttest Patient Aftercare**

1. In counsel with other clinicians, counsel patient and interpret test outcomes, discussing further testing and possible treatment.
2. Monitor for complications, which include hematuria (more common in uncontrolled hypertension and uremia), hematomas (presence of a local mass), infection, and laceration of other organs.

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**Clinical Alert**

Death (although very rare) has occurred in 0.12% of patients.

**Respiratory Tract: Cell (Cytologic) and Tissue (Histologic) Study**

The lungs and the passages that conduct air to and from the lungs form the respiratory tract, which is divided into the upper and lower respiratory tracts. The upper respiratory tract consists of the nasal cavities, the nasopharynx, and the larynx; the lower respiratory tract consists of the trachea and the lungs. Sputum is composed of mucus and cells. It is the secretion of the bronchi, lungs, and trachea and is therefore obtained from the lower respiratory tract (bronchi and lungs). Sputum is ejected through the mouth but originates in the lower respiratory tract. Saliva produced by the salivary glands in the mouth is not sputum. A specimen can be correctly identified as sputum in microscopic examination by the presence of dust cells (carbon dust–laden macrophages). Although the glands and secretory cells in the mucous lining of the lower respiratory tract produce up to 100 mL of fluid daily, the healthy person normally does not cough up sputum. Cytologic studies of sputum and bronchial specimens are important as diagnostic aids because of the frequency of cancer of the lung and the relative inaccessibility of this organ. Also detectable are cell changes that may be related to the future development of malignant conditions and to inflammatory conditions.

**Reference Values**

**Normal**
- Negative for abnormal cells or tissue
- No pathogenic organisms

**Procedures**

1. Procedure for obtaining sputum
   a. Be aware that the preferred material is an early-morning specimen. Usually, three specimens are collected on 3 separate days.
   b. Have the patient inhale air to the full capacity of the lungs and then exhale the air with an expulsive deep cough.
   c. Have the patient cough the specimen directly into a wide-mouthed, clean container containing 50% alcohol. (Some cytology laboratories prefer the specimen to be fresh if it will be delivered to the laboratory immediately.) If microbiologic studies are also ordered, the container must be sterile and no fixative should be added.
   d. Cover the specimen with a tight-fitting, clean lid.
   e. Label the specimen with the patient's name, age, date, diagnosis, and number of specimens (one, two, or three) and send immediately to the laboratory.
2. Procedure for obtaining bronchial secretions
   a. Obtain bronchial secretions during bronchoscopy (see Chapter 12). Diagnostic bronchoscopy involves removal of bronchial secretions and tissue for cytologic and microbiologic studies.
b. Collect secretions obtained in a clean container and take to the cytology laboratory. If microbiologic studies are ordered, the container must be sterile.

3. Procedure for obtaining bronchial brushings
   a. Obtain bronchial brushings during bronchoscopy.
   b. Smear the material collected directly on all-frosted slides and immediately fix, or place the actual brush in a container of 50% ethyl alcohol or saline and deliver to the cytology laboratory (check with the laboratory for their preference).

4. Procedures for bronchopulmonary lavage
   a. Use bronchopulmonary lavage to evaluate patients with interstitial lung disease.
   b. Inject saline into the distal portions of the lung and aspirate back through the bronchoscope into a specimen container. This essentially "washes out" the alveoli.
   c. Take the fresh specimen directly to the laboratory. A total cell count and a differential cell count are performed to determine the relative numbers of macrophages, neutrophils, and lymphocytes.

For all procedures, see Chapter 1 guidelines for intratest care.

**Clinical Implications** Abnormalities in sputum and bronchial specimens may sometimes be helpful in detecting the following:

1. Benign atypical changes in sputum, as in:
   a. Inflammatory diseases
   b. Asthma (Curschmann's spirals and eosinophils may be found, but they are not diagnostic of the disease.)
   c. Lipid pneumonia (Lipophages may be found, but they are not diagnostic of the disease.)
   d. Asbestosis (ferruginous or asbestos bodies)
   e. Viral diseases
   f. Benign diseases of lung, such as bronchiectasis, atelectasis, emphysema, and pulmonary infarcts.

2. Metaplasia (the substitution of one adult cell type for another); severe metaplastic changes are found in patients with:
   a. History of chronic cigarette smoking
   b. Pneumonitis
   c. Pulmonary infarcts
   d. Bronchiectasis
   e. Healing abscess
   f. Tuberculosis
   g. Emphysema (Metaplasia often adjoins a carcinoma or a carcinoma in situ.)

3. Viral changes and the presence of virocytes (viral inclusions) may be seen in:
   a. Viral pneumonia
   b. Acute respiratory disease caused by adenovirus
   c. Herpes simplex
   d. Measles
   e. Cytomegalic inclusion disease
   f. Varicella

4. Degenerative changes, as seen in viral diseases of the lung

5. Fungal and parasitic diseases (In parasitic diseases, ova or parasite may be seen.)

6. Tumors (benign and malignant)

**Interfering Factors**

1. False-negative results may be caused by:
   a. Delays in preparation of the specimen, causing a deterioration of tumor cells
   b. Sampling error (Diagnostic cells may not have exfoliated into the material examined.)

2. The frequency of false-negative results is about 15%, in contrast to about 1% in studies for cervical cancer. This high incidence occurs even with careful examination of multiple deep cough specimens.

**Selection of Medications and Media for All Respiratory Cell and Tissue Procedures**

1. Mild sedative and analgesia or local anesthetic (or all) may be used during bronchoscopy. Analgesia is indicated for pain after bronchoscopy. See Chapter 12 for bronchoscopy care and Appendix C for sedative and analgesia precautions.

2. Sputum specimens are collected in a wide-mouthed container; 50% alcohol may be added if transportation to the laboratory will be delayed.

3. Bronchial washings may be collected in a trap tube or wide-mouthed container.

4. Bronchial brushes may be smeared directly on glass slides, which are then fixed immediately in 95% alcohol or spray fixative. Brushes may be placed in a fixative solution such as 50% alcohol.

**Clinical Alert**

The uncooperative patient is a contraindication.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the test. Tell the patient not to drink fixative liquid in specimen container.

2. Emphasize that sputum is not saliva. If a patient is having difficulty producing sputum, a hot shower before obtaining a specimen may improve the yield.

3. Advise the patient to brush the teeth and rinse the mouth well before obtaining the sputum specimen to avoid
If a bronchoscopy is performed, maintain NPO for 6 hours before the procedure. Abnormal results in studies of the pancreas, gallbladder, and duodenum may reveal malignant cells (usually adenocarcinoma). Granulomatous inflammations seen in chronic gastritis and sarcoidosis of the stomach, which is characterized by giant epithelial cells and folic acid deficiencies, characterized by giant epithelial cells. A pernicious anemia is caused by vitamin B₁₂ deficiencies, characterized by giant epithelial cells. Ulcerative colitis, characterized by hyperchromatic nuclei surrounded by a thin cytoplasmic rim. Malignant cells (usually adenocarcinoma) may be recognized because of their characteristic cellular changes. Response to radiation may also be noted from cytologic studies.

**Gastrointestinal Tract: Cell (Cytologic) and Tissue (Histologic) Study** Exfoliative cytology of the gastrointestinal tract is useful in the diagnosis of benign and malignant diseases. It is not, however, a specific test for these diseases. Many benign diseases, such as leukoplakia of the esophagus, esophagitis, gastritis, pernicious anemia, and granulomatous diseases, may be recognized because of their characteristic cellular changes. Response to radiation may also be noted from cytologic studies.

**Reference Values**

| Normal Negative for abnormal cells | Squamous epithelial cells of the esophagus may be present. |

**Procedure**

1. Give a sedative before the procedure. For esophageal studies, pass a nasogastric Levin tube approximately 40 cm (to the cardioesophageal junction) with the patient in a sitting position.
2. For stomach studies, pass a Levin tube into the stomach (approximately 60 cm) with the patient in a sitting position.
3. For pancreatic and gallbladder drainage, pass a special double-lumen gastric tube orally 45 cm, with the patient in a sitting position. Then place the patient on his or her side and pass the tube slowly 8.5 cm. It takes about 20 minutes for the tube to reach this distance. Confirm the tube location by biopsy. Lavage with physiologic salt solution is done during all upper gastrointestinal cytology procedures.
4. Be aware that specimens can also be obtained during endoscopy procedures.
5. Remember that material obtained with the use of brushes may be smeared directly on glass slides, which are fixed immediately in 95% alcohol or spray fixative. Brushes may also be placed in a fixative such as 50% alcohol. See Chapter 12 for endoscopic biopsy procedures. Washings must be delivered immediately to the laboratory and may need to be placed on ice. Check with your individual laboratory for specific instructions on handling of washings from the gastrointestinal tract.

**Clinical Implications**

1. The characteristics of benign and malignant cells of the gastrointestinal tract are the same as for cells of the rest of the body.
2. Abnormal results in cytologic studies of the esophagus may be a nonspecific aid in the diagnosis of:
   a. Acute esophagitis, characterized by increased exfoliation of basal cells with inflammatory cells and polymorphonuclear leukocytes in the cytoplasm of the benign squamous cells.
   b. Vitamin B₁₂ and folic acid deficiencies, characterized by giant epithelial cells.
   c. Malignant diseases, characterized by typical cells of esophageal malignancy.
3. Abnormal results in studies of the stomach may be a nonspecific aid in the diagnosis of:
   a. Pernicious anemia, characterized by giant epithelial cells. An injection of vitamin B₁₂ causes these cells to disappear within 24 hours.
   b. Granulomatous inflammations seen in chronic gastritis and sarcoidosis of the stomach, which is characterized by granulomatous cells.
   c. Gastritis, characterized by degenerative changes and an increase in the exfoliation of clusters of surface epithelial cells.
   d. Malignant diseases, most of which are gastric adenocarcinomas. Lymphoma cells can be differentiated from adenocarcinoma. The Reed-Sternberg cell, a multinucleated giant cell, is the characteristic cell found along with abnormal lymphocytes in Hodgkin's disease.
4. Abnormal results in studies of the pancreas, gallbladder, and duodenum may reveal malignant cells (usually adenocarcinoma), but it is sometimes difficult to determine the exact site of the tumor.
5. Abnormal results in examination of the colon may reveal:
   a. Ileitis, characterized by large, multinucleated histocytes (Bovine tuberculosis commonly manifests itself in this area.)
   b. Ulcerative colitis, characterized by hyperchromatic nuclei surrounded by a thin cytoplasmic rim.
   c. Malignant cells (usually adenocarcinoma).

**Interfering Factors** The barium and lubricant used in Levin tubes interfere with good results because they distort the cells and prevent accurate evaluation.

**Interventions**

**Pretest Patient Care**

1. If the specimen is obtained by bronchoscopy, check the patient's blood pressure and respirations every 15 minutes for 1 hour, then every 2 hours for 4 hours, then as ordered.
2. Assist and teach the patient to not eat or drink until the gag reflex returns.
3. Maintain bed rest and elevate the head of the bed 45 degrees.
4. Manage pain as indicated.
5. Auscultate the chest for breath sounds every 2 to 4 hours and then as ordered.
6. Perform postural drainage and oropharyngeal suctioning as ordered. (Refer to bronchoscopy care in Chapter 12.)
7. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.
8. Interpret test outcomes and provide support for abnormal outcomes.

**Posttest Patient Aftercare**

1. If the specimen is obtained by bronchoscopy, check the patient's blood pressure and respirations every 15 minutes for 1 hour, then every 2 hours for 4 hours, then as ordered.
2. Manage pain with sedation as indicated.
3. Provide emotional support.
4. Remember that material obtained with the use of brushes may be smeared directly on glass slides, which are fixed immediately in 95% alcohol or spray fixative. Brushes may also be placed in a fixative such as 50% alcohol. See Chapter 12 for endoscopic biopsy procedures. Washings must be delivered immediately to the laboratory and may need to be placed on ice. Check with your individual laboratory for specific instructions on handling of washings from the gastrointestinal tract.

**Interfering Factors** The barium and lubricant used in Levin tubes interfere with good results because they distort the cells and prevent accurate evaluation.
1. Tell the patient the purpose of this test, the nature of the procedure, and to anticipate some discomfort.
2. Be aware that a liquid diet usually is ordered for the 24 hours before testing. Encourage the patient to take fluids throughout the night and in the morning before the procedure.
3. Do not administer oral barium for the preceding 24 hours.
4. Remember that laxatives and enemas are ordered for colon cytologic studies.
5. Because insertion of the nasogastric tube can cause considerable discomfort, devise a system (eg, raising a hand) to indicate discomfort with the patient. (See gastric analysis procedure in Chapter 16.)
6. Inform the patient that panting, mouth breathing, or swallowing can help to ease insertion of the tube.
7. Tell patient that sucking on ice chips or sipping through a straw also makes insertion of the tube easier.
8. Remember that ballottement and massage of the abdomen are needed to release cells when a gastric wash technique is used.
9. See Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert

1. The uncooperative patient is a contraindication.
2. Immediately remove the tube if the patient shows signs of distress: coughing, gasping, or cyanosis.

Posttest Patient Aftercare

1. Interpret test results and monitor appropriately. The patient should be given food, fluids, and rest after the tests are completed.
2. Provide rest. Patients having colon studies will be feeling quite tired.
3. Be aware that potential complications of endoscopy include respiratory distress and esophageal, gastric, or duodenal perforation. Complications of proctosigmoidoscopy include possible bowel perforation. Decreased blood pressure, pallor, diaphoresis, and bradycardia are signs of vasovagal stimulation and require immediate notification of the physician.
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Papanicolaou (Pap) Smear: Cell (Cytologic) Study of the Female Genital Tract, Vulva, Vagina, and Cervix; DNA Test for Human Papilloma Virus (HPV) Characteristic physiologic cellular changes occur in the genital tract from birth through the postmenopausal years. Three major cell types occur in a characteristic pattern in normal vaginal smears:

1. Superficial squamous cells (mature squamous, usually polygonal, containing a pyknotic [thick, compact, dense] nucleus)
2. Intermediate squamous cells (mature squamous, usually polygonal, containing a clearly structured vesicular nucleus, which may be either well preserved or changed as a result of bacterial cytolysis)
3. Parabasal cells (immature squamous, usually round or oval, containing one or, rarely, more than one) as a result of degeneration or necrosis

Findings indicate that presence of human papilloma virus (HPV) may be associated with the development of cervical cancer.

Clinical Alert

Deviation from normal physiologic cell patterns may be indicative of a pathologic condition.

The Papanicolaou (Pap) cytologic smear is used principally for early detection of cervical cancer and diagnosis of precancerous and cancerous conditions of the vulva and vagina. This test is also used for diagnosis of inflammatory and infectious diseases. Because the Pap smear is of great importance in the early detection of cervical cancer, it is recommended that all women older than 20 years of age have the test at least once a year. HPV testing is recommended for all women with a Pap cytologic diagnosis of atypical squamous cells of undetermined significance (ASCUS). The value of the Pap smear depends on the fact that cells readily exfoliate (or can be easily stripped) from genital cancers. Cytologic study can also be used for assessing response to administered sex hormones. The microbiologic examination on cytology samples is not as accurate as bacterial culture, but it can provide valuable information. Specimens for cytologic examination of the genital tract are usually obtained by vaginal speculum examination or by colposcopy with biopsy. Material from the cervix, endocervix, and posterior fornix is obtained for most smears. Smears for hormonal evaluation are obtained from the vagina.

Clinical Alert

1. Cytologic findings alone do not form the basis of a diagnosis of cancer or other diseases. Often they are used to justify further procedures, such as biopsy.
2. The Pap smear has been FDA approved for diagnosis of cancer and human papilloma virus (HPV).
3. The Bethesda System—2001 recommends deleting hormonal evaluation. Hormonal evaluation is a crude measure of estrogen-like effects on squamous cells. It is not reproducible and does not correlate with symptoms or plasma estrogen levels (Stone, 1975).

In an effort to standardize reporting of cervical-vaginal cytology specimens, the Bethesda System for reporting cervical-vaginal diagnoses was developed by a 1977 National Cancer Institute workshop and slightly modified after a second workshop in 1991 and revised in 2001. This reporting system is being adapted by numerous laboratories nationwide. The terminology of this reporting system appears in Table 11.2.

Table 11.2 Bethesda System for Reporting Cervical-Vaginal Diagnoses

<table>
<thead>
<tr>
<th>Bethesda System—2001 (Specimen type:</th>
<th>Bethesda System—1991 Conventional [Pap] vs. liquid vs. other)</th>
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</thead>
<tbody>
<tr>
<td>Conventional [Pap] vs. liquid vs. other</td>
<td></td>
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</table>
**SPECIMEN ADEQUACY**

- Satisfactory for evaluation
- Satisfactory but limited by [describe limitation]
- Unsatisfactory
  - Unsatisfactory for evaluation (specify reason):
    1. Specimen rejected but processed (specify reason)
    2. Specimen processed and examined but unsatisfactory for evaluation of epithelial abnormality because of (specify reason)

**GENERAL CATEGORIZATION**

- Within normal limits
- Epithelial cell abnormality (followed by interpretation)
- Benign cellular changes
- Other

**EPITHELIAL CELL ABNORMALITIES**

**SQUAMOUS CELL TYPE**

- Atypical squamous cells of undetermined significance (ASC-US), qualify
- Low-grade squamous intraepithelial lesion (LSIL)
- HSIL
- HSIL encompassing moderate, severe, carcinoma in situ (CIS)
- Squamous cell carcinoma

**GLANDULAR CELL LESIONS**

- Endometrial cells, cytologically benign in a postmenopausal woman
- ASCUS, qualify
- Endocervical adenocarcinoma
- Endometrial adenocarcinoma
- Extrapuerine adenocarcinoma
- Adenocarcinoma NOS (not otherwise specified)

**INTERPRETATION/RESULT**

- Within normal limits
- Negative for intraepithelial lesion or malignancy
- Organisms:
  1. Trichomonas vaginalis
  2. Fungal organisms morphologically consistent with Candida species
  3. Shift in flora suggestive of bacterial vaginosis (coccobacillus)
  4. Bacteria morphologically consistent with Actinomyces species
  5. Cellular changes consistent with herpes simplex virus
- Other nonneoplastic findings:
  1. Reactive changes associated with inflammation (includes repair), radiation, IUD, atrophy, glandular cells status after hysterectomy, or endometrial cells (in women >40 years of age)

**SQUAMOUS CELL TYPE**

- Squamous cell:
  - ASCUS, cannot exclude high-grade squamous intraepithelial lesion (HSIL)(ASC-H)
  - LSIL encompassing human papilloma virus (HPV), mild dysplasia, cervical intraepithelial neoplasm (CIN) grade 1 (low-grade precursor)
  - HSIL encompassing moderate, severe, CIS/CIN 2 and CIN 3 (grades 2 and 3 are high-grade precursor)
- Squamous cell carcinoma

**GLANDULAR CELL LESIONS**

- Atypical
  1. Endocervical cells (NOS or specify in comments)
  2. Endometrial cells (NOS or specify in comments)
  3. Glandular cells (NOS or specify in comments)
  4. Endocervical cells, favor neoplastic
  5. Glandular cells, favor neoplastic
- Endocervical adenocarcinoma in situ
  1. Adenocarcinoma
  2. Endometrial
  3. Extrapuerine
  4. NOS (not otherwise specified)

The AutoPap System received preliminary approval from the U.S. Food and Drug Administration in early 1998 and is the first device of its kind to receive a recommended approval for automated initial Pap smear screening. With the AutoPap System, approximately 25% of submitted Pap smears would receive AutoPap review only and would not need to be seen by a technologist. Cytyc has taken a different approach to create a better Pap smear: ThinPrep. The Pap smear collection device for ThinPrep is rinsed in a special solution (ie, PreservCyt) and sent to the lab. A special machine prepares a uniform monolayer Pap smear. These slides are then manually screened in the usual manner. Studies have shown that these ThinPrep smears have a higher rate of detection of biopsy-proven high-grade lesions and a lower rate of false-negative results than conventional Pap smears. Human papilloma virus (HPV) has been identified as the primary causal factor in cervical cancer. The Digene Hybrid Capture HPV Test is approved in the U.S. for HPV DNA detection. Primarily, this test is useful to triage or manage women with an ASCUS or equivocal cytology result. It is an efficient, rapid test that is able to differentiate patients with high-risk versus low-risk HPV and can be performed from the same patient specimen when the ThinPrep Pap Test is used. In addition, the FDA has approved testing for Chlamydia trachomatis and Neisseria gonorrhoeae directly from the ThinPrep sample vial. If the ThinPrep Pap Test is not used, a collection kit is available from Digene. Clinicians should check with their laboratories for ordering and collection instructions for any of these tests.

**Reference Values**

- **Normal Pap** No abnormal or atypical cells No inflammation, no infection, no partially obscuring blood Major cell types within normal limits Negative for intraepithelial cell abnormality of malignancy Negative for HPV

**Procedure**

1. Ask the patient to remove clothing from the waist down.
2. Place the patient in a lithotomy position on an examining table.
Gently insert an appropriately sized bivalve speculum, lubricated and warmed only with water, into the vagina to expose the cervix.

Observe standard universal precautions (see Appendix A).

If a conventional Pap smear, as opposed to liquid base, is being taken, scrape the posterior fornix and the external os of the cervix with a wooden spatula, a cytobrush, or a cytobroom. Smear material obtained on glass slides and place immediately in 95% alcohol or spray fixative before air-drying occurs.

If a ThinPrep Pap smear is being taken, use a broomlike collection device. Insert the central bristles of the broom into the endocervical canal deep enough to allow the short bristles to contact the ectocervix fully. Push gently and rotate the broom in a clockwise direction five times. Rinse the broom with a PreservCyt solution vial by pushing the broom into the bottom of the vial 10 times, forcing the bristles apart. As a final step, swirl the broom vigorously to release material. Discard the collection device. Tighten the cap on the solution container so that the torque line on the cap passes the torque line on the vial.

Label the specimen properly with the patient's name and identifying number (if appropriate) and the area from which the specimen was obtained, and send it to the laboratory with a properly completed information sheet, including date of collection, patient's date of birth, date of last menstrual period, and pertinent clinical history.

Examination takes about 5 minutes.

See Chapter 1 guidelines for intratest care.

Clinical Alert

1. The best time to take a Pap smear is 2 weeks after the first day of the last menstrual period, definitely not when the patient is menstruating or bleeding, unless bleeding is a continuous condition.
2. Cytologic specimens should be considered infectious until fixed with a germicidal fixative. Observe standard precautions when handling specimens from all patients.

Clinical Implications

1. Abnormal Pap cytologic responses include atypical squamous cells of undetermined significance (ASCUS) and can be classified as protective, destructive, reparative (regenerative), or neoplastic.
2. Inflammatory reactions and microbes (Trichomonas vaginalis and Monilia, Coccobacilla, Candida, and Actinomyces species, cells indicative of herpes simplex virus [HSV]) can be identified to help in the diagnosis of vaginal diseases, and evidence of Chlamydia trachomatis and Neisseria gonorrhoeae.
3. Reactive cells associated with inflammation, typical surgical repair, radiation, intrauterine contraception devices (IUDs), post-hysterectomy glandular cells, atrophy, and endometrial cells in a woman 40 years of age or older
4. Positive DNA test for HPV
5. Precancerous and cancerous lesions of the cervix can be identified.

Clinical Alert

1. A cytobrush should not be used to obtain a cervical specimen from a pregnant patient.
2. Some nonpregnant patients experience heavy bleeding after a cytobrush is used.

Interfering Factors

1. Medications such as tetracycline and digitalis, which affect the squamous epithelium, alter test results.
2. The use of lubricating jelly in the vagina or recent douching interferes with test results by distorting the cells and preventing accurate evaluation.
3. Heavy menstrual flow and blood may make the interpretation of the results difficult and may obscure atypical cells.

Interventions

Pretest Patient Care

1. Explain the Pap cytology test purpose and procedure. In rape cases, vaginal swabs for forensic evidence require a chain-of-custody protocol (see Appendix L).
2. Instruct the patient not to douche for 2 to 3 days before the test because douching may remove the exfoliated cells.
3. Instruct the patient not to use vaginal medications or vaginal contraceptives during the 48 hours before the examination. Intercourse is not recommended the night before the examination.
4. Have the patient empty bladder and rectum before examination.
5. Ask the patient to give the following information:
   a. Age—indicate if adolescent, pregnant, or postmenopausal
   b. Use of hormone therapy, birth control pills, or contraceptive devices
   c. Past vaginal surgical repair or hysterectomy
   d. All medications taken, including prescribed, over-the-counter, and herbal medications
   e. Any radiation therapy
   f. Any other pertinent clinical history (eg, previous abnormal Pap smear, signs of inflammation or bleeding)
6. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Give the patient a perineal pad after the procedure to absorb any bleeding or drainage.
2. Interpret test results and counsel appropriately regarding repeat cytology testing if atypical or abnormal cells are present.
3. Explain that monitoring and management of women with atypical or abnormal cells follows ASCCP consensus guidelines. Repeat HPV DNA tests and repeat cytology Pap smears are standard.
4. Counsel that treatment may include intravaginal estrogen therapy, diagnostic excisional procedures, and/or referral to an expert. Management options may vary if the woman is an adolescent, pregnant, or postmenopausal (see Table 11.2, the Bethesda System, and Chapter 12, Endoscopic Studies for typical procedures).
5. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Further Testing After Abnormal Results of Pap Testing**

<table>
<thead>
<tr>
<th>Adolescents—LSIL</th>
<th>Postmenopausal Women—ASCUS</th>
<th>Women With HSIL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Repeat Pap at 6 and 12 months.</td>
<td>Intravaginal estrogen therapy</td>
<td>Colposcopy with endocervical assessment</td>
</tr>
<tr>
<td>Colposcopy if repeat Pap is ASC or above, or HPV DNA at 12 months</td>
<td>After treatment, repeat PAP cytology in one week.</td>
<td>If no CIN, review cytology, colposcopy, and histology. If necessary, a revised report is issued.</td>
</tr>
<tr>
<td>If positive for high-risk HPV, refer for colposcopy.</td>
<td>If Pap is negative, repeat at 4–6 months.</td>
<td>If no change found upon review, biopsy to confirm CIN.</td>
</tr>
<tr>
<td>If positive, perform another colposcopy</td>
<td>If repeat is negative, return to regular screening schedule. If either Pap is ASC or above, refer for colposcopy.</td>
<td>Manage and treat per ASCCP.</td>
</tr>
</tbody>
</table>

**Footnote**


**Anal Smears: Cell (Cytologic) Study**

The incidence of anal squamous neoplasms has been increasing, especially in homosexual and bisexual men and in women with multicentric genital tract squamous lesions. When evaluating high-risk populations, the rate of anal cancer has been reported to be as high as 70 cases per 100,000. The etiology and pathogenesis of anal squamous neoplasia are similar to that of cervical squamous neoplasia, including an association with human papilloma virus, which has been identified in 90% of anal squamous cancers in reported studies.

**Clinical Alert**

Anoscopic and histologic assessment of anal lesions is critical to classify lesions accurately. Any cytologic abnormality should be followed up with high-resolution anoscopy and any lesion should be biopsied to confirm the grade of dysplasia.

While there are no official guidelines regarding anal cytology screening for anal squamous intraepithelial lesions (ASIL), smears of the anorectal junction are being done with increasing frequency on high-risk patients. Taking an anal “Pap” is a fairly simple procedure, and samples are handled in a similar fashion to cervical/vaginal Pap smears. Clinicians should check with their laboratories for specific handling instructions.

**Reference Values**

**Normal** Negative for intraepithelial cell abnormality or malignancy Negative for HPV

**Procedure**

1. Ask the patient to remove clothing from the waist down.
2. Place the patient on the side with the knees drawn up to the chest.
3. Gently insert a Dacron swab or cytobrush into the anus to a distance of 2 to 3 cm, ensuring sampling of the anorectal junction by passing and including the dentate line.
4. Rotate the swab or cytobrush 360 degrees while gently pulling back and forth.
5. Transfer the sample by inserting the swab or brush into a vial of fixative fluid and gently agitate, or if the laboratory prefers, directly apply the sample to a glass slide, which is then placed in 95% alcohol or spray fixed.
6. Seal the sample vial in a biohazard bag and forward to the laboratory with a properly completed requisition.
7. See Chapter 1 guidelines for intratest care.

**Clinical Implications** Abnormal results are indicative of abnormal cytology, anal squamous intraepithelial lesions (ASIL)
and malignancy.

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose of the test and the collection procedure. No rectal suppositories before day of obtaining smear.
2. Advise that there may be a slight discomfort (eg, pressure sensation) during insertion and rotation of swab.
3. Follow guidelines in Chapter 1 for safe, effective, and informed pretest care.

**Posttest Patient Care**

1. Give the patient a perineal pad after the procedure to absorb any bleeding or drainage.
2. Interpret test results and counsel appropriately regarding subsequent testing (anoscopy and biopsy) if an abnormal result is received and possible need for treatment (ie, excisional procedures).
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Aspirated Breast Cysts and Nipple Discharge: Cell (Cytologic) Study, Fine Needle Aspiration (FNA) and Cytologic Study of Breast Aspirate and Biomarkers of Cancer Risk**

Nipple discharge usually is normal only during the lactation period. Any other nipple discharge is abnormal, and when it occurs, the breasts should be examined for mastitis, duct papilloma, and intraductal cancer. (However, certain situations increase the possibility of finding a normal nipple discharge, such as pregnancy, perimenopausal state, and use of birth control pills.) About 3% of breast cancers and 10% of benign lesions of the breast are associated with abnormal nipple discharge. The contents of the identified breast cyst are obtained by fine-needle aspiration biopsy and are examined to detect malignant cells. Fine needle periareolar breast aspiration, along with the Gail Risk Model and certain biomarkers, is used to predict cancer development in high-risk individuals.

**Reference Values**

**Normal** Negative for neoplasia or hyperplasia with atypia No evidence of high-risk results No expression of select biomarkers as predictors of future cancer development

**Procedure for Nipple Discharge**

1. Limit this procedure to patients who have no palpable masses in the breast or other evidence of breast cancer.
2. Wash the nipple with a cotton pledget and pat dry.
3. Gently strip, or milk, the nipple to obtain a discharge. Express fluid until a pea-sized drop appears. The patient may assist by holding a bottle of fixative beneath the breast so that the slide may be dropped in immediately.
4. Spread the nipple discharge directly on glass slides and then drop into the fixative bottle containing 95% alcohol or spray fixed.
5. Identify the specimen with pertinent data, including from which breast it was obtained, and send without delay to the laboratory.
6. For all procedures, see Chapter 1 guidelines for intrastest care.

**Procedure for FNA**

1. Administer buffered lidocaine (1%) as a local anesthetic. Use a 1.5 inch-21 GA needle with attached 10–12 mL syringe pre-wetted with tissue culture medium. Position needle directly adjacent to areola, avoiding superficial blood vessels. A number of aspirations may be performed in upper, outer, and inner quadrants of breast.
2. All cells if from multiple aspirations may be pooled in 5 mL of an ice-cold medium in an ice bath and fixed in acetone, methanol, and/or formalin until stained. Part of specimen is used for cytology (cell) study, the rest for expression of biomarkers.
3. Use sterile measures and standard precautions.

**Clinical Implications** Abnormal results are helpful in identifying:

1. Benign breast conditions, such as mastitis or intraductal papilloma
2. Malignant breast conditions, such as intraductal cancer or intracystic infiltrating cancer
3. FNA results of hyperplasia with atypia are associated with a greater risk of future development of breast cancer.
4. Expression of DNA aneuploidy (=2+ intensity), p53 expression (=2+ intensity), HER2/ neu expression (=2+ intensity), nER expression (=1+ intensity), and EGF-R expression (=2+ intensity).
5. Also see breast biopsy prognostic markers and ER, PR, and DNA ploidy.

**Interfering Factors** Use of drugs that alter hormone balance (eg, phenothiazines, digitalis, diuretics, steroids) often results in a clear nipple discharge.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the nipple discharge procedure. Oral lorazepam may be given for anxiety.
2. The nipple should be washed with a cotton pledget and patted dry.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. No special instructions are needed for nipple discharge aftercare because this is not an invasive procedure. The patient should be instructed to contact the clinician.
2. Interpret test results and counsel appropriately about possible further testing (eg, biopsy) and treatment (eg, tamoxifen, which reduces breast cancer risk, or antibiotics for infection).
3. After FNA, monitor for hematoma formation and infection. Apply cold packs for approximately 10 minutes, bind breast and chest wall with gauze, and instruct patient to wear a tight-fitting sports bra.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Clinical Alert**

1. Any nipple discharge, regardless of color, should be reported and examined. A bloody or blood-tinged discharge is especially significant.
2. After FNA, a large hematoma may require surgery, and infection may require antibiotics.

**Urine: Cell (Cytologic) Study**

Cells from the epithelial lining of the urinary tract exfoliate readily into the urine. Urine cytology is most useful in the diagnosis of cancer and inflammatory diseases of the bladder, the renal pelvis, the ureters, and the urethra. This study is also valuable in detecting cytomegalic inclusion disease and other viral diseases and in detecting bladder cancer in high-risk populations, such as workers exposed to aniline dyes, smokers, and patients previously treated for bladder cancer. A Pap stain of smears prepared from the urinary sediment, filter preparations, or cytocentrifuged smears is useful to identify abnormalities.

**Reference Values**

**Normal** Negative Epithelial and squamous cells are normally present in urine. (See also Chapter 3, especially Microscopic Examination of Urine Sediment.)

**Procedure**

1. Obtain a clean-voided urine specimen of at least 180 mL for an adult or 10 mL for a child.
2. Obtain a catheterized specimen, if possible, if cancer is suspected.
3. Deliver the specimen immediately to the cytology laboratory. Urine should be as fresh as possible when it is examined. If a delay is expected, an equal volume of 50% alcohol may be added as a preservative.
4. Collect urine specimens or bladder washings in wide-mouthed containers; add 50% alcohol if laboratory transport will be delayed. Check with your laboratory for specific instructions.
5. See Chapter 1 guidelines for intratest care.

**Clinical Implications**

1. Findings possibly indicative of inflammatory conditions of the lower urinary tract include:
   a. Epithelial hyperplasia
   b. Atypical cells
   c. Abundance of red blood cells
   d. Leukocytes
2. Findings indicative of viral disease include the following:
   a. Cytomegalic inclusion disease: large intranuclear inclusions
      1. Cytomegaloviruses or salivary gland viruses are related to the herpes varicella agents.
      2. Infected people may excrete virus in the urine or saliva for months.
      3. About 60% to 90% of adults have experienced infection.
      4. In closed populations (eg, institutionalized mentally disabled persons, household contacts), high infection rates may occur at an early age.
   b. Measles: characteristic cytoplasmic inclusion bodies may be found in the urine before the appearance of Koplik's spots.
3. Findings possible indicative of malacoplakia and granulomatous disease of the bladder or upper urinary tract include:
   a. Histocytes with multiple granules in an abundant, foamy cytoplasm
   b. Michaelis-Gutmann bodies in malacoplakia
4. Cytologic findings possibly indicative of malignancy. If the specimen shows evidence of any of the changes associated with malignancy, cancer of the bladder, renal pelvis, ureters, kidney, or urethra may be suspected. Metastatic tumor should be ruled out as well.

**NOTE**

Inflammatory conditions could be caused by benign prostatic hyperplasia, adenocarcinoma of the prostate, kidney stones, diverticula of bladder, strictures, or malformations.

**NOTE**

Cytomegalic inclusion disease is a viral infection that usually occurs in childhood but is also seen in cancer patients treated with chemotherapy and in transplantation patients treated with immunosuppressive drugs. The renal tubular epithelium is usually involved.

**Interventions**

**Pretest Patient Care**

1. Be aware that patient preparation depends on the type of procedure being done. Explain the purpose, procedure, benefits, and risks to the patient.
2. If cystoscopy is done, give the patient anesthesia (general, spinal, or local). Refer to Chapter 12 for cystoscopy care.
3. If voided urine is required, instruct the patient in the procedure for collection of a clean-catch specimen.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Clinical Alert**

The only contraindication is an uncooperative patient.

**Posttest Patient Aftercare**

1. Interpret test results and monitor appropriately. If cystoscopy is performed gently and with adequate lubrication, the patient should experience only minimal discomfort after the procedure.
2. Be aware that aftereffects may include mild dysuria and transient hematuria, but these should disappear within 48 hours after the procedure. The patient should be able to void normally after a routine cystoscopic examination. Refer to Chapter 12 for cystoscopy care.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Cerebrospinal Fluid (CSF): Cell (Cytologic) Study**

CSF obtained by lumbar puncture is examined for the presence of abnormal cells and for an increase or decrease in the normally present cell population. Most of the usual laboratory procedures for study of CSF involve an examination of the leukocytes and a leukocyte count; chemical and microbiologic studies are also done. Cell studies of the CSF also have been used to identify neoplastic cells. These studies have been especially helpful in diagnosis and treatment of the different phases of leukemia. The nature of neoplasia is such that for tumor cells to exfoliate, they must actually invade the CSF circulation and enter such areas as the ventricle wall, the choroid plexus, or the subarachnoid space.

**Reference Values**

**Normal**

- Total cell count, adult: 0–10/mm³ or 0–10 × 10⁹ cells/L (all mononuclear cells)
- Total cell count, infant: 0–20/mm³ or 0–20 × 10⁹ cells/L

Negative for neoplasia

A variety of normal cells may be seen. Large lymphocytes are most common. Small lymphocytes are also seen, as are elements of the monocyte/macrophage series. The CSF of a healthy person should be free of all pathogens. Negative for blood

**Procedure**

1. Obtain four specimens of at least 1 to 3 mL each by lumbar puncture (see Chapter 5).
2. Remember that generally, only one specimen of 1 to 3 mL goes to the cytology laboratory. Other tubes are sent to different laboratories for examination.
3. Label the specimen with the patient's name, date, and type of specimen.
4. Send the sample immediately to the cytology laboratory for processing.

**Clinical Alert**

The laboratory should be given adequate warning that a CSF specimen is being delivered. Time is a crucial factor; cells begin to disintegrate if the sample is kept at room temperature for more than 1 hour.

**Clinical Implications**

1. CSF abnormalities may indicate:
   a. Malignant gliomas that have invaded the ventricles or cortex of the brain: leukocytes, 150/mm³ or 150 × 10⁹ cells/L (The samples may be normal in 75% of patients.)
   b. Ependymoma (neoplasm of differentiated ependymal cells) and medulloblastoma (a cerebellar tumor) in children
   c. Seminoma and pineoblastoma (tumors of the pineal gland)
   d. Secondary carcinomas:
      1. Secondary carcinomas metastasizing to the central nervous system have multiple avenues to the subarachnoid space through the bloodstream
      2. The breast and lung are common sources of metastatic cells exfoliated in the CSF. Infiltration of acute leukemia is also common.
   e. Central nervous system leukemia
   f. Fungal forms:
      1. Congenital toxoplasmosis: leukocytes, 50 to 500/mm³ or 50–500 × 10⁹ cells/L (mostly monocytes present)
      2. Coccidioidomycosis: leukocytes, 200/mm³ (200 × 10⁹ cells/L)
   g. Various forms of meningitis:
      1. Cryptococcal meningitis: leukocytes, 800/mm³ or 800 × 10⁹ cells/L (lymphocytes are more abundant than polynuclear neutrophilic leukocytes)
      2. Tuberculous meningitis: leukocytes, 25 to 1000/mm³ or 25–1000 × 10⁹ cells/L (mostly lymphocytes present)
      3. Acute pyogenic meningitis: leukocytes, 25 to 1000/mm³ or 25–1000 × 10⁹ cells/L (mostly polynuclear neutrophilic leukocytes present)
   h. Meningoencephalitis (primary amebic meningoencephalitis):
      1. Leukocytes, 400 to 21,000/mm³ (400–21,000 × 10⁹ cells/L)
      2. Red blood cells are also found.
      3. Wright's stain may reveal amebas.
   i. Hemosiderin-laden macrophages, as in subarachnoid hemorrhage
   j. Lipophages from central nervous system destructive processes
2. Characteristics of neoplastic cells:
   a. Sometimes marked increase in size, most likely sarcoma and carcinoma
   b. Exfoliated cells tend to be more polymorphic as the neoplasm becomes increasingly malignant
Interfering Factors The lumbar puncture can occasionally cause contamination of the specimen with squamous epithelial cells or spindly fibroblasts.

Interventions

Pretest Patient Care

1. Explain the procedure to the patient (see Chapter 5). A local anesthetic will be used. Remember to ask whether the patient has a history of reacting to local anesthetic. CSF is collected in tubes and delivered immediately to the laboratory. No fixative is added to the specimen. Instruct the patient that the procedure may be uncomfortable and that immobilization is extremely important. The patient should be instructed to breathe normally and not to hold the breath. Provide the patient with physical and emotional support during the procedure.
2. See guidelines in Chapter 1 for safe, effective, informed pretest care.

Clinical Alert

The only contraindication is an uncooperative patient.

Posttest Patient Aftercare

1. Place the patient in a supine position. Keep the head of the bed flat for 4 to 8 hours as ordered; if headache occurs, elevate the feet 10 to 15 degrees above the head. Assist and teach the patient to turn and deep breathe every 2 to 4 hours. Blood pressure, pulse, and respiration should be checked every 15 minutes four times, then every hour four times, then as ordered. Control pain as ordered and observe the site of puncture for redness, swelling, or drainage; report any symptoms to physician.
2. Interpret test outcomes and monitor appropriately.
3. Follow guidelines in Chapter 1 for safe, effective, and informed posttest care.

Effusions (Thoracentesis and Paracentesis): Cell (Cytologic) Study

Effusions are accumulations of fluids. They may be exudates, which generally accumulate as a result of inflammation (tuberculosis, abscess, pancreatitis), lung infarct or embolus, trauma, systemic lupus erythematosus (SLE), or transudates, which are fluids not associated with inflammation (ie, cirrhosis, congestive heart failure, and nephrotic syndromes). Table 11.4 compares these two effusions.

<table>
<thead>
<tr>
<th>Exudate</th>
<th>Transudate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Accumulates in body cavities and tissues because of malignancy or inflammation</td>
<td>1. Accumulates in body cavities from impaired circulation</td>
</tr>
<tr>
<td>2. Associated with an inflammatory process</td>
<td>2. Not associated with an inflammatory process</td>
</tr>
<tr>
<td>3. Viscous; opaque to purulent</td>
<td>3. Highly fluid</td>
</tr>
<tr>
<td>4. High content of protein, cells, and solid materials derived from cells</td>
<td>4. Low content of protein (&lt;2.5–3.0 g/dL or &lt;25–30 g/L), cells, or solid materials derived from cells</td>
</tr>
<tr>
<td>5. May have high WBC content</td>
<td>5. Has low WBC content</td>
</tr>
<tr>
<td>6. Clots spontaneously (contains high concentration of fibrinogen)</td>
<td>6. Will not clot</td>
</tr>
<tr>
<td>7. Malignant cells as well as bacteria may be detected</td>
<td>7. Malignant cells may be present</td>
</tr>
<tr>
<td>8. Specific gravity &gt;1.016</td>
<td>8. Specific gravity &lt;1.016</td>
</tr>
</tbody>
</table>

Fluid contained in the pleural, pericardial, peritoneal, or abdominal cavity is a serous fluid. Accumulation of fluid in the peritoneal cavity is called ascites. Cytologic studies of effusions (exudate or transudates) are helpful in determining the cause of these abnormal collections of fluids. The effusions are found in the pericardial sac, the pleural cavities, and the abdominal cavities. The chief problem in diagnosis is in differentiating malignant cells from reactive mesothelial cells.

Reference Values

Normal Negative for abnormal cells

Procedure

1. General procedure
   a. Obtain material for cytologic examination of effusions by either thoracentesis or paracentesis.
   b. Remember that both of these procedures involve surgical puncture or a cavity aspiration of a fluid.
   c. Fluid may be obtained in syringes, vacuum bottles, or other containers, depending on the volume of accumulated fluid. Heparin may be added to prevent clotting. Check with your laboratory for specific instructions.
2. Thoracentesis procedure
   a. Ensure that chest x-rays are available at the patient's bedside so that the location of fluid may be determined.
   b. Give the patient a sedative if necessary.
   c. Expose the chest. The physician inserts a long thoracentesis needle with a syringe attached.
   d. Withdraw at least 40 mL of fluid. It is preferable to withdraw 300 to 1000 mL of fluid.
   e. Collect the specimen in a clean container and add heparin if necessary, particularly if the specimen is very bloody (5 to 10 U of heparin per milliliter of fluid). Do not add alcohol.
   f. Label the specimen with the patient's name, the date, the source of the fluid, and the diagnosis.
   g. Send the covered specimen immediately to the laboratory. (If the specimen cannot be sent at once, it may be
3. Paracentesis (abdominal) procedure
   a. Ask the patient to void.
   b. Place the patient in Fowler's position.
   c. Give a local anesthetic.
   d. Introduce a No. 20 needle into the patient's abdomen and withdraw fluid, 50 mL at a time, until 300 to 1000 mL has been withdrawn.
   e. Follow the same procedure for collection and transport of the specimen as for thoracentesis.
   f. For all procedures, see Chapter 1 guidelines for intratest care.

**Clinical Alert**

Paracentesis can precipitate hepatic coma in a patient with chronic liver disease. The patient must be watched constantly for indications of shock: pallor, cyanosis, or dizziness. Emergency stimulants should be ready.

**Clinical Implications**

1. All effusions contain some mesothelial cells. (Mesothelial cells make up the epithelial layer covering the surface of all serous membranes.) The more chronic and irritating the condition, the more numerous and atypical are the mesothelial cells. Histiocytes and lymphocytes are common.

2. Evidence of abnormalities in serous fluids is characterized by:
   a. Degenerating red blood cells, granular red cell fragments, and histiocytes containing blood. Presence of these structures means that injury to a vessel or vessels is part of the condition causing fluid to accumulate.
   b. Mucin, which is suggestive of adenocarcinoma
   c. Large numbers of polymorphonuclear leukocytes, which is indicative of an acute inflammatory process such as peritonitis
   d. Prevalence of plasma cells, which suggest parasitic infestation, Hodgkin's disease, or hypersensitive state
   e. Presence of many reactive mesothelial cells together with hemosiderin histiocytes, which may indicate:
      1. Leaking aneurysm
      2. Rheumatoid arthritis
      3. Lupus erythematosus
   f. Malignant cells

3. Abnormal cells may be indicative of:
   a. Malignancy (The most important criterion of cancer is the arrangement of chromatin within the nuclei.)
   b. Inflammatory conditions

**Interfering Factors** Vigorous shaking and stirring of specimens causes altered results.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose of the test and the procedure. The procedure varies depending on the site of fluid accumulation. General patient preparation includes measuring blood pressure, temperature, pulse, and respiration; administering sedation as ordered; preparing local anesthetic as ordered; providing emotional support; and obtaining a signed consent form.

2. Be aware that local anesthetic and sedative may be ordered to achieve a state of conscious sedation (see Appendix C).

3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Clinical Alert**

The only contraindication is an uncooperative patient.

**Posttest Patient Aftercare**

1. Monitor according to agency protocols.
2. Check blood pressure, pulse, and respirations every 15 minutes for 1 hour, then every 2 hours for 4 hours, and as ordered. Check temperature every 4 hours for 24 hours. Apply adhesive bandage or dressing to site of puncture. Check dressing every 15 to 30 minutes. Turn patient onto the unaffected side for 1 hour, then to a position of comfort. Manage pain as indicated. Measure and record the total amount of fluid removed; note its color and character.

3. See guidelines in Chapter 1 for safe, effective, informed posttest care.

**Skin/Cutaneous Immunofluorescence Biopsy: Cell (Cytologic) Study and Tissue (Histologic) Study**

Biopsy of the skin for direct epidermal fluorescent studies is indicated in the investigation of certain disorders such as lupus erythematosus, blistering disease, and vasculitis. Skin biopsies are also used to confirm the histopathology of skin lesions, to rule out other diagnoses (ie, herpes simplex and psoriasis), and to monitor the results of treatment.

**Reference Values**

**Normal** A descriptive interpretative report of the skin biopsy is made.

**Procedure**

1. Obtain a 3- to 6-mm punch biopsy or shave biopsy, excisional biopsy, or incisional biopsy specimen of involved or uninvolved skin. Scraping smears and/or aspirates also may be obtained. Take care not to crush the specimen.
2. Check with your laboratory for specific guidelines for specimen handling.
3. See Chapter 1 guidelines for intratest care.
Clinical Implications

1. Biopsy of skin shows the lesions of discoid lupus erythematosus as a bandlike immunofluorescence of immunoglobulins and complement components. Similar findings in a biopsy of normal skin are consistent with SLE and may be used to monitor the results of treatment.

2. In blistering diseases such as pemphigus and pemphigoid, in which circulating antibodies may not be present, a lesion may show intercellular epidermal antibody or pemphigus or basement membrane antibody of pemphigoid.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the skin biopsy. Local anesthesia will be used.

2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Clinical Alert

Contraindications include:

1. An uncooperative patient
2. Bleeding diathesis tendency—anticoagulant therapy

Posttest Patient Aftercare

1. Monitor biopsy site for infection or bleeding. Counsel and interpret outcomes and need for possible treatment.

2. See Chapter 1 for safe, effective, informed posttest care.

Estrogen/Estradiol Receptor (ER), Progesterone Receptor (PR) Tissue (Histologic) Study and DNA Ploidy (Tumor Aneuploidy)

Cancers have abnormal amounts of nuclear DNA. The higher the grade of tumor cells, the more likely the DNA content will be abnormal. The determination of tumor ploidy (the number of chromosome sets in a cell; ie, diploid, two sets, triploid, three sets) by various methods: flow cytometry (Fc), histograms, and image analysis divide cells into triploid/diploid (slowly replicating cells) or aneuploid (rapidly replicating cells). ER and PR in the cells of breast and endometrial cancer tissues are measured to determine whether the cancer is likely to respond to endocrine therapy or to removal of the ovaries. DNA ploidy measures cell turnover (replication) in a specimen identified as cancer and predicts progress, shorter survival, and relapse in some patients with cancer: bladder, breast, colon, endometrial, prostate, kidney, and thyroid. The predictive value is greater for breast, prostate, and colon.

Reference Values

Normal

ER: negative; <3 femtomoles (fmol)/mg (<3.0 nmol/kg) of protein

PR: negative; <5 fmol/mg (<5.0 nmol/kg) of protein

DNA index (DI): 0.9–1.0 is normal DNA ploidy (content) or the diploid state. An interpretive histogram by flow cytometry classifies the stained nucleic as DNA diploid, DNA aneuploid, DNA tetraploid, or DNA uninterpretable.

Procedure

1. Obtain a fresh specimen by biopsy, keep on ice, and deliver immediately to the histology laboratory.

2. Examine a 1-g specimen of quickly frozen tumor for saturation and express in a Scatchard's plot. Do not place the specimen in formalin. Some laboratories can perform ERA/PRA studies on paraffin-embedded tissue. Check with your laboratory for specific instructions.

3. Classify specimens for DNA ploidy on the basis of the percentage of epithelial cells that contain diploid (2n) DNA content and nondiploid DNA (aneuploid). DNA content is calculated as the DNA index.

4. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. A positive test for ER occurs at levels 10 fmol (10 nmol/kg) and for PR binding at levels of 10 fmol (10 nmol/kg). The frequency of positive ER and PR occurs more often in postmenopausal women.

2. Approximately 50% of ER-positive tumors respond to antiestrogen therapy, and 60%–70% respond in patients with both ER- and PR-positive tumors.

3. ER-negative tumors rarely respond to antiestrogen therapy.

4. The finding of positive progesterone increases the predictive value of selecting patients for hormonal therapy. There is some evidence to suggest that progesterone receptor synthesis is estrogen dependent.

5. The presence of aneuploid peaks in the replicative activity of neoplastic cells may be prognostically significant, independent of tumor grade and stage.

6. The greater the amount of cells in S phase (DNA synthesis) of the cell cycle, the more aggressive the tumor. Positive aneuploidy points to a favorable prognosis in some conditions, such as acute lymphoblastic lymphoma and neuroblastoma and perhaps transitional cell bladder cancer.

Interventions

Pretest Patient Care

1. Explain purpose and procedure of testing. See Tissue (Histologic) Biopsy Studies: Overview; Prognostic and Predictive Markers; and Breast Biopsy: Cell (Cytologic) and Tissue (Histologic) Study and Prognostic Markers on pages 766 and 770. Obtain appropriate clinical history so that this information can be provided with the specimen.

2. Be aware that positive ER and PR means that antiestrogen drug therapy may be beneficial.

3. See Chapter 1 for safe, effective, informed pretest care.
Clinical Alert

Contraindications include:
1. An uncooperative patient
2. Bleeding diathesis (tendency to spontaneous bleeding due to coagulation defect)—anticoagulant therapy

Posttest Patient Aftercare

1. Interpret test results and counsel appropriately about possible treatment.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

GENETIC STUDIES

Overview of Genetic Studies

Genetics is concerned with the components and function of biologic inheritance. Genetic testing investigates the presence, absence, or activity of genes through direct and indirect means and by chemical analysis, microscopic methods, submicroscopic techniques, and molecular biology studies.

Insight into causes of developmental problems, birth defects, and heritable disorders often involves genetic studies. Basic technology counts the chromosomes in a person's cells or measures the amount of specific proteins and enzymes. At the other end of the spectrum, cellular DNA can be assayed with molecular probes designed to identify a unique genetic sequence. Genetic testing is a rapidly evolving field that has shown expanding possibilities, an ever-increasing number of tests, and a unique set of limitations and dilemmas.

Many disease states reflect hereditary components even though general clinical studies usually focus on the disorder itself rather than on its genetic components. This section addresses circumstances that may require biochemical analysis (enzymes, organic acids, amino acids) and DNA tests of cytogenetic (chromosome) studies for proper diagnosis and/or management.

Testing may be done before birth, neonatally, during childhood or adult life, or postmortem.

Clinical Alert

1. Cytogenetics is the part of genetics concerned with the structure and functions of cells, especially chromosomes. Genetic tests are done to identify inborn errors of metabolism, to determine sex when ambiguous genitalia are present as well as to detect chromosome aberrations such as Down syndrome.
2. Biochemical analysis and tests to detect carrier status of inborn errors of metabolism are being done more and more often, primarily through detection of abnormal accumulation in body fluids and tissue. Advances in the study of molecular genetics has been affected by the completion of the Human Genome Project. Tests in this category include diagnosis of neoplastic disease (eg, Philadelphia chromosome in chronic myelocytic leukemia and N-MYC gene in neuroblastoma) and inherited disorders (eg, cystic fibrosis, spinal cerebellar ataxia).

Indications for Testing

1. Prenatal care: medical management of potentially problematic pregnancies identified through abnormal maternal screening, ultrasound, or family history may include specific genetic testing and realistic information about fetal abnormalities to allow parents to make informed decisions about pregnancy continuation.
2. Newborn screening, to detect preventable, common, or treatable disease
3. Investigation of fetal death, stillbirth, or miscarriage
4. Decisive diagnosis: diagnostic and/or presymptomatic studies may be done to investigate certain syndromes or diseases related to chromosomal or single gene disorders; for gene status in the diagnosis of cancer; for inherited diseases in carrier status; and to test asymptomatic relatives at risk for developing significant medical or reproductive problems. As a general rule, presymptomatic testing is not offered to minors.
5. Cytogenetic analysis or DNA probes are used to study bone marrow and to check for diagnostic translocations, particularly in leukemias (cells with genetic changes).
6. Assignment of gender in the presence of ambiguous genitalia
7. Genetic counseling: to address prognosis and diagnosis as well as causes and recurrence risks in the context of the family as well as individuals. Genetic counseling often depends on precise testing or chromosomal analysis.
Inheritance in Human Disorders

Genetic information is coded within DNA. This information is packaged into chromosomes that are present in cell nuclei. In humans, 46 chromosomes contain an estimated 30,000 to 50,000 gene pairs. DNA contains four distinct molecules (base pairs). These four base pairs code the information that controls growth, development, and function by providing a template for message molecules called RNA. RNA molecules are involved in the process of transcription (changing a DNA message into a protein) as well as in providing molecules that regulate expression, make the hardware within the cell for building proteins (ribosomes), and also perform many housekeeping functions.

With the exception of red blood cells and egg/sperm cells, there are 23 pairs of chromosomes in human cell nuclei. One of each pair comes from each parent. Twenty-two of these pairs match up and contain copies of the same genes (although the copies may not be identical). These chromosomes are assigned numbers, and they are called autosomes. Then, there are two sex chromosomes, X and Y. Females have two X-chromosomes; males have one X-and one Y-chromosome. The Y chromosome is unique to males and contains genes that determine male structure and function and also affect fertility. The Y chromosome contains very few genes, but its presence or absence determines male or female development.

Genes, like chromosomes, come in pairs, except for genes on the sex chromosomes.

1. **Autosomal dominant inheritance.** Within one gene pair, an abnormality in a single copy of the gene may produce a disorder. A person with such a gene combination would have a theoretical 50-50 chance of passing this gene on to any offspring. Dominant disorders may therefore be inherited from a parent, or they may arise as a new mutation in an egg or sperm cell that participates in fertilization. For many dominantly inherited conditions, manifestations of the disorder are not consistent. This observation is known as variable expression. Examples of dominantly inherited disorders include Huntington's disease, neurofibromatosis, familial hypercholesterolemia, and hereditary colon cancer (Chart 11.2).

2. **Autosomal recessive inheritance.** Both copies of the gene pair must not function correctly for a problem to be apparent. If both parents carry the same nonfunctional gene, there is a 1-in-4 chance that any child could inherit two nonfunctional copies, leading to possible disease. Examples of autosomal recessively inherited diseases include cystic fibrosis, sickle cell disease, Tay-Sachs disease, some nonsyndromic early-onset hearing loss, and recurrent pyogenic infections.

3. **X-linked recessive inheritance.** Males have only one X chromosome, so that abnormal genes on the X chromosome can cause problems. Females have a second X chromosome, which usually masks the affects of an abnormal gene, although not always completely. A woman with a disease-causing gene on one X chromosome would have a 50-50 chance of passing this gene to any child, and this is independent of her 50-50 chance of having a son. Examples of X-linked disorders include hemophilia and Duchenne's muscular dystrophy.

4. **Multifactorial inheritance.** Some developmental processes, as well as some adult disease states, are influenced by the interactions of many genes associated with environmental factors. Examples of multifactorial disorders include pyloric stenosis, cleft lip and palate, spinal bifida, and schizophrenia.

5. **Cytogenic inheritance.** Chromosomal abnormalities may include abnormal numbers of chromosomes (eg, Down syndrome is caused by three copies of chromosome 21). Chromosomal rearrangements, called translocations, can be unbalanced, causing multiple congenital abnormalities. A molecular abnormality within a single gene can cause structural differences like fragile-X mental retardation syndrome. Submicroscopic deletions of chromosomes can be studied using fluorescent in situ hybridization. Examples of human syndromes caused by microdeletions include Williams' syndrome and DiGeorge's syndrome.

6. **Mitochondrial inheritance.** Separate from the nucleus of the cell are the energy-processing organelles called mitochondria. These organelles possess a unique set of genes on a single chromosome. Mutations in these genes can cause a wide variety of disorders, including neuromuscular disorders. Examples include Kearns-Sayre syndrome and Leber's hereditary optic neuropathy. Mitochondria are inherited exclusively from the mother.

7. **Nontraditional inheritance.** Some human genes are sensitive to modification (known as imprinting or methylation).
that alters gene expression, depending on the sex of the parent in which the gene originates. Some human syndromes are caused by the presence of two copies of a gene or chromosome originating from one parent, and none from the other (called uniparental disomy, or UPD). Examples include Beckwith-Wiedemann syndrome and Prader-Willi syndrome.

### Chart 11.2 Types of Genetic Disorders

<table>
<thead>
<tr>
<th>Autosomal Dominant</th>
<th>Autosomal Recessive</th>
<th>X-Linked Recessive</th>
</tr>
</thead>
<tbody>
<tr>
<td>Familial breast cancer</td>
<td>Sickle cell anemia</td>
<td>Hemophilia A and B</td>
</tr>
<tr>
<td>Huntington's disease</td>
<td>Thalassemia β and α</td>
<td>Duchenne's and Becker's muscular dystrophy</td>
</tr>
<tr>
<td>Adult polycystic disease (some types)</td>
<td>Cystic fibrosis</td>
<td>Fragile X syndrome</td>
</tr>
<tr>
<td>Myotonic dystrophy</td>
<td>Alpha 1-antitrypsin deficiency</td>
<td>Omithene transcarbamylase deficiency (OTC)</td>
</tr>
<tr>
<td></td>
<td>Tay-Sachs disease</td>
<td></td>
</tr>
</tbody>
</table>

### Direct Detection of Abnormal Genes by DNA Testing

Many genetic diseases continue to be detected by the effects they produce in abnormal body structure, function, or chemistry. With the elucidation of gene structure and the cataloging of human gene mutations, direct detection of hundreds of mutations is possible. For known genetic disorders for which specific mutational analysis is not available, indirect analysis using varied techniques, including protein expression, may be applicable. A database of medical genetics information resources for physicians and other health care providers and researchers maintains daily updates of available testing worldwide at www.genetests.org. Registration is required. Genetic testing for diagnostic purposes requires patient education and consent, physician request, and coordination of sample collections. Diagnostic tests must be done in a Clinical Laboratory Improvement Act (CLIA)-approved laboratory. Research labs cannot provide this service and should not be contacted for clinical testing. Sensitivity of testing in genetic disease needs to be addressed because many diseases may have different causes, and many genetic tests are not capable of finding all mutations in complex genes. For example, nearly 1000 gene mutations have been linked to cystic fibrosis, but still an estimated 3% to 10% of mutations cannot be found. Interpretation of results can be a challenge also, especially in situations in which a gene change can be demonstrated but it is not known whether it is a harmless change or not. An example of this is polymorphism in the BRCA1 gene. In such situations, interpretation may rely on comparison to gene changes found in known affected relatives. A variety of morbid genetic changes have been discovered, including gain or loss of a single base pair or larger group of base pairs as well as repetitive sequences that get copied over and over so many times that they disable the function of the gene. Detection strategies are tailored to the type of mutation present or suspected.

### Procedure

1. Establish availability and sensitivity of clinical testing and inform the patient of the benefits, limitations and consequences of testing (see Genetic Counseling, next). Informed consent may be required. Prepayment may be required. Test results may take weeks or months.
2. Obtain samples or specimens of body fluids or tissues as specified by the receiving laboratory. Overnight shipment must usually be arranged.

### Clinical Implications

1. Improved diagnosis of types of cancer may have therapeutic implications.
2. Discovery of hereditary disease or cancer may have implications for other family members.
3. Precise DNA tests can be done for some inherited diseases (eg, cystic fibrosis, Duchenne's and Becker's muscular dystrophy, some polycystic kidney diseases).
4. Paternity identity testing and forensic testing
5. Identification of microbes in infectious diseases (eg, chlamydia, cytomegalovirus)
6. Prediction of progression in neuromuscular disorders (eg, Huntington's disease, myotonic dystrophy, cerebellar ataxia)
7. Identification of comorbid disease risks (eg, progressive kidney failure in some hearing-loss syndromes)
8. Identification of reproductive risks
9. Explanation of miscarriage and stillbirth
10. Potential associations with common diseases of aging (eg, cardiovascular disease, Alzheimer's disease)

### Genetic Counseling

Genetic counseling is the process of providing individuals and families with information on the nature, inheritance, and implications of genetic disorders in order to help them make informed medical and personal decisions. Risk assessment, family history, and genetic testing to clarify genetic status of family members may be part of the genetic counseling process.

Genetic counselors are health care professionals with specialized education, training, and experience in medical genetics. They frequently work as part of a team that includes physicians and other specialists in biochemistry and genetics, and they coordinate activities with many medical specialties, including prenatal care, pediatrics specialties, neurology, hematology, and laboratory testing.

Genetics services are available at or through most major medical centers in the United States and serve the medical and lay communities as sources of information, clinical evaluation, management of genetic conditions and birth defects, and coordination with appropriate testing services. Geographic listings of genetic clinics and genetic counselors can be found...
When testing for genetic disease is being considered, pretest counseling may include additional attention to issues of realistic usefulness of currently available tests and consideration of personal, family, privacy, and insurance implications of testing. Just because a test is available does not mean it is appropriate—unwanted information can be generated by genetic testing, tests may cost thousands of dollars, and ambiguous results are possible.

Posttest counseling not only presents test results but reviews medical and psychological implications for the family and potentially may be expanded to include other family members for counseling and testing. Experience with rare genetic diseases once identified will add to the direction of specific medical care and therapy, and patient education and counseling can assist the process of identifying options and resources.

The number of specific genetic tests is increasing rapidly, although availability may be limited and the cost may be well over $500 and not covered by health insurance. An additional dilemma is the lack of usefulness of testing in many disorders to rule out a specific diagnosis. (For example, a tall thin individual with some heart findings like mitral valve prolapse may be thought to have Marfan's syndrome. Currently, testing for the gene that causes Marfan's syndrome can be done, but it does not find many mutations, even in individuals who are known to have the syndrome.)

Often, it is necessary to study an affected family member to determine what gene mutation is present in a family. This can be problematic in diseases like breast cancer because the affected persons may be deceased, unavailable, or uncooperative because of family dynamics. If a family wishes to be studied but no gene mutation is identified, linkage studies might be considered to estimate risks within a family. Because the possibility of identifying gene changes associated with human disease now exists, so does the expectation and challenge of improving treatment and understanding of both rare and common diseases.

**CYTOGENETICS**

Chromosomal Analysis

The karyotype, a study of chromosome distribution for an individual, determines chromosome numbers and chromosome structure (Chart 11.3); alterations in either of these can produce problems. The standard karyotype can be a diagnostic precursor to genetic counseling. Additional or missing pieces of most chromosomal material cause developmental problems. Despite much speculation, it is not known exactly how these abnormalities translate into structural or functional anomalies. Predictions almost always depend on comparisons with clinical findings from other similar cases that present the same evidence.

**Chart 11.3 Definition and Nomenclature of Karyotype**

**Background**

The karyotype is an arrangement of the chromosomes on a cell into a specific order, from the largest size to the smallest, so that their number and structure can be analyzed. This is routinely done through banding, a technique that permits detection of the differences in structure between the different pairs. Before banding, it was often impossible to pair chromosomes correctly; instead, they were arranged in groups according to size and structure and labeled A through G. The X chromosomes were part of group C, and the Y chromosomes belonged to group E. Now, they are usually placed with each other, apart from the other pairs.

The pairs of chromosomes are differentiated according to the following characteristics:

1. Their length
2. The location of the centromere, the constriction that divides chromosomes into long (q) and short (p) arms
3. Ratio of the long and short arms to each other
4. Secondary constrictions
5. Satellites, which are small, variable pieces of DNA seen at the ends of the arms of some chromosomes
6. Staining or banding patterns. A variety of different stains and techniques can be used. The most common is Giemsa banding. Most of the other methods, such as centromeric or fluorescent staining, are restricted to specific situations.

**Nomenclature of the Karyotype**

The standard conventions for listing karyotypes is as follows:

1. The first number denotes the total number of chromosomes.
2. Second, the sex chromosome complement follows (usually XX for normal females and XY for normal males).
3. Third, the missing, extra, or abnormal chromosomes are identified.
4. The letter "p" refers to the short arm, "q" to the long arm.
5. Bands are numbered from the centromere out. As techniques evolve, these are further subdivided. For example, in the two-digit number 32, the first number (3) is the band and the second number (2) is the subdivision of that band (band 32). Decimal points indicate further division under the same system; for example (working backward), 32.41 is the first subdivision (1) of the fourth subdivision (4) of the second subdivision of the third band.
6. A three-letter code at the end designates the banding technique. The first letter is the type of banding; the second letter denotes the general technique; the third letter indicates the stain. Probably the most common code is GTG: band type G, banding by trypsin, using Giemsa stain. Special or unusual techniques are used only in selected circumstances.
Standard chromosome studies can be helpful in evaluation of the following clinical situations:

1. Multiple malformations of structure and function
2. Failure to thrive
3. Mental retardation
4. Ambiguous genitalia or hypogonadism
5. Recurrent miscarriages
6. Infertility
7. Primary amenorrhea or oligomenorrhea
8. Delayed onset of puberty
9. Stillbirths or miscarriages (particularly with associated malformations)
10. Prenatal diagnosis of potential or actual abnormalities related to chromosome disorders (eg, Down syndrome, especially in offspring of mothers > 35 years of age)
11. Detection of parents with chromosomal mosaicism or translocations, who may be at high risk for transmitting genetic abnormalities to their children
12. Sex determination
13. Selected cancers and leukemias in which abnormalities of the chromosomes may reveal prognosis or disease stage

Reference Values

Normal 46 chromosomes Women: 44 autosomes + 2 X chromosomes (karyotype 46, XX) Men: 44 autosomes + 1 X and 1 Y chromosome (karyotype 46, XY) A photograph of representative karyotype is included with report.

Procedure Specimens for chromosome analyses are generally obtained as follows, using aseptic procedures and special kits and containers:

1. Be aware that heparinized venous blood leukocytes from peripheral vascular blood samples are used most frequently because they are the most easily obtained. Preparation of the cells takes at least 3 days. The time required is directly proportional to the complexity of the analytic process.
2. Collect bone marrow in a green-topped tube, at least 5 mL in a heparinized syringe (20–25 units of heparin). Biopsies can sometimes be completed within 24 hours. Bone marrow analysis is often done to diagnosis certain categories of leukemias.
3. Remember that fibroblasts from skin or other surgical specimens can be grown and preserved in long-term culture mediums for future studies. Growth of a sufficient amount of the specimen for studies usually requires at least 1 week. These specimens are especially helpful in detecting mosaicism (different chromosome constitutions in different tissues) and in the study of stillbirths, neonatal death, and spontaneous abortion.
4. Be aware that amniotic fluid in the prenatal period obtained through amniocentesis stored in a sterile container requires at least 1 week to produce a sufficient amount of cell growth for analysis. These studies are often done for prenatal detection of chromosomal abnormalities (see Chapter 15).
5. Remember that chorionic villus sampling (CVS) can be done at earlier stages of pregnancy (about 9 weeks) than can amniocentesis. Some initial CVS studies can be done almost immediately after conception. Occasional false-positive results represent mosaicism of the placenta (the presence of several cell lines, some of which may not be found in the fetus). These studies need confirmation of findings through long-term culture (see Chapter 15).
6. Grow cells from fetal tissue or from early-trimester products of conception to determine causes of spontaneous abortion. Cells from the fetal surface of the placenta may be easiest to grow and are the most likely to be successful.
7. Take the buccal smear, for detecting sex chromosomes, from the inner cheek and use fluorescent in situ hybridization with probes specific for the X or Y chromosome.
8. Take dried blood spot from heel of newborn.
9. Place specimens of lymph nodes or solid tumors in sterile containers.
10. Remember that chromosome analysis is often performed using other specimens, such as skin, fascia, lung tissue, kidney, or the placenta. At least 2 mm of volume is needed for an adequate specimen.
11. See Chapter 1 guidelines for intratest care.

Clinical Implications Many chromosomal abnormalities can be placed into one of two classes; some examples follow:

1. Abnormalities of number
   a. Autosomal:
      1. Trisomy 21 (Down syndrome)
      2. Trisomy 18 (Edward's syndrome)
      3. Trisomy 13 (Patau's syndrome)
   b. Sex chromosome syndrome:
      1. Turner-Ulrect syndrome (45 single X)—short stature, webbed neck, and renal and C anomalies
      2. Klionsel's syndrome (47 XXXY)—hypogonadism, infertility, learning disabilities, undeveloped secondary characteristics
      3. XXX, 47 XXY—tall, increased risk for behavior problems
      4. Triple XXX—increased risk for infertility and behavior problems
2. Abnormalities of structure
   a. Deletions:
      1. Cri du chat/cat's cry syndrome: the distal part of the chromosome 5 short arm is deleted
      2. Missing short arm of chromosome 18: 18p- is deleted
      3. Prader-Willi syndrome: 15 Q is deleted in some cases

More than 80 other abbreviations can be used to label other structural findings. Some of the more common ones are mentioned in clinical implications of chromosome analyses.
b. Duplications: extra material from the second band in the long arm of the third chromosome: 3q2 trisomy (Cornelia de Lange's syndrome resemblance)
c. Translocations: translocation of chromosomes 11 and 22: t(11;22) or 14 and 21
d. Isochromosomes: a single chromosome with duplication of the long arm of the X chromosome: i(Xq) (a variant of Turner's syndrome)
e. Ring chromosomes: a chromosome 13 with the ends of the long and short arms joined together, as in a ring: r(13)
f. Mosaicism: two cell lines, 1 normal female and the other for Turner's syndrome: 46, X, 45, X

Interventions

Pretest Patient Care

1. Provide information and referrals for appropriate genetic counseling and treatment if necessary.
2. Explain the purpose, procedure, and limitations of the genetic test together with the known risks and benefits. This education process should be done by a genetic counselor.
3. Obtain informed, signed, and witnessed consent. This is required for most genetic tests.
4. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. If an amniotic fluid specimen or CVS is obtained for analysis, follow the same precautions as listed in Chapter 15.
2. Provide timely information and compassionate support and guidance for parents, children, and significant others.
3. See Chapter 1 for guidelines for safe, effective, informed posttest care.

Clinical Alert

1. Occasionally, it is possible to line a certain chromosomal pattern with specific genes and to then understand the clinical picture from analyzing these results. However, for the most part, the association between specific chromosomal abnormalities and specific sets of findings is not yet well understood. Interpretations from karyotype studies usually come from correlations with similar cases rather than from any theoretical considerations. Therefore, because many variables exist, predictions must be made cautiously and judiciously.
2. Most laboratories provide interpretations of results. However, it may be necessary to talk directly with laboratory personnel to fully understand the meaning of an unusual karyotype.

Special Chromosomal Studies

The fragile-X syndrome is one of the most common genetic causes of mental retardation. An X-linked trait, it is more commonly seen in males. Females may carry this gene without exhibiting any of its characteristics; however, they can also be as severely affected as males. This syndrome takes its name from the small area on the long arm of the X chromosome that looks like a break in the arm (although it actually is not). The cells need to be grown in a special medium to reveal this pattern; a regular karyotype will miss it. Even with the special medium, not all cells show the characteristic. In female carriers of this trait, the syndrome becomes harder to detect as the woman ages. Accurate detection of fragile-X syndrome at a molecular level is now available.

Rare conditions such as excess chromosome breakage (Fanconi's anemia) or abnormal centromeres (Roberts' syndrome) merit special analytic processes and procedures. Chromosome and molecular analyses are done using a venous blood sample (5 mL with EDTA tube) to identify the fragile-X mental retardation syndrome and possible carrier status.

Newborn Screening for Congenital Disorders

All North American states and provinces and most industrialized countries of the world require screening of newborns to detect congenital and metabolic disorders within the first week of life (eg, before the baby is discharged from the hospital or maternity home, or after home birth, specimens are collected by physicians, nurses, or birth attendant [midwife]). Providing and performing the testing and following up on all abnormal results is the responsibility of state health departments. Some state health departments also supply required types of formulas. For those babies with a confirmed congenital disorder, genetic counseling, treatment, and long-term care are also provided in most states (eg, special dietary formula for parents with PKU children is free of charge).

Procedure

1. Newborn blood is sampled using a heel-stick procedure in the first week of life (see Chapter 2). A special kit is used.
2. Apply the blood drops (less than 0.5 mL) to a piece of special filter paper, which usually contains three to five printed circles.
3. Fill all the printed circles with blood, which in most cases can be done with one drop of blood per circle.
4. Be sure the attached coverslip does not come into contact with the blood until completely dry. Do not permit the blood-soaked portion of the collection kit to come in contact with another surface (eg, desktop, absorbent paper). Proper collection procedures are based on the National Committee on Clinical Laboratory Standard (NCCLS) document LA4-A4: Blood Collection on Filter Paper for Newborn Screening Programs; Approved Standard.
5. In addition to the filter paper, the collection kit also contains a multipart form requesting information regarding the baby’s name, mother’s name, birth date and time, specimen collection data and time, birth weight, and so forth. It is important that this form be filled out completely because the information is critical for the laboratory staff in the interpretation of test results (see example of form used in Appendix H). For example, differentiating normal from
abnormal thyroid-stimulating hormone (TSH) results for potential hypothyroidism may be dependent on the age (in hours) of the infant at the time of special collection.

6. After the specimen is collected, allow the blood card to air-dry in a horizontal position for a minimum of 3 hours at room temperature. After the blood is dried, the specimen should be forwarded (mail or courier) to the screening laboratory within 24 hours.

Processes vary among newborn screening laboratories. Most testing begins by punching a 1/8-inch blood spot into a 96-well microplate (or some other vessel such as a test tube or dimple tray). The technology used to test the blood can be described under general headings.

1. Basic chemistry procedures, such as measurement of phenylalanine or galactose metabolites
2. Electrophoresis technology, or liquid chemotherapy, is used to separate hemoglobin fractions and to quantify the amount of each hemoglobin based on difference in electrical charge and mobility in an electric field.
3. Immunochemistry (eg, antigen-antibody reactions) is used to measure thyroid hormones (hypothyroidism), 17-hydroxyprogesterone (congenital adrenal hyperplasia), and immunoreactive trypsinogen enzyme (cystic fibrosis).
4. Gene mutation analysis to detect the genetic mutation causing the disorder's (cystic fibrosis, hemoglobin) alternate assay, are tested using mutation assays.
5. Tandem mass spectrometry (MS/MS) has allowed newborn screening programs to expand significantly the number of disorders screened using a single 1/8-inch spot (eg, fatty acid oxidation [FAO], organic acidemia [OA], urea cycle [UC], and aminoacidopathies [AA]). The FAO and OA disorders are detected by measuring acylcarnitines (an intermediate compound containing fatty acids or organic acids combined with carnitine that occurs from blocked metabolic pathways). At least 20 disorders can be detected by this technology.

Chart 11.4 is a list of disorders that currently can be detected through newborn screening. It should be noted that the information provided is generally not considered all-inclusive, and results are considered presumptive (requiring confirmation before a formal diagnosis is made and treatment implemented). The State Public Health Laboratory or Health Department in specific states or regions will provide information about the status of newborn screening in a state or region. Abnormal results are indicated by change in paper color, as follows: blue—possibly abnormal, gold—definitely abnormal. Abnormal values represent the absence of expected enzyme activity, elevated or decreased hormone values, presence of abnormal or variant hemoglobins, and abnormal levels of amino acids and evidence of fatty acid disorders or genetic mutation.

<table>
<thead>
<tr>
<th>Chart 11.4 Disorders Currently Detectable Through Newborn Screening</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Biotinidase</strong></td>
<td></td>
</tr>
<tr>
<td>Prevalence:</td>
<td>1:100,000</td>
</tr>
<tr>
<td>Substance measured:</td>
<td>Biotinidase enzyme activity</td>
</tr>
<tr>
<td>Determined by:</td>
<td>Basic chemistry</td>
</tr>
<tr>
<td>Abnormal:</td>
<td>No enzyme activity. Birth weight and age in hours dependent</td>
</tr>
<tr>
<td>Interfering factors:</td>
<td>Transfusions may cause a false-negative result</td>
</tr>
<tr>
<td>Additional diagnostic testing:</td>
<td>Serum/plasma enzyme quantification</td>
</tr>
<tr>
<td>Treatment:</td>
<td>Biotin—daily supplements</td>
</tr>
<tr>
<td>Clinical symptoms (not treated):</td>
<td>Seizures, dermatitis, hair loss</td>
</tr>
</tbody>
</table>

**Congenital Adrenal Hyperplasia (CAH)**

| Prevalence | 1:10,000 |
| Substance measured: | 17-Hydroxyprogesterone (17-OHP) |
| Determined by: | Immunochemistry |
| Abnormal (cutoff): | Elevated 17-OHP: typical cutoff range: >90 ng/dL (>2.7 nmol/L) are considered critical in full term babies. Cutoffs will vary by program in low-birth-weight babies. |
| Interfering factors: | False positives can be expected in low-birth-weight babies and early sample collections (<24 hours). |
| Diagnostic testing: | Refer to a pediatric endocrinologist for CAH workup |
| Treatment: | Glucocorticoid replacement, 9-a-fluorohydrocortisone |
| Clinical symptoms (not treated): | Salt-losing crisis in males that often results in death. Virilization (ambiguous genitalia) in females |

**Congenital Hypothyroidism**

| Prevalence | 1:3,000 |
| Substance measured: | Thyroid-stimulating hormone (TSH) and/or thyroxine (T₄) |
Substance measured: Thyroid-stimulating hormone (TSH) and/or thyroxine (T4)
Determined by: Immunochemistry
Abnormal (cutoff): Elevated TSH: typical cutoff range: 20–50 µg/dL (258–645 nmol/L) (program dependent)
Decreased T4: typical cutoff range: 6–8 µg/dL (77–103 nmol/L) (program dependent)
Interfering factors: False positives can be expected in early (<24 hour) discharges. False negatives can occur in very-low-birth-weight babies.
Additional diagnostic testing: Serum T4 and TSH measurements
Treatment: Synthroid—daily supplements
Clinical symptoms (not treated): Mental retardation, cretinism, liver failure

Cystic Fibrosis (CF)

Prevalence: 1:4000 (Caucasians)
Substance measured: Immunoreactive trypsinogen (IRT)
Mutant alleles
Determined by: Immunochemistry
Mutation analysis
Interfering factors: False negatives can occur because some CF mutations may not cause an IRT elevation.
Abnormal (cutoff): Elevated IRT: typical cutoff range: 140–180 ng/mL (program dependent)
Mutation analysis: detection of mutant allele(s)
Additional diagnostic testing: Pilocarpine iontophoresis sweat chloride test
Treatment: Care at a CF foundation–approved center
Clinical symptoms (not treated): Persistent diarrhea, malnutrition, chronic cough, respiratory diseases (infections)

Hemoglobinopathies (Sickle Cell Disease)

Prevalance: 1:400 (African Americans)
Substance measured: Hemoglobin fractions (eg, fetal, sickle, adult, hemoglobin C)
Determined by: Electrophoresis and/or high performance liquid chromatography (HPLC)
Abnormal: Detection of hemoglobin(s) other than fetal and adult
Interfering factors: Transfusions will invalidate testing for up to 60 days
Additional diagnostic testing: Hemoglobin detection and quantification on whole blood
Treatment: Penicillin—daily supplements
Clinical symptoms (not treated): Sepsis, pain crisis, death (25% of babies)

Fatty Acid Oxidation (FAO) Disorders

<table>
<thead>
<tr>
<th>Substance measured:</th>
<th>Acylcarnitines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Determined by:</td>
<td>Tandem mass spectrometry</td>
</tr>
<tr>
<td>Abnormal:</td>
<td>Each FAO disorder has a distinctive acylcarnitine profile. The exact profile is program dependent.</td>
</tr>
<tr>
<td>Interfering factors:</td>
<td>False negatives may occur if specimen collection is delayed (&gt;14 days).</td>
</tr>
<tr>
<td>Diagnostic testing:</td>
<td>Consult with a certified biochemical geneticist. Urine organic acids; mutation analysis</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Substance measured:</th>
<th>Fatty Acid Oxidation (FAO) Disorders</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medium-chain acyl-CoA dehydrogenase (MCAD)</td>
<td>1:20,000</td>
</tr>
<tr>
<td>Short-chain acyl-CoA dehydrogenase (SCAD)</td>
<td>1:10,000</td>
</tr>
<tr>
<td>Long-chain 3-hydroxyacyl-CoA dehydrogenase (LCHAD)</td>
<td>1:50,000</td>
</tr>
<tr>
<td>Very-long-chain acyl-CoA dehydrogenase (VLCAD)</td>
<td>1:50,000</td>
</tr>
<tr>
<td>Glutaric acidemia type II (GAI)</td>
<td>1:100,000</td>
</tr>
<tr>
<td>Carnitine palmityltransferase deficiency type II (CPT-II)</td>
<td>Unknown</td>
</tr>
<tr>
<td>2,4-Dienoyl-CoA reductase deficiency</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

Prevalence:
- Medium-chain acyl-CoA dehydrogenase (MCAD) 1:20,000
- Short-chain acyl-CoA dehydrogenase (SCAD) 1:10,000
- Long-chain 3-hydroxyacyl-CoA dehydrogenase (LCHAD) 1:50,000
- Very-long-chain acyl-CoA dehydrogenase (VLCAD) 1:50,000
- Carnitine palmityltransferase deficiency type II (CPT-II) Unknown
- 2,4-Dienoyl-CoA reductase deficiency Unknown
Diagnostic testing: Consult with a certified biochemical geneticist. Urine organic acids; mutation analysis
Treatment: Diet restrictions that are disorder dependent
Clinical symptoms (not treated): Vomiting, lethargy, hypoglycemia, hypotonia; sudden death or permanent neurologic damage can occur

**Galactosemia**

Prevalence: 1:50,000
Substance measured: Total metabolites (galactose and galactose-1-phosphate) and/or galactose-1-phosphate uridyl transferase (GALT)
Determined by: Basic chemistry
Abnormal (cutoff) Elevated metabolites
Typical cutoff range: 10–15 mg/dL or 555–832 µmol/L (program dependent)
No GALT activity
Interfering factors: False negatives may occur if there hasn't been a lactose load before specimen collection. Transfusions may cause false negatives also.
Additional diagnostic testing: Serum/plasma metabolite levels, GALT activity quantification
Treatment: Lactose restriction diet
Clinical symptoms (not treated): Sepsis, milk intolerance; mental retardation, sudden death can occur

**Organic Acidemias**

<table>
<thead>
<tr>
<th>Substance measured:</th>
<th>Acylcarnitines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Determined by:</td>
<td>Tandem mass spectrometry</td>
</tr>
<tr>
<td>Abnormal:</td>
<td>Each OA disorder has a distinctive acylcarnitine profile. The exact profile is program dependent.</td>
</tr>
<tr>
<td>Interfering factors:</td>
<td>False negatives may occur if specimen collection is delayed (&gt;14 days).</td>
</tr>
<tr>
<td>Additional diagnostic testing:</td>
<td>Consult with a certified biochemical geneticist.</td>
</tr>
<tr>
<td>Treatment:</td>
<td>Diet restrictions that are disorder dependent</td>
</tr>
<tr>
<td>Clinical symptoms (not treated):</td>
<td>Vomiting, metabolic acidosis, ketosis, dehydration or coma, hyperammonuria, hypoglycemia, hypotonia, sepsis, developmental delay. Sudden death or permanent neurologic damage can occur.</td>
</tr>
</tbody>
</table>

**Phenylketonuria (PKU)**

Prevalence: 1:15,000
Substance measured: Phenylalanine
Determined by: Basic chemistry or tandem mass spectrometry
Abnormal (cutoff): Elevated phenylalanine: typical cutoff range: 2.0–4.0 mg/dL or 121.1–242.2 µmol/L (program dependent)
Interfering factors: False negatives may occur due to lack of protein.
Diagnostic testing: Serum/plasma amino acid quantification
Treatment: Dietary restriction of phenylalanine
Clinical symptoms (not treated): Mental retardation
Maple Syrup Urine Disease (MSUD)

Prevalence: 1:100,000
Substance measured: Leucine, isoleucine, valine
Determined by: Basic chemistry or tandem mass spectrometry
Abnormal (cutoff): Elevated leucine/isoleucine/valine. Typical cutoff range: 4.0–6.0 mg/dL or 304.9–457.4 µmol/L (program dependent)
Interfering factors: False negatives may occur due to lack of protein.
Diagnostic testing: Serum/plasma amino acid quantification
Treatment: Dietary restrictions of branched-chain amino acids
Clinical symptoms (not treated): Lethargy, vomiting, coma, mental retardation

Homocystinuria

Prevalence: 1:150,000
Substance measured: Methionine
Determined by: Basic chemistry or tandem mass spectrometry
Abnormal (cutoff): Elevated methionine. Typical cutoff range: 1.0–2.0 mg/dL or 67–134 µmol/L (program dependent)
Interfering factors: False negatives may occur due to lack of protein.
Diagnostic testing: Serum/plasma amino acid quantification
Treatment: Dietary restrictions of methionine. Cystine supplementation, folic acid, betaine
Clinical symptoms (not treated): Dislocated lenses, cataracts, muscle weakness, arterial and venous thrombosis, developmental delay

Tyrosinemia

Prevalence: 1:150,000
Substance measured: Tyrosine
Determined by: Basic chemistry or tandem mass spectrometry
Abnormal (cutoff): Elevated tyrosine: typical cutoff range: 4.0–6.0 mg/dL or 220.8–331.0 µmol/L (program dependent)
Interfering factors: False negatives may occur due to lack of protein
Additional diagnostic testing: Serum/plasma amino acid quantification
Treatment: Dietary restrictions of phenylalanine and tyrosine Liver transplants
Clinical symptoms (not treated): Vomiting, diarrhea, renal dysfunction, chronic liver disease, speech delays

Citrullinemia

Prevalence: 1:150,000
Substance measured: Citrulline
Determined by: Tandem mass spectrometry
Abnormal (cutoff): Elevated citrulline: typical cutoff range: 1.0–2.0 mg/dL or 57.1–114.2 µmol/L (program dependent)
Interfering factors: False negatives may occur due to lack of protein.
Additional diagnostic testing: Serum/plasma amino acid quantification Urine—normal levels of argininosuccinic acid
Treatment: Dietary protein restriction of arginine
Clinical symptoms (not treated): Vomiting, lethargy, coma, seizures, anorexia, death

Argininosuccinic Acidemia
Argininosuccinic Acidemia

Prevalence: 1:150,000
Substance measured: Citrulline
Determined by: Tandem mass spectrometry
Abnormal (cutoff): Elevated citrulline: typical cutoff range: 1.0–2.0 mg/dL or 57.1–114.2 µmol/L (program dependent)
Interfering factors: False negatives may occur due to lack of protein.
Additional diagnostic testing: Serum/plasma amino acid quantification
Urine elevations of argininosuccinic acid
Treatment: Dietary restriction of protein; arginine supplement
Clinical symptoms (not treated): Lethargy, coma, progressive neurologic deterioration, atoxia

Clinical Implications

1. Most of the congenital disorders are autosomal recessive genetic disorders (the exception being hypothyroidism). This means that for the baby to have one of these disorders, both the mother and father have to carry the abnormal gene that causes the disorder. In this case, there is a 1-in-4 chance that with each pregnancy, the couple will have an affected baby.
2. Although the symptoms of the various disorders can be quite varied, there are common issues. All the disorders are relatively rare. The most frequently detected disorder is hypothyroidism, which occurs in 1 of 3000 births in most states or regions. Some disorders have frequency rates of 1 in 100,000 or less. All of the disorders, if not detected early and treated promptly, will cause very severe complications. These complications include mental retardation, neurologic problems, or death.
3. All of the disorders can be detected by laboratory tests in the first few days of life before there are any clinical symptoms.
4. If detected early and promptly treated, the baby can develop essentially normally, both mentally and physically.
5. The treatments are relatively simple and inexpensive when compared with lifetime institutional care. For example, several of the metabolic disorders are treated by changes in diet and vitamin supplements.

Interventions

Pretest Patient Care

1. Remember that it is important that the parents be informed about the timing and importance of newborn screening. The state newborn screening programs provide, free of charge, educational brochures regarding newborn screening. Be sure the parents receive this material.
2. Be aware that most states do not require informed consent to perform newborn screening and have limited reasons for parental refusals. If the parent refuses to have the baby screened and the reason is valid under specific state criteria, have the parents sign a waiver for the baby's health care record.
3. Complete the form associated with the newborn screening blood collection kit. Be sure the name on the blood collection card matches the baby whose blood is being drawn.
4. Collect the newborn screening specimen before discharge from the hospital. If the initial specimen was collected before 24 hours of age, obtain a repeat in about 14 days as recommended by the American Academy of Pediatrics.
5. For premature and sick infants, collect an initial specimen as soon as medically possible but no later than the first week of life. Be familiar with your local newborn screening laboratory procedures and policies regarding initial and repeat testing.
6. Collect initial specimen before transfusion, if possible. Be familiar with your local newborn screening laboratory procedures and policies regarding repeat testing when a baby has been transfused.
7. Record the date the specimen was sent to the screening laboratory.

Posttest Patient Aftercare

1. After the testing is complete, the screening laboratory will send a report back to the hospital. Record the receipt and review of the report. The newborn screening laboratory should be contacted if a report is unreasonably delayed (within 10 days of specimen being sent). All reports should be added to the baby's health care records as soon as possible after receipt.
2. Be aware that most newborn screening programs report two types of abnormal results. One type is considered "borderline" or "possible," when the test results are unequivocal or only marginally indicative of a disorder. In most cases, the recommendation is to repeat the newborn screening testing. If the repeat test results are normal, no further action is necessary. Care must be taken to neither alarm the parents nor trivialize the importance of repeat testing. The second type of report issued is when the screening test result is highly indicative of a particular disorder. In this case, the screening laboratory will contact the clinician directly, provide recommendations, and often refer the baby and parents to a specialty clinic for evaluation.
3. Do not institute disorder-specific intervention (eg, diet changes, antibiotics) until directed by clinician after consultation with a specialist. The state newborn screening program can provide contact for appropriate specialty clinics or experts in the disorders.
4. Ensure that genetic counseling is provided.
1. If there is a family history of one of the disorders screened for, notify the newborn screening laboratory so that the specialist can be alerted.
2. The most frequent reason for a retest is that the first specimen was unsatisfactory (inadequate amount of blood or improper use of capillary collection tubes).
3. Be sure the baby gets the newborn screen before hospital discharge or by the seventh day of life for extended hospital stays.
4. Ensure there is a positive correlation between the name written on the blood collection card and the name of the baby being screened.
5. Ensure there is a newborn screening test report in the medical report.
6. Check newborn screening results (including calling the newborn screening laboratory) on babies being readmitted to hospital with severe jaundice, anemic, failure to thrive, seizures, and so forth.
7. Follow up:
   a. Determine whether family of affected children are compliant with appropriate care.
   b. Additional newborn testing may be done (eg, electroencephalogram procedure for evoked auditory response).

**Clinical Alert**

**1. Specific Genetic Tests**

**Genetics of Type 1 and Type 2 Diabetes Mellitus** The cause of diabetes mellitus is deficient insulin action (insulin action is equal to the product of insulin concentration [B-cell control] and insulin sensitivity [target cell function]). Deficient insulin action leads to disordered carbohydrate, lipid, and protein metabolism. Type 1 results from insulin deficiency; type 2 results from a combination of insulin resistance and relative insulin deficiency. **Type 1:** Autoimmune diabetes may result from an interaction of genetics and the environment and results in an absolute insulin deficiency. Type 1 diabetes resulting from autoimmune destruction of pancreatic B cells is not inherited, but susceptibility to type 1 disease is. The major genetic loci indicative of susceptibility to type 1 is located in the HLA complex: DRB1, DQA1, and DQB1. **Type 2:** Patients with type 2 diabetes (due to insulin resistance and B-cell failure) often have a first-degree relative with the disease and a genetic predisposition to type 2 resulting in a restricted ability of the B cells to secrete insulin. Type 2 diabetes mellitus is inherited as a dominant gene, although not all cases are hereditary. Persons at risk include those with a family history and those who develop gestational diabetes.

**Genetics of Hereditary Hemochromatosis (Primary Iron Overload)** Hemochromatosis is a relatively common disorder that leads to liver dysfunction at 40 to 50 years of age; it is not a rare genetic disease as once thought. Prevalence in the United States is 1 in 200 Caucasians (possibly also other ethnic groups). This disorder, treated by phlebotomy at regular intervals, is fatal if not diagnosed early. At the present time, genotyping is done of the HFE gene to include the C282Y and H63D mutations.

**Biochemical Genetics** Testing for hereditary metabolic disorders identifies inborn errors of metabolism (IEM) and enzyme disorders. Included are amino acids, carbohydrates, cholesterol, cofactors and vitamins, lysosomal shortage, lactic acids, fatty acids, carnitines, organic acids, porphyrins, purines and pyrimidine, and urea. Molecular genetics includes the diagnosis of neoplastic disorders (eg, Philadelphia chromosome and neuroblastoma) and the carrier identification and prenatal diagnosis of various inherited disorders (eg, thalassemia, cystic fibrosis, hemophilia A). In general, it is the mature tissue product of a gene that carries out its function.

**Population Genetics** Population genetics is the study of genes in populations and of factors that maintain or change the frequency of genes and genotypes from generation to generation. Multifactorial inherited disease deals with traits or diseases not inherited in factors believed to play an important role in causation; examples of these disorders are hypertension, schizophrenia, diabetes mellitus, and common birth defects such as cleft lip, cleft palate, and neural tube defects.

**Treatment of Genetic Diseases** Environmental therapy to restrict potentially toxic agents, use of diet or medications, and gene therapy are the main treatments for genetic diseases. The following is a brief summary of the therapies.

1. **Dietary and drug restrictions.** Examples include cholesterol and saturated fats in familial hypercholesterolemia, antimalarial drugs and certain antibiotics in glucose-6-phosphate dehydrogenase deficiency; in alpha -antitrypsin deficiency, no cigarette smoking; evidence of alcohol in hemochromatosis.
2. **Replacement therapy.** Examples include factor VIII in hemophilia A, liver transplantation in homozygous hypercholesterolemia, bone marrow transplantation in sickle cell anemia, and kidney transplantation in cystinosis.
3. **Removal of toxic agents.** Examples include abnormal copper accumulation treated by penicillin phlebotomy to remove iron in hemochromatosis, and statins (eg, Lovastatin) to treat and lower cholesterol in familial hypercholesterolemia.
4. **Gene therapy.** Gene therapy is the treatment of disease by transfer of genetic material (DNA or RNA) into the patient. Gene therapy interventions are based on the type of tissue involved. In germline tissue therapy, the gene modification would be introduced into all cells (including ovaries and testes), resulting in the modified gene passed on to subsequent generations, but this type of therapy is not currently done.
5. **Somatic gene therapy.** The genetic modification is restricted to somatic cells (not into ovaries and testes) and does not transmit genetic alterations to subsequent generations. It is in this area that most research is currently involved. Difficulties with delivery of the gene to target cells and obtained adequate expression are abstracts to use of gene therapy in humans.

**Pharmacogenomics** Pharmacogenomics studies genetic variations and drug metabolism to match the best drug for phenotype (in specific diseases) before beginning therapy. Matching effective drugs to DNA-based diagnostic and predictive markers to individualize treatment. Examples of the forefront of treatment and diagnostic test companies include: diagnostic tests, genetic markers and drug targets for schizophrenia, hemochromatosis, peripheral arterial occlusive disease, rheumatoid arthritis, obesity, severe anxiety, stroke, type 2 diabetes (Roche and deCode Genetics Co.), progression of advanced heart disease (Pharmacia and deCode Genetic Companies), obesity and diabetes (Bayer Corp. and CuraGen Corp.), and markers for colon, breast, and ovarian cancer.
BIBLIOGRAPHY


DeMay RM: Practical Principles of Cytopathology. Chicago, ASCP Press, 1999


Digene Corp.: Clinician Q & As. January 2003. (Online.) Accessible at http://www.digene.com/clinician_3_1.html


Kiesner F: Oncotech announces collaborative research with Southwest Oncology Group (SWOG) to evaluate the recently patented Oncotech radiation resistance assay as a predictor of clinical outcome in cervical carcinoma patients. Press Release, November 9, 2001

Kiesner F: Oncotech and the Johns Hopkins Cancer Center Institute research collaboration to eliminate unnecessary exploratory surgery. Press Release, November 2, 2001


Titus K: Breast specimens: FNA, core, more. CAP Today, 16(2):2002


OVERVIEW OF ENDOSCOPIC STUDIES

Clinical Alert

Mediastinoscopy

Bronchoscopy

Thoracoscopy

Esophagogastroduodenoscopy (EGD); Upper Gastrointestinal (UGI) Study; Endoscopy; Gastroscopy

Esophageal Manometry

Endoscopic Retrograde Cholangiopancreatography (ERCP) and Manometry

Enteroscopy

Colposcopy

Loop Electrosurgical Excision Procedure (LEEP), Cone Biopsy, Cervical Conization

Flexible Proctoscopy; Sigmoidoscopy; Proctosigmoidoscopy

Colonoscopy

Peritoneoscopy; Laparoscopy; Pelviscopy; Fertiloscopy

Cystoscopy (Cystourethroscopy)

URODYNAMIC STUDIES

Cystometrogram (CMG); Urethral Pressure Profile (UPP); Rectal Electromyogram (EMG); Cystourethrogram

Arthroscopy

Clinical Alert

Sinus Endoscopy

BIBLIOGRAPHY

OVERVIEW OF ENDOSCOPIC STUDIES

Endoscopy is the general term given to all examination and inspection of body organs or cavities using endoscopes. These instruments can also provide access for certain kinds of surgical procedures or treatments. Endoscopes, known generally as fiberoptic instruments, are used for direct visual examination of certain internal body structures by means of a lighted lens system attached to either a rigid or flexible tube. The newest instruments transmit signals from the tip of the scope to a TV monitor. Light travels through an optic fiber by means of multiple reflections. Fiberoptic instruments, composed of fiber bundle systems, redirect and transmit light around twists and bends in cavities and hollow organs of the body. An image fiber and a light fiber allow visualization at the distal tip of the scope. Separate ports allow instillation of drugs, lavage, suction, and insertion of a laser, brushes, forceps, or other instruments used for excision, sampling, or other diagnostic and therapeutic procedures. The flexible scope can be inserted into orifices or other areas of the body not easily accessible or directly visualized by rigid scopes or other means. Procedures are done for health screening, diagnosis of pathologic conditions, or therapy, such as removal of tissue (polyps) or foreign objects. Sedatives or analgesia (to achieve a state of conscious sedation) or local or general anesthetics may be used. The use of video documentation and endoscopic sonography also aid in cancer diagnosis, staging of cancer, and operability. Biopsy tissue is submitted to the laboratory for histologic examination (see Chap. 11).

Clinical Alert

Endoscopically related bacteremia infections may result from tissue manipulations, blood stream invasion by pathogens, or a contaminated endoscope, usually due to improper cleansing and disinfection. After endoscopic procedures, assess for fever, elevated white blood cells, signs of bloodstream infection, and signs of sepsis. It is important that strict infection control guidelines be followed by persons who clean and disinfect the endoscopes. Hospitals and clinics should follow the infection control policy for their institution, which should include documentation of all endoscopic procedures, including name of patient, type of procedure, date and time of procedure, and serial number of the endoscope used in each procedure. A log documenting the time, date, and serial number of each endoscope cleaned and disinfected should also be maintained. These records allow for tracing an infection back to a specific instrument. Any infections suspected to have been caused by a contaminated instrument should be reported immediately to the appropriate infection control and risk management departments for investigation.

Clinical Alert

1. Observe standard precautions and latex precautions for all endoscopic procedures. See Appendix A and Appendix B.
2. Endoscopic procedures have not proved useful in U.S., but screenings are helpful in Japan and China where death and cure rates of esophageal and stomach cancer are improved by endoscopic detection.
3. Some investigators and clinicians have concerns about tissue damage, immunosuppression, and postsite metastases after endoscopic procedure (DeVita, Hellman, Rosenberg, 2001, pp. 740–741).

Mediastinoscopy

Mediastinoscopy, performed under general anesthesia, requires insertion of a lighted mirror-lens instrument, similar to a
bronchoscope, through an incision at the base of the anterior neck, to examine and biopsy mediastinal lymph nodes. Because these nodes receive lymphatic drainage from the lungs, mediastinal biopsy specimens can allow identification of diseases such as carcinoma, granulomatous infection, sarcoidosis, coccidioidomycosis, or histoplasmosis. Mediastinoscopy is used to stage lung tumors, diagnose sarcoidosis, biopsy mediastinal lymph nodes directly, and assess hilar adenopathy of unknown origin. It has virtually replaced scalene fat pad biopsy for examining suspicious nodes on the right side of the mediastinum. It is the routine method of establishing tissue diagnosis and staging of lung cancer and for evaluating the extent of lung tumor metastasis done just before thoracotomy. Nodes on the left side of the chest are usually resected through left anterior thoracotomy (mediastinoscopy) or occasionally by scalene fat pad biopsy. This procedure is performed by a thoracic surgeon.

Reference Values
Normal No evidence of disease Normal lymph glands

Procedure
1. Mediastinoscopy is considered a surgical procedure and is usually performed under general anesthesia in a hospital.
2. Biopsy is performed through a suprasternal incision in the neck (2–3 cm or 3–4 cm), just above the sternal notch. When the Chamberlain procedure is performed, a small transverse incision is done in the second intercostal space or over the 2nd or 3rd costal cartilage.
3. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications
1. Abnormal findings may include the following conditions:
   a. Sarcoidosis
   b. Tuberculosis
   c. Histoplasmosis
   d. Hodgkin's disease
   e. Granulomatous infections and inflammatory processes
   f. Carcinomatous lesions
   g. Coccidioidomycosis
   h. Pneumocystis carinii infection
2. Results assist in defining the extent of metastatic process, staging of cancer (N2 and N3–IIIa and IIIb), and possibility of successful surgical resectability.

Interventions
Pretest Patient Preparation
1. Explain purpose, procedure, benefits, and risks of the test. It is usually used after CT scan and indicates enlarged mediastinal nodes (>1 cm).
2. Be aware that a legal surgical consent form must be appropriately signed and witnessed preoperatively (see Chap. 1).
3. Remember that preoperative care is the same as that for any patient undergoing general anesthesia and surgery.
4. Have the patient fast for 8 or more hours before the test.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare Care is the same as for any patient who has had surgery under general anesthesia.
1. Evaluate breathing and lung sounds; check wound for bleeding and hematoma.
2. At time of discharge, monitor for complications (eg, breathing difficulties, coughing up blood). Instruct the patient to call physician if problems occur.
3. Interpret test outcomes, monitor appropriately, and explain any need for follow-up tests and/or treatment (medication for TB, antibiotics).
4. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Clinical Alert
1. Previous mediastinoscopy contraindicates repeat examination because adhesions make satisfactory dissection of nodes extremely difficult or impossible.
2. Complications can result from the risks associated with general anesthesia and from preexisting conditions, pneumothorax, and subcutaneous emphysema.
3. Damage to major vessels can occur during this procedure.

Bronchoscopy
Bronchoscopy permits visualization of the trachea, bronchi, and select bronchioles. There are two types of bronchoscopy: flexible fiberoptic (Fig. 12.1), which is almost always used for diagnostic purposes, and rigid, which is less frequently used. This procedure is done to diagnose tumors, coin lesions, or granulomatous lesions; to find hemorrhage sites; to evaluate trauma or nerve paralysis; to obtain biopsy specimens; to take brushings for cytologic examinations; to improve drainage of secretions; to identify inflammatory infiltrates; to lavage; and to remove foreign bodies. Bronchoscopy can determine resectability of a lesion as well as provide the means to diagnose bronchogenic carcinoma. A transbronchial needle biopsy may be performed during this procedure, thus obviating the need for diagnostic open-lung biopsy. A flexible needle is passed through the trachea or bronchus and is used to aspirate cells
from the lung. This procedure is performed on patients with suspected sarcoidosis or pulmonary infection.

**Indications**

1. **Diagnostic:**
   a. Staging of bronchogenic carcinoma
   b. Differential diagnosis in recurrent unresolved pneumonia
   c. Evaluation of cavitary lesions, mediastinal masses, and interstitial lung disease
   d. Localization of bleeding and occult sites of cancer
   e. Evaluate immunocompromised patients (eg, human immunodeficiency virus [HIV]-infected patients, bone marrow or lung transplant recipients)
   f. Differentiate rejection from infection in lung transplantation
   g. Assess airway damage in thoracic trauma
   h. Evaluate underlying etiology of nonspecific symptoms of pulmonary disease such as chronic cough (>6 months), hemoptysis, or unilateral wheezing

2. **Therapeutic:**
   a. Removal of mucus plugs and polyps
   b. Removal of an aspirated foreign body and to relieve endobronchial obstruction
   c. Brachytherapy (radioactive treatment of malignant endobrachial tumors)
   d. Placement of a stent to maintain airway patency
   e. Drainage of lung abscess
   f. Decompression of bronchogenic cysts
   g. Laser photoresection of endotracheal lesions
   h. Bronchoalveolar lavage to remove intraalveolar proteinaceous material
   i. Alternative for difficult endotracheal intubations
   j. Control bleeding and airway hemorrhage in the presence of massive hemoptysis
   k. Debridement of tumors using newer drugs, eg, Photofren II (to achieve a state of conscious sedation)

The examination is usually done under local anesthesia combined with some form of sedation in an outpatient setting, diagnostic center, or operating room. It also can be done in a critical care unit, in which case the patient may be unresponsive or ventilator dependent.

**Reference Values**

**Normal** Normal trachea, bronchi, nasopharynx, pharynx, and select bronchioles (conventional bronchoscopy cannot visualize alveolar structures)

**Procedure**

1. Spray and swab topical anesthetic (eg, 4% lidocaine) onto the back of the nose, the tongue, the pharynx, and the epiglottis. Give an antialagogue (eg, atropine) to reduce secretions. If the patient has a history of bronchospasms, administer a bronchodilator (eg, albuterol) via a hand-held nebulizer.

2. Insert the flexible or rigid bronchoscope carefully through the mouth or nose into the pharynx and the trachea (Fig. 12.2). The scope also can be inserted through an endotracheal tube or tracheostomy. Suctioning, oxygen delivery, and biopsies are accomplished through bronchoscope ports designed for these purposes.

3. Be advised that because of sedation, usually with diazepam (Valium), midazolam (Versed), or meperidine (Demerol), the patient is usually comfortable when a state of conscious sedation is achieved. However, when the bronchoscope is advanced, some patients may feel as if they cannot breathe or are suffocating.

4. The right lung, by convention, is normally examined before the left lung.

5. Bronchoscopic procedures include any one or a combination of the following:
   a. Bronchial washings for cytology and staining for fungi and mycobacteria
   b. Bronchoalveolar lavage (BAL) for infectious (eg, alveolar proteinosis, eosinophilic granuloma) diseases
   c. Bronchial brushings of both visible and peripheral (under fluoroscopy) endobronchial lesions and/or transbronchial biopsies, both visible and peripheral

6. Follow guidelines in Chapter 1 for **safe, effective, informed intratest care**.
Clinical Alert

1. Morphine sulfate is contraindicated in patients who have problems with bronchospasm or asthma because it can cause bronchospasm. Analgesics, barbiturates, tranquilizers-sedatives, and atropine may be ordered and administered 30 minutes to 1 hour before bronchoscopy. The patient should be as relaxed as possible before and during the procedure but also needs to know that anxiety is normal. The patient may need additional intravenous sedatives during the procedure. Refer to conscious sedation precautions in Appendix C.
   a. Arterial blood gas measurement during and after bronchoscopy may be ordered, and arterial blood oxygen may remain altered for several hours after the procedure. Sputum specimens taken during and after bronchoscopy may be sent for cytologic examination or culture and sensitivity testing. These specimens must be handled and preserved according to institutional protocols (see Chap. 14).
2. Continuous monitoring of electrocardiogram, blood pressure, pulse oximetry, and respirations is routinely performed. Monitoring of pulse oximetry is especially important to indicate levels of oxygen saturation before, during, and after the procedure.

Clinical Implications Abnormalities revealed through bronchoscopy include the following conditions:

1. Abscesses
2. Bronchitis
3. Carcinoma of the bronchial tree (occurs in the right lung more often than the left)
4. Tumors (usually appear more often in larger bronchi)
5. Tuberculosis
6. Alveolitis
7. Evidence of surgical nonresectability (eg, involvement of tracheal wall by tumor growth, immobility of a main-stem bronchus, widening and fixation of the carina)
8. Pneumocystis carinii infection
9. Inflammatory processes
10. Cryptococcus neoformans infection
11. Histoplasmosis
12. Blastomycosis
13. Aspergillosis
14. Coccidioidomycosis
15. Cryptococcus neoformans infection
16. Coccidioidomycosis
17. Histoplasmosis
18. Blastomycosis
19. Phycomycosis

Clinical Considerations The following data must be available before the procedure: history and physical examination, recent chest x-ray film, recent arterial blood gas values, and, if the patient is > 40 years of age or has heart disease, electrocardiogram (ECG). Appropriate blood work (coagulation), urinalysis, pulmonary function tests, and sputum studies (especially for acid-fast bacilli) must be done as well. Bronchoscopy is often done as an ambulatory surgical procedure.

Interventions

Pretest Patient Preparation for Traditional Bronchoscopy

1. Reinforce information related to the purpose, procedure, benefits, and risks of the test. Record signs and symptoms (eg, dyspnea, bloody sputum, coughing, hoarseness).
2. Emphasize that pain is not usually experienced because lungs do not have pain fibers.
3. Explain that the local anesthetic may taste bitter, but numbness will occur in a few minutes. Feelings of a thickened tongue and the sensation of something in the back of the throat that cannot be coughed out or swallowed are not unusual. These sensations will pass within a few hours following the procedure as the anesthetic wears off.
4. Be aware that an informed consent form must be properly signed and witnessed (see Chap. 1).
5. Have the patient fast for at least 6 hours before the procedure to reduce the risk for aspiration. Gag, cough, and swallowing reflexes will be blocked during and for a few hours after surgery.
6. Ensure that the patient removes wigs, nail polish, makeup, dentures, jewelry, and contact lenses before the examination.
7. Use relaxation techniques to help the patient relax and breathe more normally during the procedure. The more relaxed the patient is, the easier it is to complete the procedure.
8. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare for Traditional Bronchoscopy

1. Be certain that swallow, gag, and cough reflexes are present before allowing food or liquids to be ingested orally. Usually the patient has fasted for at least 2 hours before the procedure.
2. Provide gargles to relieve mild pharyngitis. Monitor ECG, blood pressure, temperature, pulse, pulse oximeter readings, skin and nail bed color, lung sounds, and respiratory rate and patterns according to institution protocols. Document observations.
3. Be aware that the following may be ordered:
   a. Oxygen by mask or nasal cannula. Humidified oxygen at specific concentrations up to 100% by mask may be necessary.
   b. A chest x-ray film. This will check for pneumothorax or to evaluate the lungs.
   c. Sputum specimens. These must be preserved in the proper medium or solution.
4. Elevate the head of the bed for comfort.
5. Interpret test outcomes, monitor appropriately, and explain need for other tests or treatment. Follow-up procedures may be necessary. Computed tomography (CT)-guided fine-needle cytology aspiration may be done when bronchoscopy is not diagnostic.
6. Refer to intravenous sedation precautions in Appendix C.
7. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Contraindications to Bronchoscopy**

Contraindications to bronchoscopy include the following conditions:

1. Severe hypoxemia
2. Severe hypocapnia (carbon dioxide retention)
3. Certain cardiac arrhythmias, cardiac states
4. History of being hepatitis B carrier
5. Bleeding or coagulation disorders
6. Severe tracheal stenosis

**Clinical Alert**

A. Observe for possible complications of traditional bronchoscopy, which may include the following conditions:

1. Shock
2. Bleeding following biopsy (rare, but can occur if there is excessive friability of airways or massive lesions, or if patient is uremic or has a hematologic disorder)
3. Hypoxemia
4. Partial or complete laryngospasm (inspiratory stridor) that produces a "crowing" sound; may be necessary to intubate
5. Bronchospasm (pallor and increasing dyspnea are signs)
6. Infection or gram-negative bacterial sepsis
7. Pneumothorax
8. Respiratory failure
9. Cardiac arrhythmias
10. Anaphylactic reactions to drugs
11. Seizures
12. Febrile state
13. Hypoxia, respiratory distress
14. Empyema
15. Aspiration

B. Virtual noninvasive bronchoscopy using spinal CT technology requires no sedation or analgesics. Indications include pulmonary embolism and staging of lung cancer.

**Special Pediatric Considerations**

Bronchoscopy instruments can decrease an already small airway lumen even more by causing inflammation and edema. Consequently, a child can rapidly become hypoxic and desaturate oxygen very quickly. Resuscitation, oxygen administration equipment, and drugs must be readily accessible when this procedure is performed on a child. Close monitoring of respiratory and cardiac status is imperative during and after the procedure. The same precautions and treatment apply to children and adults. Most children suffer cardiac arrest because of respiratory problems, not cardiac problems.

**Thoracoscopy**

Thoracoscopy is an examination of the thoracic cavity using an endoscope. Video-assisted thoracoscopy (VAT) is a recent addition to the procedures available for diagnosing intrathoracic diseases. This procedure is making a comeback because it can be used as a diagnostic device when other methods of diagnosis fail to present adequate and accurate findings. Moreover, the discomfort and many of the risks associated with traditional diagnostic thoracotomy procedures are reduced with thoracoscopy versus other procedures. Thoracoscopy allows visualization of the parietal and visceral pleura, pleural spaces, thoracic walls, mediastinum, and pericardium without the need for more extensive procedures. It is used most frequently to investigate pleural effusion and can be used to perform laser procedures; diagnose and stage lung disease; assess tumor growth, pleural effusion, emphysema, inflammatory processes, and conditions predisposing to pneumothorax; and perform biopsies of pleura, mediastinal lymph nodes, and lungs.

**Reference Values**

**Normal** Thoracic cavity and tissues normal and free of disease

**Procedure**

1. Be advised that thoracoscopy is considered an operative procedure. The patient's state of health, the particular positioning needed, and the procedure itself determine the need for either local or general anesthesia. The incision is usually made at the midaxillary line and the sixth intercostal space.
2. Schedule admission the morning of the procedure. Many patients are discharged the following day, provided the lung has reexpanded properly and chest tubes have been removed.
3. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

**Clinical Implications**

Abnormal findings can include the following conditions:

1. Carcinoma or metastasis of carcinoma
2. Empyema
3. Pleural effusion
4. Conditions predisposing to pneumothorax or ulcers
5. Inflammatory processes
6. Bleeding sites
7. Tuberculosis, coccidioidomycosis, or histoplasmosis

**Interventions**
Pretest Patient Preparation

1. Reinforce and explain the purpose, procedure, benefits, and risks of the examination and describe what the patient will experience. Record preprocedure signs and symptoms.
2. Be aware that a surgical consent form must be appropriately signed and witnessed before the procedure begins (see Chap. 1).
3. Complete and review required blood tests, urinalysis, recent chest x-ray film, and ECG (for certain individuals) before the procedure.
4. Have the patient fast for 8 hours before the procedure.
5. Insert an intravenous line for the administration of intraoperative intravenous fluids and intravenous medication.
6. Perform skin preparation and correct positioning in the operating room.
7. Place a chest tube and connect to negative suction or sometimes to gravity change after the thoracoscopy is completed.
8. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Take a postoperative chest x-ray film to check for abnormal air or fluid in the chest cavity.
2. Monitor vital signs, amount and color of chest tube drainage, fluctuation of fluid in the chest tube, bubbling in the chest bottle, and respiratory status, including arterial blood gases. Promptly report abnormalities to the physician.
3. Administer pain medication as necessary. Encourage relaxation exercises as a means to lessen the perception of pain. Monitor quality and rate of respirations. Be alert to the possibility of respiratory depression related to narcotic administration or intrathecal narcotics.
4. Encourage frequent coughing and deep breathing. Assist the patient in splinting the incision during coughing and deep breathing to lessen discomfort. Promote leg exercises while in bed and assist with frequent ambulation if permitted.
5. Use open-ended questions to provide the patient with an opportunity to express concerns.
7. Interpret test outcomes and monitor appropriately.
8. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care. Provide written discharge instructions.

Clinical Alert

1. Do not clamp chest tubes unless specifically ordered to do so. Clamping chest tubes may cause a tension pneumothorax. Sudden onset of sharp pain, dyspnea, uneven chest wall movement, tachycardia, anxiety, and cyanosis may indicate pneumothorax. Notify the physician immediately.
2. Possible wound and pulmonary complications include the following:
   a. Acute respiratory distress, hypoxia
   b. Infection
   c. Hemorrhage (watch for unusually large outputs of blood in a relatively short period of time into the chest bottle and notify physician immediately)
   d. Empyema
   e. Atelectasis
   f. Aspiration
3. Nerve damage may occur during the procedure.

Esophagogastroduodenoscopy (EGD); Upper Gastrointestinal (UGI) Study; Endoscopy; Gastroscopy

Endoscopy is a general term for visual inspection of any body cavity with an endoscope. Endoscopic examination of the upper gastrointestinal (UGI) tract (mouth to upper jejunum) is referred to when the following examinations are ordered: panendoscopy, esophagoscopy, gastroscopy, duodenoscopy, esophagastroscope, or esophagogastroduodenoscopy (EGD).

Esophagogastroduodenoscopy allows direct visualization of the interior lumen of the upper gastrointestinal tract with a fiberoptic instrument designed for that purpose. EGD is indicated for patients with dysphagia; reflux symptoms; weight loss; hematemesis; melena; persistent nausea and vomiting; persistent epigastric, abdominal, or chest pain; and persistent anemia. EGD can confirm suspicious x-ray findings and establish a diagnosis of symptomatic patients with negative x-ray reports. EGD can be used to diagnose and treat many abnormalities of the UGI tract, including hernias, gastroesophageal reflux disease (GERD), esophagitis, gastritis, strictures, varices, ulcers, polyps, and tumors. It can be used to remove foreign bodies (such as a swallowed coin in a small child) and for placement of a percutaneous gastric or duodenal feeding tube. For patients who require some form of UGI surgery, it provides a safe way to perform presurgical screening and postsurgical surveillance.

Reference Values

Normal UGI tract within normal limits

Procedure

1. Remember that this examination is usually performed in an outpatient setting of a hospital or ambulatory clinic. It also may be performed in the operating room or in a critical care setting.
2. Use a topical spray to anesthetize the patient's throat.
3. Start an intravenous line and use for administration of sedation alone or in combination with analgesics. These medications are given to achieve a state of conscious sedation (see Appendix C). Resuscitation equipment must be available.
4. Perform continuous monitoring of the patient’s vital signs, ECG, and oxygen saturation (pulse oximetry).
5. Remove partial dental plates or dentures. Insert a mouthpiece to prevent the patient from biting the endoscope and to prevent injury to the patient's teeth, tongue, or other oral structures.
6. Lubricate the endoscope well. Gently insert through the mouthpiece into the esophagus and advance slowly into the stomach and duodenum. Insufflate air through the scope to distend the area being examined so that optimal visualization of the mucosa is possible. Obtain tissue biopsy specimens and brushings for cytology. Take photos to provide a permanent record of observations.
7. Inform the patient that he or she may have an initial gagging sensation that quickly subsides. During the procedure, the patient may belch frequently. Sensations of abdominal pressure or bloating are normal, but the patient should not experience actual pain.
8. Immediately after the examination is completed, ask the patient to remain on his or her left side until fully awake.

Clinical Implications Abnormal results may indicate the following conditions:

1. Hemorrhagic areas or erosion of an artery or vein
2. Hiatal hernia
3. Esophagitis, gastritis
4. Neoplastic tissue
5. Gastric ulcers (benign or malignant)
6. Esophagitis, gastritis, duodenitis
7. Esophageal or gastric varices
8. Esophageal, pyloric, or duodenal strictures

Interventions

Pretest Patient Preparation

1. Explain the purpose and procedure of the examination, the sensations that may be experienced, and the benefits and risks of the test. Refer to intravenous conscious sedation precautions in Appendix C. Reassure the patient that the endoscope is thinner than most food swallowed. Inform the patient that he or she may be quite sleepy during the EGD and may not recall much or any of the experience. Record preprocedure signs and symptoms (eg, vomiting, melena, dysphagia, and persistent upper GI pain).
2. Patients should be instructed to fast before the procedure, according to the hospital or clinic policy. Generally, adult patients should fast 6–8 hours before the examination, and children may have clear liquids up until 2 hours before the procedure; however, each patient should be assessed on an individual basis, according to age, size, and general health status. Inpatients may have intravenous fluids to prevent dehydration. Outpatients need education about potential risks for aspiration and possible cancellation of the procedure if fasting is not maintained.
3. Confirm informed consent. A legal consent must be signed and witnessed before the procedure.
4. Encourage the patient to urinate and defecate if possible before the examination.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Do not permit food or liquids until the patient's gag reflex returns.
2. Monitor blood pressure, pulse, respirations, and oxygen saturation according to the hospital or clinic policy, usually every 15–30 minutes, until the patient is fully awake.
3. Ask the patient to remain on his or her left side with side-rails raised until fully awake. This position usually prevents aspiration.
4. Encourage the patient to belch or expel air inserted into the stomach during the examination.
5. Remember that the patient should not experience discomfort or side effects once the sedative has worn off. Occasionally, the patient may complain of a slight sore throat. Sucking on lozenges after swallowing reflexes return may be helpful if these are permitted.
6. Interpret test outcomes and monitor appropriately.
7. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

Complications are rare; however, the following complications can occur:

1. Perforation
2. Bleeding or hemorrhage
3. Aspiration
4. Infection
5. Complications from drug reaction (leading to hypotension, respiratory depression or arrest, allergic or anaphylactic response)
6. Complications from unrelated diseases (such as myocardial infarction, cerebrovascular accident)
7. Death (very rare)

Esophageal Manometry

Esophageal manometry measures the movement, coordination, and strength of esophageal peristalsis as well as the function of the upper and lower esophageal sphincters. The test consists of recording intraluminal pressures at various levels in the esophagus and at the upper and lower esophageal sphincters. Intraluminal pressures can be measured with the use of a manometric catheter, which is passed intranasally to the patient and then attached to an infusion pump, transducer, and recorder. The intraluminal pressures produce waveform readings (somewhat similar to ECG readings),
which can be used to assess esophageal function.

**Indications for Testing**

1. Abnormal esophageal muscle function
2. Difficulty swallowing (dysphagia)
3. Heartburn
4. Noncardiac chest pain
5. Regurgitation
6. Vomiting
7. Esophagitis

Another test, often done in conjunction with manometry, is the Bernstein test (discussed later). This procedure is useful for evaluating heartburn, esophagitis, and noncardiac chest pain.

**Reference Values**

**Normal**
- Normal esophageal and stomach pressure readings
- Normal contractions
- No acid reflux

**Procedure**

1. Remember that the examination is usually performed in an outpatient setting, such as an ambulatory clinic or physician's office.
2. Attach the manometric catheter to the infusion pump. Set up the transducer and recording equipment and calibrate according to manufacturer's recommendations.
3. Assess the patient's nasal passage for adequate size and patency. Generously apply a topical anesthetic to the selected nostril.
4. Lubricate the manometric catheter and pass it through the nostril, down the esophagus, and just below the lower esophageal sphincter with the patient in a sitting position. Facilitate this with the patient drinking sips of water through a straw.
5. Begin recording. Pull the catheter through the lower esophageal sphincter, then the esophageal body, and finally the upper esophageal sphincter. Different techniques may be used to obtain recordings. The patient may be asked to swallow, not swallow, take sips of water, or hold his or her breath while the catheter is pulled through.
6. Be aware that the **Bernstein test** evaluates for acid reflux by means of a nasogastric tube passed to a point 5 cm above the gastroesophageal junction. Concentration of hydrochloric acid (0.1 normal HCl) is infused for 10 minutes into the esophagus to reproduce symptoms of heartburn or chest discomfort. In the first 5 minutes of testing, 0.9% sodium chloride (NaCl) is infused as a control. Testing takes about 15 minutes. The patient may lie down or sit up.
7. Follow guidelines in Chapter 1 regarding **safe, effective, informed intratest care**.

**Clinical Implications**

Abnormal recordings reveal the following conditions:

1. Primary esophageal motility disorders, such as achalasia, nutcracker esophagus, or diffuse esophageal spasm.
2. Hypertensive lower esophageal sphincter
3. Acid reflux

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose, procedure, benefits and risks of the test.
2. Obtain an informed consent that is properly signed and witnessed.
3. Confirm that the patient has fasted for 6 hours before testing.
4. Instruct the patient on the techniques of swallowing, sipping water, and so forth to facilitate accurate recordings.
5. Follow guidelines in Chapter 1 regarding **safe, effective, informed pretest care**.

**Posttest Patient Aftercare**

1. Advise the patient that a sore throat and nasal passage irritation are common for 24 hours after the examination. Sensations of heartburn may also persist. Administer antacids if ordered.
2. Observe for or instruct patient to watch for nasal bleeding, gastrointestinal bleeding, or unusual pain.
3. Interpret test outcomes, counsel, and monitor appropriately as above.
4. Follow guidelines in Chapter 1 for **safe, effective, informed posttest care**. Provide written discharge instructions.

**Clinical Alert**

Complications are rare; however, the following can occur: aspiration; perforation of nasopharynx, esophagus, or stomach; epistaxis.

**Endoscopic Retrograde Cholangiopancreatography (ERCP) and Manometry**

This examination of the hepatobiliary system is done through a side-viewing flexible fiberoptic endoscope by instillation of contrast medium into the duodenal papilla, or ampulla of Vater. This allows for radiologic visualization of the biliary and pancreatic ducts. It is used to evaluate jaundice, pancreatitis, persistent abdominal pain, pancreatic tumors, common duct stones, extrahepatic and intrahepatic biliary tract disease, malformation, and strictures and as a follow-up study in confirmed or suspected cases of pancreatic disease.

ERCP manometry can be done to obtain pressure readings in the bile duct, pancreatic duct, and sphincter of Oddi at the papilla. Measurements are obtained using a catheter that is inserted into the endoscope and placed within the sphincter.
Reference Values

Normal appearance and patent pancreatic ducts, hepatic ducts, common bile ducts, duodenal papilla (ampulla of Vater), and gallbladder Manometry: Normal pressure readings of bile and pancreatic ducts and sphincter of Oddi

Procedure

1. Remember that this examination is usually performed in a hospital or outpatient setting where fluoroscopy and x-ray equipment are available.
2. Have the patient gargle with or spray his or her throat with a topical anesthetic.
3. Start an intravenous line and use for administration of sedatives and analgesics. These medications are given to achieve a state of conscious sedation (see Appendix C). In some situations, general anesthesia may be used. Resuscitation equipment must be available.
4. Perform continuous monitoring of the patient's vital signs, ECG, and oxygen saturation (pulse oximetry).
5. Remove partial dental plates or dentures. Insert a mouthpiece to prevent the patient from biting the endoscope and to prevent injury to the patient's teeth, tongue, or other oral structures.
6. Have the patient assume a left lateral position with the knees flexed. The endoscope is well lubricated and inserted via the mouthpiece, down the esophagus and stomach, and into the duodenum. At this point, have the patient assume a prone position with the left arm positioned behind him or her.
7. Instill simethicone to reduce bubbles from bile secretions. Give glucagon or anticholinergic agents intravenously to relax the duodenum so that the papilla can be cannulated. (Atropine increases the heart rate.)
8. Pass a catheter into the ampulla of Vater and instill a contrast agent through the cannula to outline the pancreatic and common bile ducts. Perform fluoroscopy and x-rays at this time.
9. Take biopsy specimens or cytology brushings before the endoscope is removed.
10. Monitor for side effects and drug allergy reactions (eg, diaphoresis, pallor, restlessness, hypotension).
11. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications

Abnormal results reveal stones, stenosis, and other abnormalities that are indicative of the following conditions:

1. Biliary cirrhosis
2. Primary sclerosing cholangitis
3. Cancer of bile ducts, gallstones
4. Pancreatic cysts
5. Pseudocysts
6. Pancreatic tumors
7. Cancer of head and pancreas
8. Chronic pancreatitis
9. Pancreatic fibrosis
10. Cancer of duodenal papilla
11. Papillary stenosis
12. Peptic ulcer disease

Clinical Alert

Contraindications include:

1. Acute pancreatitis, pancreatic pseudocysts, and cholangitis
2. Obstructions or strictures within the esophagus or duodenum
3. Acute infections
4. Recent myocardial or severe pulmonary disease
5. Coagulopathy
6. Recent barium x-rays of the GI tract (barium obscures views during ERCP)

Interventions

Pretest Patient Preparation

1. Explain the purpose, procedure, benefits, and risks of the test. If done as an outpatient procedure, the patient should arrange for a ride home and should leave all valuables at home. Blood work, urinalysis, x-ray films, and scans should be reviewed and charted before the procedure. Record baseline vital signs and preprocedure signs and symptoms (eg, jaundice, persistent abdominal pain, and signs of pancreatic cancer).
2. Be aware that an informed consent form must be properly signed and witnessed.
3. Have the patient fast for 8–12 hours before ERCP.
4. Inform the patient to expect the following:
   a. The patient may be quite sleepy during the ERCP and may not recall much of the experience.
   b. The patient should swallow when requested to do so and should not attempt to talk (to prevent damage to the oral pharynx).
   c. Initially, the patient may experience a gagging or choking sensation that quickly subsides. Slow, deep breathing may help with this feeling. Sensations of abdominal pressure or bloating are normal.
   d. The patient will have to lie quietly while x-rays are being taken.
   e. Encourage the patient to urinate and defecate before the procedure.
5. Refer to conscious sedation precautions in Appendix C.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Do not permit food or liquids until the patient's gag reflex returns.
2. Monitor the blood pressure, pulse, respirations, oxygen saturation, and temperature according to institutional policy.
3. Observe the patient for signs of complications such as infection, urinary retention, cholangitis, or pancreatitis. Check for temperature elevation, which may be the first sign of inflammation. Monitor the white blood cell count and assess for signs of sepsis.
4. Infection may result from obstructed and infected biliary systems and/or contaminated endoscopes used during the procedure.
5. Monitor for respiratory and central nervous system depression from narcotics (naloxone may be used to reverse narcotic effects, and flumazenil is used for reversing diazepam-like drugs).
6. Explain that some abdominal discomfort may be experienced for several hours after the procedure.
7. Advise patient that drowsiness may last up to 24 hours. During this time, the patient should not perform any tasks that require mental alertness, and legal documents should not be signed.
8. Tell patient that a sore throat can be relieved by gargles, ice chips, fluids, or lozenges if permitted.
9. Notify physician of any of the following signs or symptoms:
   a. Prolonged, sharp abdominal pain; abnormal weakness; faintness
   b. Fever
   c. Nausea or vomiting
10. Interpret test outcomes and counsel appropriately.
11. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care. Provide written instructions to outpatients. Outpatients should be discharged to the care of a responsible adult.

Clinical Alert

Observe for possible complications:

1. Pancreatitis and sepsis (the most common complications of ERCP)
2. Hemorrhage
3. Perforation
4. Aspiration
5. Respiratory depression or arrest
6. Medication reaction

Enteroscopy

Enteroscopy is the endoscopic examination of the small bowel with fiberoptic endoscope. The endoscope is about 250–300 cm long, depending on the manufacturer. This long instrument is passed down the esophagus, through the stomach, through the distal duodenum, and then into the jejunum. Once in the jejunum, the endoscopist uses a series of movements to advance the endoscope as far as possible. A device known as an overtube may be applied to the endoscope to prevent it from looping in the stomach and inhibiting deep intubation of the small intestine. Fluoroscopy may also be useful in determining the position of the endoscope in the small bowel.

The main indication for enteroscopy is unexplained gastrointestinal bleeding. It may also be used to help diagnose patients with unexplained chronic diarrhea or suspicious x-ray findings. It is very useful in diagnosing a small bowel abnormality out of reach of a standard endoscope that might otherwise be done surgically.

Reference Values

Normal Small intestinal tract within normal limits

Procedure

1. This examination is usually performed in an outpatient setting of a hospital or ambulatory clinic. It also may be performed in the operating room or in a critical care setting.
2. Use a topical spray to anesthetize the patient's throat.
3. Start an intravenous line and use for administration of sedation alone or in combination with analgesics. These medications are given to achieve a state of conscious sedation (see Appendix C). Resuscitation equipment must be available.
4. Perform continuous monitoring of the patient's vital signs, ECG, and oxygen saturation (pulse oximetry).
5. Remove partial dental plates or dentures. Insert a mouthpiece to prevent the patient from biting the endoscope and to prevent injury to the patient's teeth, tongue, or other oral structures.
6. Be aware that depending on the endoscopist's preference, an overtube may be back-loaded onto the endoscope. The endoscope is well lubricated and gently inserted through the mouthpiece into the esophagus and advanced into the stomach and duodenum. To advance into the distal duodenum and jejunum, the endoscopist may use a series of pushing and pulling movements that serve to pleat the small bowel onto the endoscope, allowing deeper intubation. Fluoroscopy is useful to determine location in the small bowel.
7. Obtain biopsy specimens and brushing for cytology. Take photos to provide a permanent record of observations.
8. Inform the patient that he or she may initially have a strong gagging or choking sensation. During the procedure, the patient may belch frequently and have a sensation of abdominal pressure and bloating.
9. Immediately after the procedure, have the patient remain on his or her left side until fully awake.

Clinical Implications Abnormal results would indicate the following:

1. Vascular abnormalities, such as angiodysplasia or varices
2. Ulcerative lesions, such as in Crohn's disease
3. Diverticula, such as Meckel's diverticulum
4. Tumors

Interventions

Pretest Patient Preparation

1. Explain the purpose and procedure of the examination, the sensations that may be experienced, and the benefits and risks of the test. Refer to the conscious sedation and analgesia precautions in Appendix C.
2. Inform the patient that the procedure may be several hours long, depending on the ease of passing the endoscope, diagnosis, and treatment.
3. Inform the patient that he or she might be quite sleepy during the test and may not recall much of the experience.
4. Instruct the patient on fasting 10–12 hours before the procedure to avoid the risks for aspiration and possible cancellation of the procedure.
5. Confirm informed consent. A legal consent form must be signed and witnessed before the procedure.
6. Encourage the patient to urinate and defecate if possible before the examination.
7. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. No food or liquids are permitted until the patient's gag reflex returns. Sucking on ice chips or throat lozenges may be helpful to relieve a sore throat.
2. Monitor blood pressure, pulse, respirations, and oxygen saturation according to the hospital or clinic policy, until the patient is fully awake.
3. The patient should remain on his or her left side with side-rails raised until fully awake.
4. Encourage the patient to belch to expel air inserted during the procedure.
5. Interpret test outcomes and monitor appropriately.
6. Follow guidelines in Chapter 1 regarding safe, effective, and informed posttest care. Provide written discharge instructions to outpatients. Outpatients should be discharged to the care of a responsible adult.

Education Alert

New technology in the form of a video capsule may aid diagnosis of small bowel abnormalities. The video capsule is basically a wireless virtual endoscope that transmits video images to a recorder. The patient fasts for 12 hours, then swallows the video capsule (which is the size of a large pill) and wears a recorder around the waist. The patient can begin drinking and eating several hours later. The video capsule travels through the entire GI tract via normal peristalsis, recording images. The images from the recorder can then be downloaded and reviewed for diagnostic purposes.

Clinical Alert

Potential complications include:

1. Shearing or stripping of gastric mucosa (which may arise from use of the overtube)
2. Pancreatitis
3. Hemorrhage
4. Perforation

Colposcopy

Colposcopy permits examination of the vagina and cervix with the colposcope, an instrument with a magnifying lens. The colposcope is also used to examine male genital lesions suspected in sexually transmitted diseases, condylomas, or human papillomavirus. Indications for this procedure in women include abnormal Papanicolaou (Pap) smear results and/or other cervical lesions, leukoplakia, and other cancerous lesions. Biopsy specimens and cell scrapings are obtained under direct visualization. Colposcopy is also valuable for assessing women with a history of exposure to diethylstilbestrol. Advantages of colposcopy include the following:

1. Lesions can be localized and their extent determined.
2. Inflammatory processes can be differentiated from neoplasia.
3. Invasive or noninvasive disease processes can be differentiated.

Colposcopy cannot readily detect endocervical lesions. Cervicitis and other changes can produce abnormal findings. When combined with findings from Pap smears, colposcopy can be a means of enhancing diagnostic accuracy. Tables 12.1 and Tables 12.2 present correlation of findings and advantages and disadvantages of Pap smears and colposcopy. See Chapter 11 for Pap smear procedure.

<table>
<thead>
<tr>
<th>Table 12.1 Correlation of Colposcopic and Histologic (Tissue) Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Colposcopic Term</strong></td>
</tr>
<tr>
<td>Original squamous epithelium</td>
</tr>
<tr>
<td>Columnar epithelium</td>
</tr>
</tbody>
</table>
Transformation zone: Tongues of squamous metaplasia; gland openings; nabothian cysts

White epithelium: White, sharp-bordered lesion visible only after application of acetic acid; no vessels visible

Punctuation: Sharp-bordered lesion; red stippling; epithelium whiter after application of acetic acid

Mosaic: Sharp-bordered lesion, mosaic pattern; epithelium whiter after application of acetic acid

Hyperkeratosis: White patch; rough surface; already visible before application of acetic acid

Atypical vessel: Horizontal vessels running parallel to surface; constrictions and dilatations of vessels; atypical branching, winding course

**Table 12.2 Pros and Cons of Colposcopy and Cytology (Examination of Cells)**

<table>
<thead>
<tr>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>COLPOSCOPY</strong></td>
<td></td>
</tr>
<tr>
<td>Localizes lesion</td>
<td>Inadequate for detection of endocervical lesions</td>
</tr>
<tr>
<td>Diagnostic biopsy reveals cause of cancer</td>
<td>More intensive training is necessary</td>
</tr>
<tr>
<td>Evaluates extent of lesion</td>
<td>Cervicitis and regenerative changes may produce abnormal findings</td>
</tr>
<tr>
<td>Differentiates between inflammatory atypia and neoplasia</td>
<td></td>
</tr>
<tr>
<td>Differentiates between invasive and noninvasive cervical lesions</td>
<td></td>
</tr>
<tr>
<td>Enables follow-up</td>
<td></td>
</tr>
<tr>
<td>CYTOLOGY</td>
<td></td>
</tr>
<tr>
<td>Ideal for mass screening</td>
<td>Cannot localize lesion</td>
</tr>
<tr>
<td>Economical</td>
<td></td>
</tr>
<tr>
<td>Detection of HPV by DNA for cervical cancer</td>
<td>Inflammation, atrophic changes, or folic acid deficiency may produce suspicious changes</td>
</tr>
<tr>
<td>Specimen can be obtained by most health care personnel</td>
<td>Many steps between patient and cytopathologist allow misdiagnosis</td>
</tr>
<tr>
<td>Detects lesion in endocervical canal</td>
<td>Value of single smear is limited</td>
</tr>
<tr>
<td>Detects endocervical and endometrial carcinoma</td>
<td>False-negative rate is 5%–10%</td>
</tr>
<tr>
<td>High correlation with biopsy material (&gt;90%)</td>
<td></td>
</tr>
</tbody>
</table>

See Chapter 11 for more information on cytology and histology.

Whitish areas of epithelium (leukoplakia), mosaic staining patterns, irregular blood vasculature, hyperkeratosis, and other abnormal-appearing tissues can be seen using colposcopy. Leukoplakia vulva is a precancerous condition characterized by white to grayish infiltrated patches on the vulvar mucosa. The colposcope has a definite advantage for detecting atypical epithelium, designated in the literature as basal cell activity. Atypical epithelium cannot be called benign and yet does not fulfill all criteria for carcinoma in situ. Its early detection promotes cancer prophylaxis.

Patients receiving colposcopy may often be spared having to undergo surgical conization (the removal of a cone of tissue from the cervix).

Another gynecology procedure, a hysteroscopy, can be done to determine the cause of abnormal uterine bleeding, size and shape of the uterine cavity, location of a misplaced intrauterine device (IUD), and uterine abnormalities. A hysteroscopy is performed early in the menstrual cycle in a physician's office. A local anesthetic is usually administered into the cervix and paracervical area before insertion of the hysteroscope.

**Reference Values**

*Normal* Normal vagina, cervix, vulva, and genital areas Normal pink squamous epithelium and capillaries Normal color, tone, and surface contours

**Procedure**

1. Place the patient in the modified lithotomy position. Expose the vagina and cervix with a speculum after the internal and external genitalia have been carefully examined. Do not insert any part of the colposcope into the vagina.
2. Swab the cervix, vagina, or male genital areas with 3% acetic acid as needed during the procedure to improve visibility of epithelial tissues (it precipitates nuclear proteins within the cells). Remove the cervical mucus completely. Do not use cotton-wool swabs because fibers left on the cervix interfere with proper visualization.
3. Begin actual visualization with the colposcope with a field of white light and decreased magnification to focus on sites of white epithelium or irregular cervical contours. The light is then switched to a green filter for magnification of vascular changes.
   a. Diagram suspicious lesions and take photographs for the permanent health care record.
   b. The transformation zone and squamocolumnar junction (where the squamous epithelium meets the columnar epithelium of the cervix) are areas where many women exhibit atypical cells. It is imperative that these zones be visualized completely, especially in older women, because of changes associated with aging.
4. Obtain biopsy specimens of the lesions using a fine biopsy forceps. Some patients note discomfort at this time.
5. Place specimen in proper preservative, label accurately, and route to the appropriate department.
   a. Endocervical curettage must be performed before colposcope-directed biopsy so that epithelial fragments dislodged during colposcopy do not cause false-positive results in the endocervical curettage. The endocervical smear (curettage biopsy samples) should be placed on a slide in formalin.
   b. Sterile saline or sterile water should be used to cleanse and rinse acetic acid from the vaginal area to prevent burning or irritation. Bleeding can be stopped by applying toughened silver nitrate cautery sticks or ferric subsulfate (Monsel's solution).
6. Alert patient that a small amount of vaginal bleeding or cramping for a few hours is not abnormal.
7. Be aware that a paracervical block may be necessary in patients who are extremely anxious and afraid.
8. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

**Education Alert**
Before undergoing this procedure, the patient knows that the cervical (Pap) smear is abnormal. Women often believe that they do have cancer unless it is explained that they do not.

**Clinical Implications**

1. Abnormal lesions or unusual epithelial patterns include the following:
   a. Leukoplakia
   b. Abnormal vasculature
   c. Slight, moderate, or marked dysplasia
   d. Abnormal-appearing tissue is classified by punctuation (ie, sharp borders, red stippling, epithelium whiter with acetic acid); mosaic pattern (ie, sharp borders, mosaic pattern, epithelium whiter with acetic acid); or hyperkeratosis (ie, white epithelium, rough, visible without acetic acid)
2. Extent of abnormal epithelium (with acetic acid) and extent of nonstaining with iodine
3. Development of cervical changes and potential cervical carcinoma is a greater risk in these patients. An annual Pap smear is mandatory for those who have undergone colposcopy.
4. Acute inflammation with human papillomavirus or bacterial infections (eg, chlamydia), bacterial vaginosis and gonorrhea

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure. Record preprocedure signs and symptoms (eg, abnormal Pap, cervical or vaginal drainage or bleeding).
2. Obtain a urine specimen and a pertinent gynecologic history.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Clinical Alert**

1. Patients may experience a vasovagal response. Watch for bradycardia and hypotension and treat accordingly. Have the patient sit for a short while before standing.
2. Anti-inflammatory agents such as ibuprofen may relieve cramping.
3. Cervical scars from previous events may prevent satisfactory visualization.
4. Monitor for complications, including heavy bleeding, infection, or pelvic inflammatory disease.
5. Development of cervical changes and potential cervical carcinoma is a greater risk in these patients. An annual Pap smear is mandatory for those who have undergone colposcopy.
6. A paracervical block may be used with the procedure.

**Posttest Patient Aftercare**

1. Instruct the patient to abstain from sexual intercourse and to not insert anything into the vagina for 2 to 7 days (per physician's orders) after the procedure.
2. If specimens are taken, slight vaginal bleeding may occur. Excessive bleeding, pain, fever, or abnormal vaginal discharge should be reported immediately. Ibuprofen may relieve cramps.
3. Interpret test outcomes and counsel appropriately regarding follow-up treatment such as cone biopsy and loop electrosurgical excision procedure (LEEP). If radiation treatment is prescribed, cervical tumor tissue may be tested for the presence of glutathione as a possible indicator of radiation resistance.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care. Provide written discharge instructions.

**Loop Electrosurgical Excision Procedure (LEEP), Cone Biopsy, Cervical Conization**

These procedures are done as a follow-up for an abnormal Pap smear and colposcopy findings, to enhance accuracy of colposcopy, and to investigate squamous intraepithelial lesions (SILs). They are done to exclude invasive cancer, determine extent of noninvasive lesions, and treat (LEEP and cone biopsy) and remove abnormal cervical dysplasia, based on lesion size, distribution, and grade, when there is lack of correlation between Pap smear, previous biopsy, and
colposcopy.

Reference Values

Normal Normal cervix cells, which flatten as they grow

Procedure

1. Place the patient’s feet in stirrups and insert a speculum, as with a Pap test and colposcopy.
2. Apply a local anesthetic to the cervix and a mild vinegar (acetic acid) or iodine, depending on the procedure type. For LEEP procedures, insert a fine wire loop with a special high-frequency current to remove a small piece of cervical tissue.
3. Apply a paste to the cervix to reduce bleeding. This may cause a dark vaginal discharge. A laser or a cone biopsy may also be one of the procedures.

Clinical Alert

Complications may include heavy bleeding, severe cramping, infection, and accidental cutting or burning of normal tissue. Cervical stenosis may be an untoward effect of this procedure.

Clinical Implications Abnormal findings include dysplasia and invasive cancer into deeper parts of the cervix.

Interventions

Pretest Patient Preparation

1. Explain purpose and procedure and equipment used for procedures. Tell patient that the most common symptom of cervical cancer is abnormal vaginal discharge or bleeding.
2. Provide support and take measures to relieve fear and anxiety about possible diagnosis of cervical cancer.
3. Refer to Chapter 1 guidelines regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Instruct patient to call physician if heavy or bright-red bleeding or clots, chills, aching, severe abdominal pain (not relieved by pain medication), foul-smelling discharge, or unusual swelling occurs. Water discharge and white, dark, and light spotting may last approximately 4 weeks. The heaviest discharge occurs about 1 week after treatment. Do not use tampons.
2. No douching or bubble baths. Delay sexual intercourse for approximately 4 weeks. Check with physician.
3. No douching or bubble baths. Delay sexual intercourse for approximately 4 weeks. Check with physician.
4. Be sure to stress the importance of returning for follow-up appointment to evaluate satisfactory healing.
5. Follow Chapter 1 guidelines regarding safe, effective, informed posttest care.

Flexible Proctoscopy; Sigmoidoscopy; Proctosigmoidoscopy

These tests involve the examination of the rectum, anal canal, and sigmoid colon, up to 65 cm, with a proctosigmoidoscope. Rigid scopes are not as commonly used since the advent of flexible fiberoptic instruments, which are more comfortable for patients. Their main use is for the investigation of rectal bleeding, evaluation of colonic symptoms, detection and diagnosis of cancers and other abnormalities such as diverticula in this area of the gastrointestinal tract. These examinations should be routine (every 3–5 years) for cancer screening of individuals >50 years of age. These tests can also evaluate hemorrhoids, polyps, blood or mucus in the stool, unexplained anemia, and other bowel conditions. Sigmoidoscopy is used along with air-contrast barium studies.

Reference Values

Normal Normal anal, rectal, and sigmoid colon mucosa

Procedure

1. Have the patient assume the knee-to-chest position for rigid proctoscopy (inserted 25 cm). When the flexible proctoscope is used, the patient must be in the left lateral position. Carefully insert the proctoscope (inserted 35–60 cm) or sigmoidoscope into the rectum.
2. Remember that the examination can be done with the patient in bed or positioned on a special tilt-table.
3. Inform the patient that he or she may feel a very strong urge to defecate or pass gas. The patient may also experience a feeling of bloating or cramping, which is normal.
4. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications Examination may reveal the following: edematous, red, or denuded mucosa; granularity; friability; ulcers; polyps; cysts; thickened areas; changes in vascular pattern; pseudomembranes; spontaneous bleeding; or normal mucosa. These findings may help to confirm or to rule out the following conditions:

1. Inflammatory bowel disease
   a. Chronic ulcerative colitis
   b. Crohn's disease
   c. Proctitis (acute and chronic)
   d. Pseudomembranous colitis
   e. Antibiotic-associated colitis
2. Polyps
   a. Adenomatous
   b. Familial
   c. Diminutive
3. Cancer and tumors
   a. Adenocarcinoma
b. Carcinoids

c. Other tumors such as lipomas

4. Anal and perianal conditions
   a. Hemorrhoids
   b. Abscesses and fistulas
   c. Strictures and stenoses
   d. Rectal prolapse
   e. Fissures
   f. Contractures

Interventions

Pretest Patient Preparation

1. Explain test purpose, procedure, and benefits (when used with annual fecal occult blood test) and risks (a safe procedure). Record pertinent preprocedure signs and symptoms (eg, rectal bleeding). Obtain a signed, witnessed informed consent, if required.

2. Be aware that there is no need for the patient to fast. However, a restricted diet such as clear liquids the evening before the test may be prescribed.

3. Remind patient that laxatives and enemas may be taken the night before the examination. Enemas or a rectal laxative suppository may be administered the morning of the procedure. For patients of all ages, one or two phosphate (Fleet) enemas are frequently ordered to be performed about 1 to 2 hours before the examination. This is considered ample preparation by many endoscopy departments.

4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Clinical Alert

1. Patients with acute symptoms, particularly those with suspected ulcerative or granulomatous colitis, should be examined without any preparation (ie, without enemas, laxatives, or suppositories).

2. Perforation of the intestinal wall can be an infrequent complication of these tests.

3. Notify the patient's physician before administering laxatives or enemas to a pregnant woman.

4. Notify physician immediately of any instance of decreased blood pressure, diaphoresis, or bradycardia.

Posttest Patient Aftercare

1. Interpret test outcomes. Monitor and counsel appropriately about possible further testing (colonoscopy).

2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care. Provide written discharge instructions.

Colonoscopy

Colonoscopy visualizes, examines, and photographs the large intestine with a flexible fiberoptic or video-colonoscope inserted through the anus and advanced to the ileocecal valve. Air introduced through an accessory channel of the colonoscope distends the intestinal walls to enhance visualization. Colonoscopy can differentiate inflammatory disease from neoplastic disease and can evaluate polypoid lesions that are beyond the reach of the sigmoidoscope. Polyps, foreign bodies, and biopsy specimens can be removed via the colonoscope. Photographs of the large intestine lumen can also be taken. Before colonoscopy was available, major abdominal surgery was the only way to remove polyps or suspicious tissue to determine malignancy or nonmalignancy. Periodic colonoscopy is a valuable adjunct to the follow-up of persons with previous polyps, colon cancer, family history of colon cancer, or high risk factors. It is also helpful in locating the source of lower gastrointestinal bleeding. It provides a safe way to perform presurgical screening and postsurgical surveillance of suture lines and anastomoses. Colonoscopy, along with stool occult blood testing, is recommended as the primary diagnostic tool for first-degree relatives in colon cancer families.

Reference Values

Normal Normal large intestine mucosa

Procedure

1. Be aware that a clear liquid diet is usually ordered for 48 to 72 hours before examination. Have the patient fast for 8 hours before the procedure. Laxatives may be ordered to be taken for 1 to 3 days before the test; enemas may be ordered to be given the night before the test.

2. For an oral saline iso-osmotic and isotonic laxative, have the patient drink 3 to 6 liters of the prescribed solution over a 2- to 3.5-hour period. The typical volume taken is 1 gallon (4 liters), and this volume of fluid can be administered by nasogastric tube if necessary. Expect initial results in 30 minutes to 1 hour. Ingestion of the washout solution continues until feces expelled are nothing but clear liquid. Notify the physician before administering >6 liters of this solution. Be aware that patients with congestive heart failure or renal failure may be at risk for fluid volume overload if this preparation is used. See Patient Preparation for other preparation measures.

3. Start an intravenous line and use for administration of sedatives and narcotics. These medications are given to achieve a state of conscious sedation (see Appendix C). Ensure that the patient is responsive enough to inform the doctor of any subjective reactions during the examination. Ensure that resuscitation equipment is available.

4. Perform continuous monitoring of the patient's vital signs, ECG, and oxygen saturation (pulse oximetry).

5. Be aware that, on occasion, intravenous anticholinergic agents and glucagon may be used to relax bowel spasms.

6. Have the patient assume the left-sided or Sims' position, and drape properly. Insert a well-lubricated colonoscope about 12 cm into the bowel. Ask the patient to take deep breaths through the mouth during this time. Introduce air into the bowel through the special port on the colonoscope to aid viewing. As the colonoscope advances, the patient may need to be repositioned several times to aid in proper visualization of the colon. Sensations of pressure, mild pain, or cramping are not unusual.
7. Remember that the best views are obtained during withdrawal of the colonoscope. Therefore, a more detailed examination is usually performed during withdrawal than during advancement.
8. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Alert

A virtual colonoscopy preceded by traditional bowel cleansing is a noninvasive procedure (no sedation or analgesia is required for virtual colonoscopy) using the latest CAT scan equipment to rule out abnormalities. The same abnormal conditions can be identified. However, the traditional invasive procedure must be repeated if there are abnormal findings and it is necessary to remove polyps and take biopsy specimens.

Clinical Implications Abnormal findings may reveal the following conditions:
1. Polyps
2. Tumors (benign or malignant)
3. Areas of ulceration
4. Inflammation
5. Colitis, diverticula
6. Bleeding sites
7. Strictures
8. Discovery and removal of foreign bodies

Clinical Alert

To be effective, a purgative must produce fluid diarrhea. This shows that unaltered small intestinal contents are emerging and colonic residue has been cleared. Enemas must be repeated until solid matter is no longer expelled (clear liquid returns). Soapsuds enemas are contraindicated because they cause increased mucus secretion as a result of irritant stimulation.

Clinical Considerations
1. Keep colon electrolyte lavage preparations refrigerated; however, the patient may drink the solution at room temperature. Use within 48 hours of preparation, and discard unused portions.
2. Before testing, the complete blood count, prothrombin time, platelet count, and thromboplastin time results should be reviewed and charted.
3. Preparation for patients with a colostomy or who are paralyzed is the same whether or not the patient is taking aspirin or any blood thinners.
4. Persons with known heart disease may receive prophylactic antibiotics before testing.
5. Patients should not mix or drink anything with the washout preparation. Do not add ice or glucose to the solution.
6. Diabetic persons are usually advised not to take insulin before the procedure but to bring insulin with them to the clinic.

Interventions

Pretest Patient Preparation

1. Explain the purpose, procedure, benefits and risks of the test. Record preprocedure signs and symptoms (eg, GI bleeding). If done as an outpatient procedure, the patient should arrange for a ride home and should leave valuables at home. Blood work, urinalysis, x-ray films, and scans should be reviewed and charted before the procedure. Record baseline vital signs.
2. When ordered, have the patient take one 12-ounce glass of liquid preparation every 10 minutes before the examination. (Each gallon holds 10.7 12-ounce [360 mL] glasses). The entire gallon should be taken in 2 hours, if possible. Timing is important. Slower drinking does not clean the colon properly. Some patients will receive another type of preparation when ordered (eg, Propulcid capsules and liquid Fleet laxatives and enemas).
3. Remember that some patients will be on a clear-liquid diet for 72 hours before the test, then fasting, except for medications, after a clear-liquid supper the evening before the test. No solid food, milk, or milk products are permitted. Strained fruit juices without pulp (eg, apple, white grape), lemonade, Hi-C drink, water, clear liquid, Gatorade, Kool-Aid, Jell-O, Popsicles, and hard candy are permitted, but no red or purple fluids are allowed.
4. Administer purgatives and cleansing enemas as ordered. Preparation is complete when fecal discharge is clear. If returns are not clear after 4 liters of solution have been ingested, continue until returns are clear, up to 6 liters total (see previous note under Procedure).
5. Be aware that a legal consent form must be signed and properly witnessed (see Chap. 1) after patient has received proper instruction about the test.
6. Discontinue iron preparations 3 or 4 days before examination because iron residues produce an inky, black, sticky stool that interferes with visualization, and the stool can be viscous and difficult to clear. Aspirin and aspirin-containing products should also be discontinued 1 week before the examination because they may cause bleeding problems or localized hemorrhages.
7. Inform the patient to expect the following:
   a. The patient may feel quite sleepy during the test and may not recall much of the procedure.
   b. The patient may experience abdominal pressure, mild pain, or cramping.
   c. The patient may pass gas (expel flatus) or have the urge to defecate, which is normal.
   d. The patient may be asked to assume various positions to aid with passing the colonoscope.
8. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.
1. The patient may have liquids or a light meal when fully awake.
2. Stools should be observed for visible blood. The patient should be instructed to report abdominal pain or other unusual symptoms because perforation and hemorrhage are possible complications.
3. Monitor the blood pressure, pulse, respirations, and oxygen saturation, according to institutional policy, until the patient is fully awake.
4. The most frequent adverse reactions to oral purgatives include nausea, vomiting, bloating, rectal irritation, chills, and feelings of weakness.
5. The patient may expel large amounts of flatus after the procedure.
7. Follow guidelines in Chapter 1 for safe, effective, informed posttest care. Provide written discharge instructions to outpatients. Outpatients should be discharged to the care of a responsible adult.

### Clinical Alert

1. Solid food should never be taken within 2 hours before the oral cleansing regimen is begun.
2. Orally administered colon lavage is contraindicated in the following conditions:
   a. Actual or suspected ulcers
   b. Gastric outlet obstruction
   c. Weight < 20 kg
   d. Toxic colitis
   e. Megacolon
3. Relative contraindications for colonoscopy include the following conditions:
   a. Perforating disease of the colon
   b. Peritonitis
   c. Radiation enteritis
   d. Recent abdominal or bowel surgery
   e. Acute conditions of the anus and rectum
   f. Serious cardiac or respiratory problems (eg, recent myocardial infarction)
   g. Situations in which the bowel cannot be adequately prepared for the procedure (ie, fulminant granulomatous or irradiation colitis)
4. Observe for the following possible complications:
   a. Perforations of the bowel
   b. Hypotensive episodes
   c. Cardiac or respiratory arrest, which can be provoked by the combination of oversedation and intense vagal stimulus from instrumentation
   d. Hemorrhage, especially if polypectomy has been performed
   e. Death (extremely rare)
5. If colon preparations are administered by lavage to an unconscious patient or to a patient with impaired gag reflexes, observe for aspiration or regurgitation, especially if a nasogastic tube is in place. Keep the head of the bed elevated. If this is not possible, position the patient on his or her side. Have continuous suction equipment and supplies readily available.
6. No barium studies should be done during the preparation phase for colonoscopy.
7. Signs of bowel perforation include malaise, rectal bleeding, abdominal pain, distention, and fever.
8. Bloating, nausea, and occasional vomiting after oral laxatives is common. Advise patient to adhere to instructions if at all possible.

### Peritoneoscopy; Laparoscopy; Pelviscopy; Fertiloscopy

These examinations of the intraabdominal and pelvic cavities are performed using a laparoscope or pelviscope inserted through a slit in the anterior abdominal wall. The pelvic organs, as well as abdominal organs such as the greater curvature of the stomach or the liver, can be viewed. The use of laparoscopic intracorporeal ultrasound probe (LICU) in the evaluation of GI malignancies is an important new technology. The different types of examinations include peritoneoscopy, laparoscopy (intraabdominal), pelviscopy (gynecologic), and fertiloscopy (gynecologic). These procedures are frequently performed under general anesthesia in a surgical setting; however, many are also done with local anesthesia.

Peritoneoscopy is most commonly done to evaluate liver disease and to obtain biopsy specimens when the liver is too small, when previous liver biopsy proves inadequate, when contraindications to percutaneous liver biopsy exist (eg, ascites), when there is unexplained portal hypertension or liver function abnormalities, and when the liver cannot be properly palpated for a conventional liver biopsy. It does away with the need for a blind liver biopsy. Other indications for peritoneoscopy include unexplained ascites, staging of lymphomas, or staging and follow-up of ovarian cancer or abdominal masses. Sometimes patients with advanced chest, gastric, pancreatic, endometrial, or rectal tumors are evaluated by peritoneoscopy before surgical intervention is attempted.

Indications for laparoscopy include diagnosis and staging of cancer, evaluating cause of ascites, and examination of abdomen with ultrasound probes. Biopsies of abdominal and lymph node masses and hepatic lesions can also be done using a core needle biopsy, wedge biopsy using electrocautery, and cup forceps biopsy.

Gynecologic laparoscopy and pelviscopy are used to diagnose cysts, adhesions, fibroids, malignancies, inflammatory processes, or infections in persons with pelvic and abdominal pain. Evaluation of the fallopian tubes can be done for infertile patients. These procedures also provide a means to release adhesions, to obtain biopsy specimens, to do select operative procedures such as tubal ligations, or to perform laser treatments for endometriosis. Gynecologic laparoscopy
or pelviscopy is commonly performed under general anesthesia as a same-day surgical procedure.

These techniques can frequently replace laparotomy. They are less stressful to the patient; require only small incisions; can be done in shorter periods of time; can be done using local, spinal, or general anesthetics; reduce potential for formation of adhesions; and hasten healing and recovery time.

Pelviscopy differs from laparoscopy in two major respects—endocoagulation as a method for controlling bleeding and endoligation as a technique that permits suturing using extracorporeal (outside the body) or intracorporeal (inside the body) ligating and suturing methods by means of special instruments.

The pelviscope is angled at 30 degrees for better visualization. A videocamera attachment offers the physician a choice of viewing the process on a videoscreen instead of through the scope. Printouts and videotapes of the pelviscopy can be produced. Thus, pelviscopy is both a diagnostic and an operative modality.

Fertiloscopy, using a new instrument called a fertiloscope, is used to examine the entire female reproductive system. It is used to diagnose infertility and replaces laparoscopic tests done to diagnose infertility. Local anesthetics are used. Future applications include testing to rule out ovarian cancer, biopsy of the ovary, and drainage of cysts.

Reference Values

Normal
Gynecologic examination: normal size, shape, and appearance of uterus, fallopian tubes, and ovaries.

Intraabdominal examination: normal liver, gallbladder, spleen, and greater curvature of the stomach

Procedure

1. Have the patient lie supine during all procedures except gynecologic laparoscopy, in which case the patient is placed in a lithotomy position.
2. Cleanse the skin and, if the procedure is to be performed under local anesthesia, inject a local anesthetic into areas where the scope will be introduced. Otherwise, prep the patient as for an abdominal procedure under general anesthesia. Maintain a sterile field.
3. Place an intravenous line so that medications may be given intravenously as needed.
4. Place an indwelling catheter into the bladder to reduce the risk for bladder perforation.
5. Make a small incision near the umbilicus through which a trocar is introduced, followed by passage of the pelviscope or laparoscope. Sometimes, more than one puncture site will be made so that accessory instruments can be used during the procedure. Carbon dioxide introduced into the peritoneal cavity causes the omentum to rise away from the organs and allows for better visualization. A few stitches or Steri-Strips are usually needed to close the incisions. Apply adhesive bandages as dressings.
6. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications

Abnormal findings can reveal the following conditions:

1. Endometriosis
2. Ovarian cysts
3. Pelvic inflammatory disease
4. Metastasis stage of cancer
5. Uterine fibroids
6. Abscesses
7. Tumors (benign and malignant)
8. Enlarged fallopian tubes (hydrosalpinx)
9. Ectopic pregnancy
10. Infection
11. Adhesions or scar tissue
12. Ascites
13. Cirrhosis
14. Liver nodules (often an indication of cancer)
15. Engorged peritoneal vasculature (correlates with portal hypertension)

Clinical Alert

1. These procedures may be contraindicated in persons known to have the following conditions:
   a. Advanced abdominal wall cancer
   b. Severe respiratory or cardiovascular disease
   c. Intestinal obstruction, dilated bowel loops
   d. Palpable abdominal mass
   e. Large abdominal hernia
   f. Chronic tuberculosis
   g. History of peritonitis
   h. Noncorrectable coagulation disorders
2. Possible complications include the following:
   a. Bleeding from the puncture injury or from liver biopsy
   b. Misplacement of gas
   c. Thermal burns
3. The endoscopy should be aborted in favor of a laparotomy in the event of uncontrolled bleeding or suspected malignancy.

Interventions

Pretest Patient Preparation
1. Complete laboratory tests and other appropriate diagnostic modalities before these endoscopies.
2. Remember that bowel preparation may include an enema or suppository.
3. Ensure that a legal permit is properly signed and witnessed (see Chap. 1).
4. Maintain sensitivity to cultural, sexual, and modesty issues as an important part of psychological support.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Check blood pressure frequently according to institutional policies.
2. Observe for infection, hemorrhage, and bowel or bladder perforation.
3. Advise the patient that shoulder and abdominal discomfort may be present for 1 to 2 days because of residual carbon dioxide gas in the abdominal cavity. This can be controlled with mild oral analgesics. Sitting or resting in a semi-Fowler's position can also alleviate discomfort.
4. If the patient has had a general or spinal anesthetic, follow the usual cautions and protocols for the care of any person having undergone those types of anesthesia.
5. Interpret test outcomes and counsel appropriately.
6. Follow guidelines in Chapter 1 for safe, effective, informed posttest care. Provide written discharge instructions.

Cystoscopy (Cystourethroscopy)

Cystoscopy and cystourethroscopy are used to diagnose and treat disorders of the lower urinary tract. They provide views of the interior bladder, urethra, male prostatic urethra, and ureteral orifices by means of tubular, lighted, telescopic lens instruments called cystoscopes or cystourethroscopes. These scopes come in many sizes and variations as well as in flexible fiberoptic instruments. Urethroscopy is an important part of this examination because it allows visualization of the male prostate gland.

Cystoscopy is the most common of all urologic diagnostic procedures. It may be indicated in the following conditions:

1. Unexplained hematuria (gross or microscopic)
2. Recurrent or chronic urinary tract infection
3. Infection resistant to medical treatment
4. Unexplained urinary symptoms such as dysuria, frequency, urgency, hesitancy, intermittency, straining, incontinence, enuresis, or retention
5. Bladder tumors (benign and malignant)
6. Pediatric considerations include the above and the following:
   a. Posterior urethral valves, ureteroceles in females, and other congenital anomalies
   b. Complete workup of children with daytime incontinence usually done in conjunction with urodynamic studies
   c. Removal of foreign objects and stents placed in previous surgeries

Because intravenous pyelogram (IVP) does not allow proper visualization of the area from the neck of the bladder to the end of the urethra, cystoscopy makes it possible to diagnose and to treat abnormalities in this area.

Cystoscopy may be used to perform meatotomy and to crush and retrieve small stones and other foreign bodies from the urethra, ureter, and bladder. Biopsy specimens can be obtained. Bladder tumors can be fulgurated, and strictures can be dilated through the cystoscope. In conjunction with cystoscopy, ureteroscopy can be done to determine the cause of hematuria, to detect tumors and stones, and to manipulate stones.

Reference Values

Normal Normal structure and function of the interior bladder, urethra, ureteral orifices, and male prostatic urethra

Procedure

1. The examination can be performed in a special operating room designed for that purpose in a clinic or in the urologist's office. The patient's age, state of health, and extent of surgical procedure necessary determine the setting. Pediatric cystoscopy is done in the operating room under general anesthesia.
2. The external genitalia are prepped with an antiseptic solution such as povidone-iodine after the patient is properly grounded, padded, and draped.
3. Local anesthetic jelly is instilled into the urethra. For males, the anesthetic is retained in the urethra by a clamp applied near the end of the penis. For best results, the local anesthetic should be administered 5 to 10 minutes before passage of the cystoscope.
4. The scope is connected to an irrigation system, and fluid is infused into the bladder throughout the procedure. Solutions used are nonconductive and retain clarity during the procedure (eg, glycine, sterile water). The solution also distends the bladder to allow better visualization. The infusion is stopped and the bladder drained when it becomes filled with 300 to 500 mL of fluid.
5. Should blood or other matter be present in the bladder, the fiberoptic cystoscope will not provide as clear a view as a rigid cystoscope because it is more difficult to flush.
6. Institutional policies dictate general perioperative care and procedures. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.
During transurethral resection procedures, venous sinuses may be opened, and irrigation fluid may enter the circulatory system, causing water intoxication. Therefore, isotonic solutions such as sorbitol, mannitol, or glycine must be used.

Abnormal conditions revealed by cystoscopy include the following:

1. Prostatic hyperplasia or hypertrophy
2. Cancer of the bladder
3. Bladder stones
4. Urethral strictures or abnormalities
5. Prostatitis
6. Ureteral reflux (shown on cystogram)
7. Vesicle neck stenosis
8. Urinary fistulas
9. Ureteroceles
10. Diverticula
11. Abnormally small or large bladder capacity
12. Polyps

Interventions

Pretest Patient Preparation

1. Explain the purpose and procedure of the test. Special sensitivity to concern for cultural, social, sexual, and modesty issues are an important part of psychological support. Emphasize that there is little pain or discomfort from cystoscopy; however, a strong desire to void may be experienced.
2. Facilitate bowel preparation and other laboratory and diagnostic tests if extensive procedures are planned.
3. Remember that if cystoscopy is performed in the hospital, a properly signed and witnessed surgical permit must be obtained (see Chap. 1).
4. Allow the patient to take a full liquid breakfast at times. Liquids may be encouraged until the time of the examination to promote urine formation if the procedure is a simple cystoscopy done under local anesthesia. Fasting guidelines are followed when spinal or general anesthesia is planned.
5. Start an intravenous line for the administration of drugs to achieve a state of conscious sedation. Medications such as diazepam (Valium) or midazolam (Versed) are used to relax the patient. Amnesia may be a side effect. Younger men may experience more pain and discomfort than older men. Women usually require less sedation because the female urethra is shorter. The patient should be instructed to relax the abdominal muscles to lessen discomfort. See Appendix C regarding sedation and analgesia precautions.
6. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. After cystoscopy, voiding patterns and bladder emptying should be monitored. Check vital signs as necessary.
2. The intake of fluids should be encouraged.
3. Clots may form and may cause difficulty in voiding.
4. Report unusual bleeding or difficult urination to the physician promptly.
5. Urinary frequency, dysuria, pink to light-red urine, urethral burning, and posttest bladder spasms are common after cystoscopy.
6. Antibiotics may be prescribed before and after cystoscopy to prevent infection. Rectal opium suppositories may also be administered.
7. The potential for gram-negative shock is always present with urologic procedures because the urethra is such a vascular organ that any break in the tissues can allow bacteria to enter the bloodstream directly. Onset of symptoms can be rapid and may actually begin during the procedure if it is fairly lengthy. Observe for and promptly report chills, fever, increasing tachycardia, hypotension, and back pain to the physician. Blood cultures are usually ordered, followed by an aggressive regimen of antibiotic therapy.
8. Ureteral catheters may be left in place to facilitate urinary drainage, especially if there is concern about edema.
9. Routine catheter care is necessary for retention of urethral catheters. Follow institutional protocols. The patient may need instructions if discharged with catheter in place.
10. Interpret test outcomes and counsel appropriately.
11. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care. Provide written discharge instructions.

If urethral dilatation has been part of the procedure, the patient is advised to rest and to increase fluid intake.

Monitor patient's voiding patterns and bladder emptying (or instruct to self-monitor).

Evaluate and instruct the patient to watch for edema. Edema may cause urinary retention, hesitancy, weak urinary stream, or urinary dribbling any time within several days after the procedure. Warm sitz baths and mild analgesics may be helpful; however, an indwelling catheter may sometimes be necessary for relief.

URODYNAMIC STUDIES

Cystometrogram (CMG); Urethral Pressure Profile (UPP); Rectal Electromyogram (EMG); Cystourethrogram
These tests evaluate bladder, urethral, and sphincter function, identify abnormal voiding patterns, check status of neuroanatomic connectives between brain, spinal cord, and bladder, and consist of two main components: the cystometrogram (CMG) and the sphincter electromyogram (EMG). The combined measurement of the CMG and the EMG provides information about how the bladder adapts to being filled as well as how it reacts to the filling itself. These studies are indicated in an incontinent person and when there is evidence of neurologic disease (neurogenic bladder), spinal cord injury, dysuria, enuresis, infection, or specific neuropathies such as those found in multiple sclerosis, diabetes, and tabes dorsalis.

**Reference Values**

**Normal** Normal bladder sensations of fullness, heat, and cold  **Adult:** Normal bladder capacity of 400 to 500 mL, residual urine less than 30 mL, desire to void is at 175 to 250 mL; sensation of fullness felt at 350 to 450 mL; stream is strong and uninterrupted. Normal voiding pressures and muscle coordination Normal rectal EMG readings; urethral pressure profile readings normal  **Pediatric:** Bladder capacity varies with age. Compliant bladder: stretches to capacity without pressure increase. Bladder stability: no involuntary contractions

**Procedures**

1. **Cystometrogram (CMG)**
   a. Have the patient void and record urine flow rate, voiding pressure, and residual amount of urine voided.
   b. Insert a nonlatex double-lumen catheter into the bladder. Place adhesive patch electrodes parallel on each side of the anus. Measure residual urine. Connect the catheter to the cystometer. (A cystometer evaluates the neuromuscular mechanism of the bladder by measuring bladder capacity and pressure.) The bladder is gradually filled with sterile saline or sterile water or carbon dioxide gas in predetermined increments, and pressure readings are taken at these increments. Water or saline offers a more physiologic result and is less irritating.
   c. Make observations during the CMG about the patient's perception of heat and cold, bladder fullness, urge to void, and ability to inhibit voiding when bladder contractions occur.
   d. Remove the catheter and patch electrodes when the bladder is completely emptied of fluid.
   e. Inject cholinergic and/or anticholinergic drugs (eg, methantheline bromide [Banthine, atropine] or bethanechol chloride [Urecholine]) to determine their effects on bladder function (after CMG procedure).
   f. Perform the cystometric study as a control, followed by repeat study 20 to 30 minutes after injection of the drugs.
   g. Be aware that a change in posture from supine to standing or walking may be required during the examination.
   h. Remember that sleep studies may be performed in conjunction with an electroencephalogram (EEG) to evaluate persons having nocturnal incontinence (see Chap. 16 for EEG study).

2. **Rectal electromyogram (EMG)—monitors the pelvic floor muscles responsible for holding urine in the bladder.**
   a. Apply electrodes next to the anus, and attach a ground to the thigh, or introduce a needle electrode into the periurethral striated muscle. These electrodes record electromyographic activity during voiding and produce a simultaneous recording of urine flow rate. (See Chap. 16 for EMG study.)
   b. Pediatric rectal EMG: Patch electrodes record the coordination of the external sphincter and the pelvic floor muscle response to filling and the ability to inhibit bladder contractions. If the child voids on the table, the sphincter relaxes during voiding (which is normal).

3. **Urethral pressure profile (UPP)**
   a. Use a specially designed catheter, coupled to a transducer, to record pressures along the urethra as it is slowly withdrawn.
   b. Pediatric UPP: This profile assesses the functional urethral length as well as general competency of the urethra and sphincter. The same double-lumen catheter is used, which has premarked lines on it for both the CMG and the UPP. Slowly withdraw the catheter, and note the pressures at the premarked spots.

4. **Cystourethrogram—evaluates bladder wall and urethral abnormalities and tumors. It can be used to assess reflux and stress incontinence in women and to identify posttraumatic urine extravasation.**
   a. Instill an x-ray contrast medium into the bladder through a catheter until the bladder fills. Clamp the catheter and take x-rays with the patient assuming several different positions.
   b. Remove the catheter and take more x-rays as the patient voids and the contrast material passes through the urethra (voiding cystourethrogram).
   c. Pediatric cystourethrogram: Rarely are voiding cystourethrogram (VCUGs) done at the same time as EMGs. VCUGs are done in children to assess vesicle urethral reflux, to identify structural abnormalities, and to evaluate for voiding dysfunction; and they are usually done as part of the workup before considering EMG.

5. See Chapter 1 guidelines for **safe, effective, informed intratest care** for all procedures.

**Clinical Alert**

In children, the bladder is filled at 10% of what the bladder is expected to hold at a specific age (ex: age capacity [in ounces] plus 2 ounces).

**Clinical Implications** Abnormal results reveal motor and sensory defects, altered pressures and/or bladder capacity, and inappropriate or absent contractions of the pelvic floor muscles and internal sphincter during voiding.

1. **Bladder noncompliance:** During filling, the bladder is stiff, does not stretch as expected, and can possibly compromise kidney function over time. A large-capacity low-pressure bladder (high compliance) may indicate chronic overdistention from infrequent voiding habits or disturbed muscle coordination.

2. **Bladder instability (hyperreflexia):** During filling, the bladder contracts involuntarily; this occurs when the pressures go up and down in a wavelike pattern during filling due to overactivity of involuntary contractions. The unstable
bladder may be asymptomatic; many times no contractions are felt, but commonly patients have frequency, urgency, and incontinence.

3. The most common cause of incontinence is a vesicle-sphincter dyssynergia (disturbance of muscular coordination). This dyssynergia is thought to be responsible for incomplete emptying of the bladder, inappropriate voiding, perineal dampness, and predisposition to urinary tract infections.

4. Detrusor hyperreflexia: The patient cannot suppress voiding on command owing to upper or lower motor neuron lesions, as in cerebrovascular aneurysm, Parkinson's disease, multiple sclerosis, cervical spondylosis, and spinal cord injury above the conus medullaris.

5. Detrusor areflexia occurs when the detrusor reflex cannot be evoked because the peripheral innervation of the detrusor muscle has been interrupted, resulting in difficulty in initiating voiding without a residual volume being present in the bladder. The cause may be associated with trauma, spinal arachnoiditis, spinal cord birth defects, diabetic neuropathy, or anticholinergic effects of phenothiazides. In postmenopausal women, the urethral pressure profile may be altered because the mucosal sphincter is deprived of estrogen.

6. Urethrovesical hyperreflexia is caused by benign prostatic hypertrophy and stress urge incontinence.

Interfering Factors Disorientation or inability of the patient to cooperate affects the test results.

Interventions

Pretest Patient Preparation

1. Explain the purpose and procedure of the bladder function test, often done before and after certain types of spinal surgery. Be sensitive to the patient's potential anxiety and embarrassment. Record signs and symptoms of incontinence and voiding problems.

2. Ensure that the patient is relaxed and cooperative for accurate results. For children, a favorite toy or book may provide security. Sedation is not given because patient participation is necessary to verify sensations and perceptions. However, the patient must avoid movement during the examination unless instructed otherwise.

3. Allow the test and filling of the bladder to continue until the patient either leaks or voids around the catheter.

4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Encourage the patient to increase oral fluid intake to dilute the urine and to minimize bladder sensitivity.

2. Explain that some minor discomfort or burning may be noted, especially if carbon dioxide is used, but it will lessen and disappear with time.

3. Interpret test outcomes and counsel appropriately (bladder capacity varies with age). Explain possible treatments (medication).

4. Follow Chapter 1 guidelines for safe, effective, informed posttest care. Provide written discharge instructions.

Clinical Alert

1. Certain patients with cervical cord lesions may exhibit an autonomic reflex that produces an elevated blood pressure, severe headache, lower pulse rate, flushing, and diaphoresis. Propantheline bromide (Pro-Banthine) alleviates these symptoms.

2. Careful use of sterile technique reduces the incidence of urinary tract infections. Preprocedural urinary tract infections can lead to sepsis as a result of bacterial spread into the bloodstream.

Arthroscopy

Arthroscopy is the direct visual examination of the interior of a joint by means of a specially designed fiberoptic endoscope and is frequently associated with a surgical procedure. It is most commonly done for the diagnosis of athletic injuries (meniscus, patella, condyle, extrasynovial area, and synovium) and for the differential diagnosis of acute or chronic joint disorders. For example, degenerative processes can be accurately differentiated from injuries. Postoperative rehabilitation programs can be initiated to shorten recovery periods. Arthroscopy can also assess response to treatment or identify whether other corrective procedures are indicated.

Although the knee is the joint most frequently examined, the shoulder, ankle, hip, elbow, wrist, and metacarpophalangeal joints can also be explored. Calcium deposits, biopsy specimens, loose bodies, bone spurs, torn meniscus or cartilage, and scar tissue can be removed during the procedure. Currently, many of these procedures are performed in an ambulatory surgical setting.

Clinical Alert

Arthroscopy is an instrument and equipment-dependent procedure (Phippen & Mells, 2000) (eg, video and irrigation failure, scratches on lens).

Reference Values

Normal Normal joint: normal vasculature and color of the synovium, capsule, menisci, ligaments, and articular cartilage

Procedure

1. Be aware that the examination is usually performed under general or spinal anesthesia for the following reasons:
   a. The joint is very painful.
   b. Definitive treatment or surgical intervention can be done at the same time if within the realm of arthroscopic surgery.
   c. An inflated tourniquet may be used during part of the procedure to minimize bleeding at the site.
   d. Complete muscle relaxation permits a thorough examination and eliminates the risk of inadvertent patient
movement while the arthroscope is in the joint.
2. Start an intravenous line. Drape and prep the surgical site according to institutional protocols. Attach proper monitoring equipment to the patient.
3. Apply a tourniquet to the appropriate area (by use of an elastic bandage or elevation), then insert an arthroscope into the joint through a small insertion. Some surgeons choose not to inflate the tourniquet unless bleeding cannot be controlled by irrigation.
4. Aspirate the joint, then perform continuous irrigation and flushing throughout the procedure.
5. Collect joint washings and examine for loose bodies or cartilage fragments.
6. Examine all parts of the joint carefully. Take photographs or videotapes of the procedure. The physician may choose to perform surgical interventions for problems that can be corrected via arthroscopy.
7. As you withdraw the arthroscope, accessory pieces, and irrigating needles slowly, compress the joint to squeeze out excess irrigation fluid.
8. Inject steroids or local anesthetics into the joint for postoperative pain control and reduction of inflammation. Close the wounds with sutures or adhesive strips, and apply small dressings to the wound or wounds (eg, two to three small incisions for the knee joint). Apply compressive dressings and splints or immobilizers.
9. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications Abnormal results reveal the following conditions:
1. Torn or displaced meniscus or cartilage (symptoms relate to clicking, locking, and/or swelling of the joint)
2. Trapped synovium
3. Loose fragments of joint contents
4. Torn or ruptured ligaments
5. Necrosis
6. Nerve entrapment
7. Fractures or nonunion of fractures
8. Ganglions
9. Infections
10. Degenerative disease. Osteochondritis dissecans: inflammation of bone or cartilage occurs when a cartilage fragment and underlying bone detach from the articular surface (common in the knee)
11. Chronic inflammatory arthritis
12. Secondary osteoarthritis caused by injury, metabolic disorders, and wearing away of weight-bearing joints
13. Chondromalacia of femoral condyle (wearing down of back of kneecap, often producing a grinding sensation)

Interfering Factors Ankylosis, fibrosis, sepsis, or presence of contrast agent from previous arthrogram may affect results.

Interventions

Pretest Patient Preparation

1. Make sure the history and physical examination, requisite laboratory work, x-ray films, and other preoperative requirements are completed, reviewed, and documented on the patient's record.
2. Explain the purpose and procedure of the test. The patient should fast from midnight before the examination unless otherwise ordered (eg, if scheduled late in the day, a liquid breakfast may be permitted).
3. Be aware that a properly signed and witnessed permit must be completed (see Chap. 1).
4. Check peripheral pulses in the operative area. The surgical site is prepped, positioned, and draped according to institutional protocols. An intravenous line is started.
5. Teach crutch-walking before the procedure if its necessity is anticipated postoperatively.
6. Follow Chapter 1 guidelines regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Assess vital signs, bleeding, neurologic status, and circulatory status of the affected extremity (eg, color, pulse, temperature, capillary refill times, sensation, and motion).
2. Apply ice immediately and, if ordered, elevate the extremity to minimize swelling and pain. Dressing changes and suture removal are performed at the physician's discretion. The dressing must be kept clean and dry. Notify the physician of unusual bleeding or swelling.
3. Administer appropriate pain medication.
4. Remember that the patient can usually be ambulatory after recovery from the anesthetic. Crutches may be used. Degree of weight-bearing and joint motion is at the discretion of the physician; however, patient should be cautioned to avoid excessive joint use for at least 24 to 48 hours.
5. Exercise and physical therapy may be ordered postoperatively. These are designed to strengthen and maximize use of the joint.
6. Make arrangements for transportation by another person preoperatively if the patient is discharged the same day as the procedure. The patient should not drive for at least 24 hours.
7. Advise the patient to consume no alcohol for 24 hours after the procedure. Progress diet from fluid to solid foods as tolerated.
8. Instruct the patient to report fever, altered sensation, numbness, tingling, coldness, dizziness (ie, bluish color), swelling, bleeding, or abnormal pain to the physician immediately. Mild soreness and a mild grinding sensation for a few days are normal.
9. Interpret test outcomes and counsel appropriately.
10. Follow guidelines in Chapter 1 for safe, effective, informed posttest care. Provide written discharge instructions.
Clinical Alert

1. Arthroscopy is usually contraindicated if ankylosis or fibrosis is present because it is very difficult to maneuver the examining instrument in this type of joint.
2. For knee arthroscopy, the posterior approach is not used because of the neurovascular structures present in that area.
3. Do not place pillows under the knee; flexion contractures can occur as a result. If the patient's leg is ordered to be elevated, make sure the entire leg is elevated in a straight position. The knee is not flexed because a flexion contracture may result. Pad pressure points such as the heel.
4. If there is risk for sepsis or if sepsis is present in any part of the body, the procedure should not be done.
5. Arthroscopy is usually not done < 7 to 10 days after arthrography because chemical synovitis caused by a contrast medium can adversely affect the visual examination. However, it may be necessary to perform arthroscopy if the patient is experiencing severe pain. In this case, the joint must be thoroughly irrigated to remove contrast medium.
6. Be alert for signs of thrombophlebitis postoperatively. Instruct patient to watch for calf tenderness, pain, and heat and to report these symptoms to the physician immediately. Warn the patient not to massage the affected area.
7. Other complications may include hemarthrosis, adhesions, neurovascular injury, pulmonary embolus, effusion, scarring, and compartmental syndrome as a result of swelling. Compartmental syndrome is a musculoskeletal complication that occurs most commonly in the forearm or leg. The compartment of fascia surrounding muscles does not expand when bleeding or edema occurs. Consequently, the neurovascular status of the extremity may be severely compromised. This presents an emergency situation that usually requires surgical intervention to release pressure. Assess the neurovascular status of an affected extremity frequently for 24 hours after the procedure.

Sinus Endoscopy

Sinus endoscopy visualizes the anterior ethmoid, middle turbinate region, and middle meatus sinus areas. Although the purposes of sinus endoscopy are primarily to relieve infection and other symptoms of inflammation and to alter structural abnormalities in these areas, it can also be a valuable diagnostic tool. Retained secretions may contribute to chronic recurrent sinus infections, which may lead to systemic infections, cyst formation, or mucoceles that can erode sinus walls into areas of the eyeball, eye orbit, or brain.

Patients having recurrent episodes of acute or chronic sinusitis that are not responsive to antibiotic and/or allergy therapy are candidates for sinus endoscopy as both a diagnostic and therapeutic modality.

Reference Values

Normal
Normal sinuses or resolution of sinus disease

Procedure
Sinus endoscopy may be performed as an outpatient or office procedure. Normally, the diagnostic procedure is performed in the office. More extensive examination and operative procedures normally require outpatient admission to a health care facility or special diagnostic center.

1. Spray a cocaine solution of select concentration into the nares to produce local anesthesia. Introduce the endoscope to permit visualization of the nasal interior; the sinus cavities are not opened. Some patients become very talkative and euphoric as a response to cocaine.
2. Be aware that sinus computed axial tomography (CT, CAT) scans and magnetic resonance imaging (MRI) may be necessary adjuncts to this procedure to permit visualization of areas not accessible through endoscopy.
3. Perform treatment for underlying disease or malformations using local or general anesthesia and medications to achieve state of conscious sedation. Diagnostic and surgical techniques vary according to preoperative findings.
4. Remember that endoscopes using a fiberoptic light delivery system are the mainstay of visualization for diagnosis and last treatment.
5. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications
Abnormalities that may be revealed include the following conditions:

1. Chronic sinusitis (edematous or polypoid mucosa)
2. Cysts
3. Mucocele
4. Sinus erosion
5. Anatomic deformities or obstructions
6. Pathologic sinus discharge (infectious process)
7. Enlarged middle turbinates

Interventions

Pretest Patient Preparation

1. Explain test purpose, benefits, risks, and procedure. (Steps 2 through 6 refer to treatment modalities.) The procedure may take place in the office or outpatient hospital setting.
2. Review and document in the health care record a properly signed and witnessed surgical consent form (see Chap. 1), appropriate laboratory and diagnostic test results, history and physical examination, current drug therapies, and allergies before the procedure.
3. Preprocedure preparation may require the patient to:
   a. Be processed through preadmission testing if procedure will be done in a hospital surgical setting
   b. Fast from midnight the day of the procedure
   c. Remove facial prostheses, dentures, hairpieces, and jewelry before the procedure
d. Have an intravenous line placed

4. Have the patient assume a supine position in the surgical suite. Prep the face and throat according to established protocols, and properly drape the area. Tape eye pads in place to protect the eyes from injury. Perform other positioning and pressure-point padding as necessary.

5. Administer intravenous sedation as needed. Spray the nose with a topical anesthetic, and inject a small amount of 1% lidocaine with 1:200,000 aqueous epinephrine into the appropriate areas (unless contraindicated because of allergy or for other reasons) to provide anesthesia and control of bleeding. Refer to Appendix C for intravenous conscious sedation precautions.

6. Fill a 10-mL syringe with antibiotic ointment at the end of the procedure. Use a small catheter attached to the syringe tip to direct ointment to the appropriate areas. Tape a small (2- × 2-inch) “mustache dressing” to the end of the nose to collect secretions and blood. Usually, this dressing can be changed as needed. Insert nasal packing into the nares.

7. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

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**Posttest Patient Aftercare**

1. Encourage oral fluids after nausea or vomiting has resolved; the patient may experience nausea or vomiting if blood is swallowed because blood is irritating to the gastrointestinal system.

2. Remember that postprocedural instructions may include the following:
   a. Take prescribed medications as ordered (usually a broad-spectrum antibiotic and pain medication). Soothing gargles may be ordered.
   b. Report excessive bleeding or sinus discharge, unusual pain, fever, nausea or vomiting, or visual problems immediately. Provide patient with phone numbers of hospital and physician and instruct him or her to contact the physician (or the outpatient surgical department or emergency department if unable to reach physician) in the event of an emergency. This process may differ according to various health insurance regulations and protocols.
   c. Do not allow the patient to drive or sign legal documents for 24 hours because the effects of anesthetics and sedation.

3. Follow the usual cautions involved in the care of any person having received sedation and analgesia. The patient who has received drugs to achieve conscious sedation may require closer monitoring, positioning on the side to prevent aspiration, and a longer recovery time than those who receive local anesthesia.

4. Interpret test outcomes and counsel appropriately about possible treatment (medications [eg, steroids, antibiotics]). Numbness of the face may continue for several weeks.

5. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care. Provide written discharge instructions.

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**Clinical Alert**

1. Sinuses are poorly visualized through routine sinus x-ray films.

2. If sinus problems appear to be related to dental problems, the patient should see a dentist or oral surgeon before sinus endoscopy is performed.

3. Severe nasal-septal deviation must be corrected before endoscopy.

4. Potential complications include periorbital bleeding, cerebrospinal fluid leak, cellulitis, visual disturbances, and subcutaneous orbital emphysema.

5. Direct trauma to the nasofrontal duct is associated with increased risk for postoperative stenosis.

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**BIBLIOGRAPHY**


Clinical News Infection Control: TB and the link to bronchoscopies. AJN 98(4): 9, April 1998; Source: JAMA 278–1077, 1093, 1111, 1997 (editorial)


Finkelmeier BA: Cardiothoracic Surgical Nursing, 2nd ed. Philadelphia, Lippincott Williams & Wilkins, 2000


Lefton HB, Pelchman J, Harnatz A: Colon cancer screening and the evaluation and follow-up of colonic polyps. Primary Care 23(3): 515–523, 1996


OVERVIEW OF ULTRASOUND STUDIES

Ultrasonography is a noninvasive procedure for visualizing soft tissue structures of the body by recording the reflection of inaudible sound waves directed into the tissues. The diagnostic procedure, which requires very little patient preparation, is now used in many branches of medicine for accurate diagnosis of certain pathologic conditions (Chart 13.1). It may be used diagnostically with the obstetric, gynecologic, or cardiac patient and in patients with abnormal conditions of the kidney, pancreas, gallbladder, lymph nodes, liver, spleen, thyroid, or peripheral blood vessels. Frequently, it is used in conjunction with radiology or nuclear medicine scans. The procedure is relatively quick (often requiring only a few minutes to an hour) and causes little discomfort. No harmful effects have yet been established at the low intensities that are used (<100 mW/cm²). However, as with any diagnostic procedure, ultrasound should not be used frivolously. The terms ultrasound and sonogram are used interchangeably.
<table>
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<tr>
<th>Chart 13.1 Uses of Ultrasound</th>
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**Principles and Techniques**

Ultrasound uses high-frequency sound waves to produce an “echo map” that characterizes the position, size, form, and nature of soft tissue organs. Echoes of varying strength are produced by different types of tissues and are displayed as a visual pattern after computer processing of the echo information. The capability of acquiring real-time images means that ultrasound can readily demonstrate motion, as in the fetus or the heart. Ultrasound, however, cannot appropriately image air-filled structures such as the lungs.

**Doppler Method** A phenomenon that accompanies movement, the Doppler effect, can be combined with diagnostic ultrasound imaging to produce duplex scans. Duplex scans provide anatomic visualization of blood vessels and a graphic representation of blood flow characteristics. Flow direction, velocity, and the presence of flow disturbances can readily be assessed. Certain equipment is capable of advanced Doppler imaging techniques, such as:

1. **Color Doppler** imaging provides a color-coded depiction of selected blood flow parameters.
2. A technology known as color Doppler energy, power Doppler, or color angiography is sensitive to very low blood velocity states and is often used to evaluate blood flow through solid organs.
3. **B-flow Doppler** images the blood itself, producing images that resemble an angiogram.

These techniques establish the patency of a given blood vessel and are useful in investigating perfusion to an organ or mass. They are also helpful in evaluating complications in transplanted organs.

**General Procedure**

1. A gel or lubricant is applied to the skin over the area to be examined to conduct the sound waves.
2. An operator, known as a **sonographer**, holds a microphone-like device called a **transducer**. The transducer is moved over a specific body part, producing a display that is viewed on the monitor.
3. Sonography of structures in the abdominal region often require that the patient control breathing patterns. Deep inspiration and exhalation may be used.
4. Selected images are recorded for documentation purposes.
5. The examination causes no physical pain. However, in certain applications, pressure may be applied to the transducer, causing some degree of discomfort. Long examinations may leave the patient feeling tired.
6. Tests usually take 20 to 45 minutes. This is the actual procedure time and does not include waiting and preparation times.
7. Some examinations require the patient to fast or to have a filled urinary bladder. Each examining department determines its own guidelines for patient preparation.

Advances in technology have allowed the development of very small, high-resolution transducers. Catheter-sized transducers are used to visualize blood vessels “from the inside out” during angiographic procedures. Endoscopic ultrasound is used to evaluate gastrointestinal lesions and may be used to visualize pancreatic biliary structures. Small transducers passed through the esophagus permit exquisite visualization of the heart during transesophageal echocardiography (TEE). Slim transducers are introduced into the vagina to visualize gynecologic anatomy. Transrectal visualization of the prostate gland is an accepted method of screening for disease in the organ. Of course, before introduction into the body, these small transducers are properly cleansed and/or draped.

**Implications of Ultrasound Studies**

**Benefits and Risks**

1. Ultrasonography is a noninvasive procedure with no radiation risk to patient or examiner.
2. It requires little, if any, patient preparation and aftercare.
3. As far as is known, the examination can be repeated as often as necessary without being injurious to the patient. No harmful cumulative effect has been seen.
4. Because ultrasound studies demonstrate structure rather than function, they may be useful for patients whose organ function is impaired.
5. Ultrasound is useful in the detection and examination of moving parts, such as the heart.
6. It does not require the injection of contrast materials or isotopes or ingestion of opaque materials.

**Disadvantages**

1. An extremely skilled examiner is required to operate the transducer. The scans should be read immediately and interpreted for adequacy. If the scans are not satisfactory, the examination must be repeated.
2. Air-filled structures (eg, the lungs) cannot be studied by ultrasonography.
3. Certain patients (eg, restless children, extremely obese patients) cannot be studied adequately unless they are specially prepared.

**Difficult-to-Study Patients** The following general categories of patients may provide some difficulties in ultrasound studies:

1. Postoperative patients and those with abdominal scars: The area surrounding an incision is to be avoided whenever possible. If a scan must be performed over an incision, the dressing must be removed and a sterile coupling agent and probe must be employed.
2. Children and agitated adults: Because the procedure requires the patient to remain still, some patients may need to be sedated so that their movements do not cause artifacts.
3. Obese patients: Certain patients cannot be studied adequately in any case. For example, it may be difficult to obtain an accurate scan on a very obese patient, owing to alteration of the sound beam by fatty tissue. There is no preparation that would help here.

**Interfering Factors**

1. Barium has an adverse effect on the quality of abdominal studies, so sonograms should be scheduled before barium studies are done.
2. If the patient has a large amount of gas in the bowel, the examination may be rescheduled because air (bowel gas) is a very strong reflector of sound and does not permit accurate visualization.

**OBSTETRIC AND GYNECOLOGIC SONOGRAMS**

**Obstetric Sonogram**

Ultrasound studies of the obstetric patient are valuable in (1) confirming pregnancy; (2) facilitating amniocentesis by locating a suitable pool of amniotic fluid; (3) determining fetal age; (4) confirming multiple pregnancy; (5) ascertaining whether fetal growth is normal, through sequential studies; (6) determining fetal viability; (7) localizing placenta; (8)
confirming masses associated with pregnancy; (9) identifying postmature pregnancy (increased amount of amniotic fluid and degree of placental calcification); and (10) as a guidance method for chorionic villus sampling (CVS), embryo transfer, intrauterine device (IUD) extraction, and percutaneous umbilical vein sampling (PUVS). A pregnancy can be dated with considerable accuracy if a sonogram is done at 20 weeks' gestation and a follow-up scan is done at 32 weeks' gestation. This validation is most important when early delivery is anticipated and prematurity is to be avoided. Conditions in which determination of pregnancy duration is useful include maternal diabetes, Rh immunization, and preterm labor (Chart 13.2).

<table>
<thead>
<tr>
<th>Chart 13.2 Major Uses of Obstetric Ultrasound—Levels I and II *</th>
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<td><strong>Indications during First Trimester</strong></td>
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<td>Confirm pregnancy</td>
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<td>Confirm viability</td>
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<tr>
<td>Rule out ectopic pregnancy in the first trimester</td>
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<td>Confirm gestational age †</td>
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<tr>
<td>Birth control pill use</td>
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<td>Irregular menses</td>
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<td>No dates</td>
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<td>Postpartum pregnancy</td>
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<td>Previous complicated pregnancy</td>
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<td>Cesarean delivery</td>
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<td>Rh incompatibility</td>
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<tr>
<td>Diabetes mellitus</td>
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<tr>
<td>Fetal growth retardation</td>
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<tr>
<td>Clarify discrepancy between dates and size</td>
</tr>
<tr>
<td>If large for dates rule out: Leiomyomas</td>
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<tr>
<td>Bicornuate uterus</td>
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<td>Adnexal mass</td>
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<td>Multiple gestation</td>
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<td>Poor estimate of dates</td>
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<td>Missed abortion</td>
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<td>Blighted ovium</td>
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<td>As guidance method for:</td>
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<td><strong>Indications during Second Trimester</strong></td>
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<tr>
<td>Establish or confirm dates †</td>
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<tr>
<td>If no fetal heart tones: Clarify discrepancy between dates and size</td>
</tr>
<tr>
<td>If large for dates, rule out: Poor estimate of dates</td>
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<tr>
<td>Molar pregnancy</td>
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<tr>
<td>Multiple gestation</td>
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<tr>
<td>Leiomyomas</td>
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<td>Polyhydramnios</td>
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<tr>
<td>Congenital anomalies</td>
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<tr>
<td>If small for dates, rule out: Poor estimate of dates</td>
</tr>
<tr>
<td>Fetal growth retardation</td>
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<td>Congenital anomalies</td>
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<tr>
<td>Oligohydramnios</td>
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<tr>
<td>If history of bleeding, rule out total placenta previa</td>
</tr>
<tr>
<td>If Rh incompatibility, rule out fetal hydrops</td>
</tr>
<tr>
<td>Evaluation and follow-up of suspected fetal anomalies</td>
</tr>
</tbody>
</table>
If Rh incompatibility, rule out fetal hydrops

Evaluation and follow-up of suspected fetal anomalies

**Indications during Third Trimester**

If no fetal heart tones:

- Clarify discrepancy between Dates and size

If large for dates, rule out:

- Macrosomia (diabetes mellitus)
- Multiple gestation
- Polyhydramnios
- Congenital anomalies
- Poor estimate of dates ‡

If small for dates, rule out:

- Fetal growth retardation
- Oligohydramnios
- Congenital anomalies
- Poor estimate of dates ‡

Determine fetal position, rule out:

- Breech
- Transverse lie

If history of bleeding, rule out:

- Placenta previa
- Abruptio placenta

Determine fetal maturity

- Amniocentesis for lecithin/sphingomyelin ratio
- Placental maturity (grade 0–3)

Evaluation and follow-up of suspected fetal anomalies

- Estimation of fetal weight

**Footnotes**

*Ultrasound is a diagnostic tool for assessment of fetal age, health, and growth. Level I ultrasound is performed to assess gestational age, number of fetuses, fetal viability, and the placenta. Level II ultrasound is used for assessment of specific congenital anomalies or abnormalities. (See also Fetal Echocardiography, page 870.) †Accuracy ± 3 days. ‡Accuracy ± 1 to 1.5 days.

The pregnant uterus is ideal for echographic evaluation because the amniotic fluid–filled uterus provides strong transmitting interfaces between the fluid, placenta, and fetus. Ultrasonography has become the method of choice for evaluating the fetus and placenta, eliminating the need for the potentially injurious radiographic studies that were used previously.

**Reference Values**

**Normal Obstetric Sonogram** Normal image of placental position, size, and structure

- Normal fetal position and size with evidence of fetal movement, cardiac activity, and breathing activity
- Adequate amniotic fluid volume
- Normal fetal intracranial, thoracic, and abdominopelvic anatomy; four limbs visualized

**Procedure**

1. Have the pregnant woman lie on her back with her abdomen exposed during the test. This may cause some shortness of breath and supine hypotensive syndrome, which can be relieved by elevating the upper body or turning the patient onto her side.

2. Perform the transabdominal scan in the second trimester while the patient has a full bladder. Exceptions are made when the scan is performed to locate the placenta before amniocentesis, for evaluation of an incompetent cervix, or during labor and delivery. A full bladder allows the examiner to assess the true position of the placenta, repositions the uterus, and acts as a sonic window to the pelvic organs.

3. Apply a coupling agent (special transmission gel, lotion, or mineral oil) liberally to the skin to prevent air from absorbing sound waves. The sonographer slowly moves the transducer over the entire abdomen to obtain a picture of the uterine contents.

4. Perform an endovaginal (transvaginal) scan during the first trimester. Most laboratories use a transvaginal approach when performing obstetric sonograms at this time. This method does not require a full bladder. A slim transducer, properly covered and lubricated, is gently introduced into the vagina. Because the sound waves do not need to traverse abdominal tissue, exquisite image detail is produced. Check with your laboratory to determine the approach to be used.

5. Tell the patient that the examining time is about 30 to 60 minutes.

6. See Chapter 1 guidelines for intratest care.

**Clinical Implications**

1. During the first trimester, the following information can be obtained:
   a. Number, size, and location of gestational sacs
   b. Presence or absence of fetal cardiac activity and body movement
   c. Presence or absence of uterine abnormalities (eg, bicornuate uterus, fibroids) or adnexal masses (eg, ovarian cyst, ectopic pregnancy)
   d. Pregnancy dating (eg, biparietal diameter, crown–rump length)
   e. Presence and location of an IUD

2. During the second and third trimesters, ultrasound can be performed to obtain the following information:
   a. Fetal viability, number, position, gestational age, growth pattern, and structural abnormalities
Amniotic fluid volume

Fetal growth:
Early diagnosis of fetal structural abnormalities makes the following choices possible:
1. Intrauterine surgery or other prenatal therapy
2. Discontinuation of pregnancy
3. Preparation of the family for care of a child with a disorder or planning of other options

Gestational age:
1. Identification of amniotic fluid measuring at least 1 cm is associated with normal fetus status. The presence of one
   pocket measuring < 1 cm or the absence of a pocket is abnormal; it is associated with increased risk for perinatal
   death.
2. Fetal heart activity can be demonstrated at 5 weeks’ gestation in most cases. This information is
   helpful in establishing dates and in the management of vaginal bleeding. Molar pregnancies and incomplete,
   complete, and missed abortions can be differentiated.
3. Fetal viability: Fetal heart activity can be demonstrated at 5 weeks’ gestation in most cases. This information is
   helpful in establishing dates and in the management of vaginal bleeding. Molar pregnancies and incomplete,
   complete, and missed abortions can be differentiated.

Fetal growth:
The conditions that serve as indicators for ultrasound assessment of fetal growth include poor
maternal weight gain or pattern of weight gain, previous intrauterine growth retardation (IUGR), chronic infection,
ingestion of drugs such as anticonvulsants or heroin, maternal diabetes, pregnancy-induced or other hypertension,
multiple pregnancy, and other medical or surgical complications. Serial evaluation of biparietal diameter and limb
length can help differentiate between wrong dates and IUGR. Doppler evaluation of the umbilical artery, uterine
artery, and fetal aorta can also assist in the detection of IUGR. IUGR can be symmetric (the fetus is small in all
measurements) or asymmetric (head and body growth vary). Symmetric IUGR may be caused by low genetic
growth potential, intrauterine infection, maternal undernutrition, heavy smoking by the mother, or chromosomal
aberration. Asymmetric IUGR may reflect placental insufficiency secondary to hypertension, cardiovascular
disease, or renal disease. Depending on the probable cause, the therapy varies.

Artifacts may be produced when the transducer is moved out of contact with the skin. This can be resolved by
adding more coupling agent to the skin and repeating the scan.

Fetal anatomy:
Depending on the gestational age, the following structures may be identified: intracranial anatomy,
neck, spine, heart, stomach, small bowel, liver, kidneys, bladder, and extremities. Structural defects may be
identified before delivery. The following are examples of structural defects that may be diagnosed by ultrasound:
hydrocephaly, anencephaly, and myelomeningocele are often associated with polyhydramnios. Potter's syndrome
(renal agenesis) is associated with oligohydramnios defects (dwarfism, achondroplasia, osteogenesis imperfecta)
diaphragmatic hernias. Other structural anomalies that can be diagnosed by ultrasound are pleural effusion
(after 20 weeks), intestinal atresias or obstruction (early pregnancy to second trimester), hydrenephrosis, and
bladder outlet obstruction (second trimester to term with fetal surgery available). Two-dimensional studies of the
heart, together with echocardiography, allow diagnosis of congenital cardiac lesions and prenatal treatment of
cardiac arrhythmias.

Detection of fetal death:
Inability to visualize the fetal heart beating and separation of bones in the fetal head are
signs of death. With real-time scanning, the absence of cardiac motion for 3 minutes is diagnostic of fetal demise.

Placental position and function:
The site of implantation (eg, anterior, posterior, fundal, in lower segment) can be
described, as can location of the placenta on the other side of midline. The pattern of uterine and placental growth
and the fullness of the bladder influence the apparent location of the placenta. For example, when ultrasound
scanning is done in the second trimester, the placenta seems to be overlying the os in 15% to 20% of all
pregnancies. At term, however, the evidence of placenta previa is only 0.5%. Therefore, the diagnosis of placenta
previa can seldom be confirmed until the third trimester. Placenta abruptio (premature separation of placenta) can
also be identified. A transverse scan through the umbilical cord confirms the number of vessels. Doppler of the cord
detects flow abnormalities.

Fetal well-being: Ultrasound findings are a major component of the biophysical profiles. The following physiologic
measurements can be accomplished with ultrasound: heart rate and regularity, fetal breathing movements, urine
production (after serial measurements of bladder volume), fetal limb and head movements, and analysis of vascular
wave forms from fetal circulation. Fetal breathing movements are decreased with maternal smoking and alcohol use
and increased with hyperglycemia. Fetal limb and head movements serve as an index of neurologic development.
Identification of amniotic fluid measuring at least 1 cm is associated with normal fetus status. The presence of one
pocket measuring < 1 cm or the absence of a pocket is abnormal; it is associated with increased risk for perinatal
death.

Assessment of multiple pregnancy:
Two or more gestational sacs, each containing an embryo, may be seen after 6
weeks. Of twin pregnancies diagnosed in the first trimester, only about 30% will deliver twins, owing to loss or
absorption of one fetus. Of value is assessment of the relative fetal growth of twins when IUGR or twin-to-twin
transfusion to suspected. One cannot unequivocally diagnose whether twins are monozygotes or heterozygotes
with ultrasound alone unless fetuses of opposite sex are evident. Routine ultrasound cannot totally be relied on to
exclude the possibility of triplets or quadruplets, instead of only twins.

If the fetal position and amniotic fluid volumes are favorable, fetal sex can be determined by visualization of the
genitalia. It must be cautioned, however, that sex determination is not the purpose of obstetric sonography.

Interfering Factors
1. Artifacts may be produced when the transducer is moved out of contact with the skin. This can be resolved by
   adding more coupling agent to the skin and repeating the scan.
2. Artifacts (reverberation) may be produced by echoes emanating from the same surface several times. This can be
   avoided by careful positioning of the transducer.
3. A posterior placental site may be difficult to identify because of the angulation of the reflecting surface or
   insufficient penetration of the sound beam owing to the patient’s size.

Interventions
Pretest Patient Preparation
1. A brief explanation of the procedure to be performed is given, emphasizing that it is not uncomfortable or painful
and does not involve ionizing radiation that might be harmful to the mother or fetus. The studies can be repeated
without harm, but the procedure is being studied carefully to determine whether there are any long-term adverse side effects. Benefits of the procedure should be explained.

2. Most studies are performed by a transabdominal approach with a full bladder. The patient is asked to drink five to six glasses of fluid (water or juice) about 1 to 2 hours before the examination. If she is unable to drink, intravenous fluids may be administered. She is asked to refrain from voiding until the examination is complete. Tell the patient that she will have a strong urge to void during the examination. Discomfort caused by pressure applied over a full bladder may be experienced. If the bladder is not sufficiently filled, three to four 8-oz glasses of water should be ingested, with rescanning done 30 to 45 minutes later.

3. Most laboratories use a transvaginal (endovaginal) approach during the first trimester of pregnancy. No patient preparation is required for this method. Contact the laboratory performing the study to determine the method to be used.

4. Explain that a liberal coating of coupling agent must be applied to the skin so that there is no air between the skin and the transducer and to allow for easy movement of the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.

5. The woman may face the screen, and the sonographer may explain the images in basic terms. In some institutions, the father is encouraged to observe the testing. A photograph or videotape for the family to keep is sometimes provided.

6. See Chapter 1 guidelines for safe, effective informed pretest care.

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**Clinical Alert**

1. A full bladder may not be needed or desired for patients in the late stages of pregnancy or active labor. However, if a full bladder is required and the woman has not been instructed to report with a full bladder, at least another hour of waiting time may be needed before the examination can begin.

2. A transvaginal (endovaginal) scan does not require the patient to have a full bladder. Contact the laboratory to determine method to be used.

3. Endovaginal studies typically involve the use of a latex condom to sheath the transducer before it is inserted into the vaginal vault. Contact the laboratory if the patient has known or suspected latex sensitivity.

4. Fetal age determinations are most accurate during the crown–rump stage in the first trimester. The next most accurate time for age estimation is during the second trimester. Sonographic dating during the third trimester has a large margin of error (up to ± 3 weeks).

5. If fetal death is suspected, careful and considerate counseling and support are offered to parents.

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**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel appropriately. Explain possible need for follow-up testing (eg, fetal echocardiography) and/or treatment: medical (stimulate early onset of labor) or surgical (fetal surgery or immediate surgery for ectopic pregnancy).

2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

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**Fetal Echocardiography (Fetal Doppler)**

Fetal echocardiography is performed after the detection of a potential cardiac abnormality during an obstetric sonogram or in patients with a strong history of congenital cardiovascular disease. Additionally, women exposed to cardiac teratogens are usually advised to have this study. Not a screening procedure, fetal echocardiograms are most commonly performed in specialized laboratories or teaching hospitals. The heart is imaged in numerous planes, using pulsed Doppler and M-mode tracings (see Heart Sonogram, page 895), similar to an electrocardiogram. Valves and other cardiac structures are measured, and blood velocities and volumes are calculated.

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**Reference Values**

**Normal Sonogram** Normal structure of heart and great vessels Normal heart rate and rhythm, with proper hemodynamic flow through heart and great valves

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**Procedure**

1. Perform the fetal echocardiogram in the same manner as a routine obstetric scan, which also requires similar patient preparation. The pregnant patient lies on her back with the abdomen exposed. Couplant is applied to the skin, and a transducer is moved across the abdomen. Unless combined with an obstetric sonogram, the fetal echocardiogram does not require the mother to have a full bladder.

2. See Chapter 1 guidelines for intratest care.

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**Clinical Implications** Abnormalities detected during fetal echocardiography include:

1. Cardiac arrhythmias
2. Septal defects, including tetralogy of Fallot
3. Hypoplastic heart syndrome
4. Valvular abnormalities, including Ebstein's anomaly
5. Cardiac tumors
6. Vessel abnormalities, including coarctation of aorta, transposition, aortic stenosis, truncus arteriosus, and pulmonary stenosis
Clinical Alert

Optimal fetal echocardiographic studies are performed between 18 and 22 gestational weeks. Before 18 weeks, the fetal heart is too small, and after 22 weeks, image quality may be degraded by overlying structures.

Interfering Factors
Same as for obstetric sonogram.

Interventions

Pretest Patient Preparation
Same as for obstetric sonogram.

Posttest Patient Aftercare
Same as for obstetric sonogram.

Pelvic Gynecologic (GYN) Sonogram; Pelvic (Uterine Mass) Ultrasound Diagnosis; Intrauterine Device (IUD) Localization

The pelvic gynecologic ultrasound study examines the area from the umbilicus to the pubic bone in women. It may be used in the evaluation of pelvic masses, to determine the position of an IUD, to evaluate postmenopausal bleeding, or to aid in the diagnosis of cysts and tumors. Information can be provided on the size, location, and structure of masses. Spectral or color Doppler can be applied to pelvic vessels, demonstrating normal flow changes associated with the menstrual cycle, and can evaluate abnormal flow patterns to masses/tumors. The examination cannot provide a definitive diagnosis of pathology but can be used as an adjunct procedure when the diagnosis is not readily apparent. It is also used in treatment planning and follow-up radiation therapy for gynecologic cancer. Additionally, follicle development after infertility treatment can be monitored.

This test may be performed by a transvaginal method whereby a slim, covered, lubricated transducer is gently introduced into the vagina. A full bladder is not required. Because the sound waves do not need to transverse abdominal tissue, exquisite image detail is produced. This approach is most advantageous for examining the obese patient, the patient with a retroverted uterus, or the patient who has difficulty maintaining bladder distention. The transvaginal method is the approach of choice in monitoring follicular size during fertility workups and during aspiration of follicles for in vitro fertilization.

For pelvic sonograms using the transabdominal approach, a full bladder is necessary. The distended bladder serves four purposes: it acts as a "window" for transmission of the ultrasound beam; it pushes the uterus away from the pubic symphysis, thereby providing a less obstructed view; it pushes the bowel out of the pelvis; and it may be used as a reference for comparison in evaluating the internal characteristics of a mass under study.

Reference Values

Normal Sonogram
Normal pattern image of bladder, uterus, fallopian tubes, vagina, and surrounding structures

Procedure

Transabdominal Method

1. Have the patient lie on the back on the examining table during the test.
2. Apply a coupling agent to the area under study.
3. Place the active face of the transducer in contact with patient's skin and sweep across the area being studied.
4. Tell the patient that the examination time is about 30 minutes.

Transvaginal (Endovaginal) Method

1. Have the patient lie on an examining table with hips slightly elevated in a modified lithotomy position. Drape the patient.
2. Lubricate and introduce a slim vaginal transducer, protected by a condom or sterile sheath, into the vagina. Some laboratories prefer that the patient insert the transducer herself. Depth < 8 cm is all that is usually required.
3. Perform scans by using a slight rotation or movement of the handle and by varying the degree of transducer insertion. Typically, the transducer is inserted only a few inches into the vaginal vault.
4. Tell patient that the examination time is about 15 to 30 minutes.
5. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. Uterine abnormalities such as fibroids, intrauterine fluid collections, and variations in structure such as bicornuate uterus can be detected. Uterine and cervical carcinomas may be visualized, although definitive diagnosis of cancer cannot be made by sonography alone.
2. Endometrial abnormalities such as polyps can be visualized by sonography. This procedure involves distention of the endometrial canal with saline and subsequent ultrasound scanning. Very small adnexal masses may not be demonstrated by ultrasound studies. Masses identified on ultrasound may be evaluated in terms of size and consistency.
3. Cysts
   a. Ovarian cysts (the most common ovarian mass detected by ultrasound) appear as smoothly outlined, well-defined masses. Cysts cannot be confirmed as either malignant or benign, but ultrasound studies can increase the suspicion that a particular mass is malignant.
   b. A corpus luteum cyst is a single, simple cyst commonly visualized in early pregnancy.
   c. Theca-lutein cysts are associated with hydatidiform mole, choriocarcinoma, or multiple pregnancy.
   d. Because normal ovaries often have numerous visible small cysts, the diagnosis of polycystic ovaries is difficult to make on the basis of ultrasound alone.
   e. Dermoid cysts or benign ovarian teratomas may be found in young adult women and have an extremely variable appearance. Because of their echogenicity, they are often missed on ultrasound. The only initial clue may be an indentation of the urinary bladder. When a dermoid cyst is suspected on ultrasound, a pelvic radiograph should
be obtained.
4. Solid ovarian tumors such as fibromas, fibrosarcomas, Brenner's tumors, dygerminomas, and malignant teratomas are not differentiated by diagnostic ultrasound. Ultrasound documents the presence of a solid lesion but can go no further in narrowing the diagnosis.
5. Metastatic tumors of the ovary are common and may be solid or cystic in ultrasonic appearance. They are variable in size and are usually bilateral. Because ascites is often present, the pelvis and remainder of the abdomen should be scanned for fluid.
6. Pelvic inflammatory disease: Ultrasound differentiation between pelvic inflammatory disease and endometriosis is difficult. Evaluation of laboratory results and the clinical history leads to correct diagnosis. Other entities that may have similar ultrasonic presentation include appendicitis with rupture into the pelvis, chronic ectopic pregnancy, posttraumatic hemorrhage into the pelvis, and pelvic abscesses from various causes (eg, Crohn's disease, diverticulitis).
7. Bladder distortion: Any distortion of the bladder raises the possibility of an adjacent mass. Tumor, infection, and hemorrhage are the major causes of increased thickness of the urinary bladder wall. Masses such as calculi and catheters may be seen within the bladder lumen. Urinary bladder calculi are highly echogenic. A urinary bladder diverticulum appears as a cystic mass adjacent to the urinary bladder. It may be mistaken for a cystic mass arising from some other pelvic structure, so attempts are made to demonstrate its communication to the bladder.
8. Ultrasound studies can help to determine whether a pelvic mass is mobile.
9. Solid pelvic masses such as fibroids and malignant tumors may be differentiated from cystic masses, which show sound patterns similar to those of the bladder.
10. Lesions may be shown to have metastasized.
11. Studies may aid in the planning of tumor radiation therapy.
12. The position of an IUD may be determined.

Interfering Factors

1. Results may be only fair, may vary with the patient's habits and preparation (as described in Clinical Implications), and can be used only in conjunction with other studies. However, masses 1 cm and smaller can be seen with high-resolution equipment.
2. The success of a transabdominal scan depends on full bladder distention.

Interventions

Pretest Patient Preparation

1. Explain the purpose, benefits, and procedure of the test. Fasting is not required.
2. Have the patient drink four glasses of water or other liquid 1 hour before transabdominal scans. Advise the patient not to void until the test is over.
3. Contact the laboratory performing the study to determine method to be used. If a transvaginal (endovaginal) approach is to be used, no patient preparation is required.
4. Explain that a liberal coating of coupling agent must be applied to the skin so that there is no air between the skin and the transducer and to allow for easy movement of the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
5. Determine whether the patient has a latex sensitivity and communicate such sensitivities to the examining laboratory, if a transvaginal (endovaginal) approach is to be used. See latex precautions in Appendix B.
6. Reassure the patient that she will have no pain or discomfort.
7. See Chapter 1 guidelines regarding safe, effective, informed pretest care.

Clinical Alert

1. If the patient is taking nothing by mouth (NPO) or in certain emergency situations, the patient may be catheterized and the bladder filled through the catheter if a transabdominal approach is required.
2. Endovaginal studies, when indicated, typically involve the use of a latex condom to sheath the transducer before it is inserted into the vaginal vault. Contact the laboratory if the patient has a known or suspected latex sensitivity. See Appendix B regarding latex precautions.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately about possible further testing (biopsy with cytologic and histologic exam) and/or treatment (medical, pharmacologic, or surgical interventions).
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

ABDOMINAL SONOGRAMS

Kidney (Renal) Sonogram

The kidney sonogram is a noninvasive test used to visualize kidney parenchyma and associated structures, including renal blood vessels. This procedure is often performed after an intravenous pyelogram (IVP) to define and characterize mass lesions or the cause of a nonvisualized kidney. Because no contrast medium is administered, renal ultrasound is valuable for visualizing the kidneys of patients with iodine hypersensitivities. This procedure is also helpful in monitoring the status of a transplanted kidney, guiding stent and biopsy needle placement, and evaluating the progression of chronic conditions. Renal sonography is the preferred method for evaluating possible hydronephrosis in spinal cord injury patients.

Reference Values
Normal pattern image indicating normal size and position of kidneys, appropriate flow in renal vessels

Procedure

1. Have the patient lie quietly on an examining table. Scans are often performed with the patient in the decubitus position.
2. Apply warm oil or gel to the patient’s skin.
3. Ask the patient to inspire as deeply as possible for visualization of the upper parts of the kidney.
4. Tell the patient that the total study time varies from 15 to 30 minutes.
5. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. Abnormal pattern readings reveal:
   a. Cysts
   b. Solid masses
   c. Hydronephrosis
   d. Obstruction of ureters
   e. Calculi
2. Results provide information on the size, site, and internal structure of a nonfunctioning kidney.
3. Results differentiate between bilateral hydronephrosis, polycystic kidneys, and the small, end-stage kidneys of glomerulonephritis or pyelonephritis.
4. Results may be used to monitor kidney development in children with congenital hydronephrosis. This approach is safer than repeated IVP studies.
5. Perineal fluid collections such as those associated with complications of transplantation may be detected. These collections include abscesses, hematomas, urinomas, and lymphoceles.
6. Solid lesions may be differentiated from cystic lesions.
7. The spread of cancerous conditions from the kidney into the renal vein or inferior vena cava can be detected.
8. If ultrasound is combined with Doppler evaluations, the patency and flow characteristics of the renal vessels may be scrutinized.

Interfering Factors

1. Retained barium from radiology studies causes poor results.
2. Obesity adversely affects tissue visualization.

Interventions

Pretest Patient Preparation

1. Explain the purpose and procedure of the test.
2. Assure the patient that there is no pain involved and that the only discomfort is that caused by lying quietly for a long period.
3. Explain that a liberal coating of coupling agent must be applied to the skin so that there is no air between the skin and the transducer and to allow for easy movement of the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
4. Explain that the patient will be instructed to control breathing patterns while the images are being made.
5. Check with your ultrasound department for guidelines about fasting. It usually is not necessary but may be required in certain laboratories.
6. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately about further testing (CAT scans, biopsies) and/or treatment of chronic conditions.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

1. Scans cannot be done over open wounds or through dressings.
2. This examination must be performed before radiographic studies involving barium. If such scheduling is not possible, at least 24 hours must elapse between the barium procedure and the renal echogram.
3. Biopsies or drainage procedures are often done with ultrasound as a guide. If an invasive procedure is to be done, a surgical permit must be signed by the patient.

Urinary Bladder Sonogram

The urinary bladder sonogram is done as part of the investigation of possible bladder tumor and provides a simple method of estimating postvoid residual urine volume. This test reduces the need for urinary catheterization and the risk for subsequent urinary tract infection.

Reference Values

Normal Normal pattern image of the exact dimensions and contour of the bladder and little residual volume

Procedure
1. Instruct the patient to lie, with bladder fully distended, on the back on an examination table.
2. Apply a coupling agent to the anterior pelvic region to allow maximum penetration of the ultrasound beam.
3. Place the active face of the transducer in contact with the patient's skin and sweep across the area being studied.
4. Instruct the patient to void. This is typically done when the full-bladder scans are completed. Take additional images then to check for residual volume.
5. Tell patient that total examination time is about 10 to 20 minutes.
6. See Chapter 1 guidelines for intratest care.

Clinical Implications Abnormal results reveal the following:
1. Tumors of bladder
2. Cancerous extension to urinary bladder
3. Thickening of bladder wall
4. Masses posterior to bladder
5. Ureterocele

Interfering Factors
1. Residual barium from previous radiology studies affects test results.
2. Overlying gas or fat tissue affects test results.

Interventions
Pretest Patient Preparation
1. Explain the purpose, benefits, and procedure of the test.
2. Ask the patient to have a full bladder at the beginning, which is then emptied to complete the examination.
3. Assure the patient that there is no pain involved. Some discomfort may be experienced from maintaining a full urinary bladder.
4. Explain that a liberal coating of coupling agent must be applied to the skin so that there is no air between the skin and the transducer and to allow for easy movement of the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Patient may return to normal routines.
2. Interpret test outcomes and counsel about bladder abnormalities and possible further tests (cystoscopy) and treatment: medical (drugs) or surgical interventions.
3. Follow Chapter 1 guidelines regarding safe, effective, informed posttest care.

Hepatobiliary Sonogram; Gallbladder (GB) Ultrasound; Liver Ultrasound
These tests are helpful in differentiating hepatic disease from biliary obstruction. Unlike the oral cholecystogram, this procedure allows visualization of the gallbladder and ducts in patients with impaired liver function. Stones and evidence of cholecystitis are readily visualized. This procedure is indicated as an initial study for patients with right upper quadrant pain. It is also useful as a guide for biopsy or other interventional procedures. Posttransplantation color Doppler sonography of the reconstructed vessels is an important diagnostic tool.

Reference Values
Normal Normal size, position, and configuration of the gallbladder and bile ducts Normal adjacent liver tissue

Procedure
1. Ask the patient to lie quietly on an examination table. Scans usually are performed with the patient in the supine and decubitus positions.
2. Cover the skin with a layer of coupling gel, oil, or lotion.
3. Ask the patient to regulate breathing patterns as instructed during the examination.
4. Tell patient that total examination time is about 10 to 30 minutes.
5. See Chapter 1 guidelines for intratest care.

Clinical Implications
1. Gallbladder abnormal patterns reveal:
   a. Size variations
   b. Thickened wall, indicative of cholecystitis, adenomyomatosis, or tumor and commonly seen as a manifestation of cholecystopathy in patients with the acquired immunodeficiency syndrome (AIDS)
   c. Benign and malignant lesions such as polyps
   d. Gallstones
2. Bile duct abnormalities reveal:
   a. Dilation of ducts
   b. Duct obstruction by calculi, tumor, or parasites
   c. Congenital abnormalities such as choledochal cysts
3. Adjacent liver pathologies may include:
   a. Parenchymal disease such as cirrhosis
b. Masses, including cysts, solid lesions, and metastatic tumors

4. If combined with Doppler evaluation, portal hypertension and hepatofugal (portal blood flow away from the liver) flow can be detected. Posttransplantation stenoses or flow variances can be monitored.

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**Clinical Alert**

Results of sonograms alone cannot differentiate cancers from benign processes.

**Interfering Factors**

1. Intestinal gas overlying the area of interest interferes with sonographic visualization.
2. Barium from recent radiographic studies compromises the study.
3. Obesity adversely affects tissue visualization.

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose, benefits, and procedure of the test.
2. Instruct the patient to remain NPO at least 8 hours before the examination to fully dilate the gallbladder and to improve anatomic visualization. Some laboratories prefer that the last meal before the study contain low quantities of fat.
3. Assure the patient that there is no pain involved. However, the patient may feel uncomfortable lying quietly for a long period.
4. Explain that a liberal coating of coupling agent must be applied to the skin so that there is no air between the skin and the transducer and to allow for easy movement of the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
5. Explain that the patient will be instructed to control breathing patterns while the images are being made.
6. See Chapter 1 guidelines for **safe, effective, informed pretest care**.

**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel appropriately about further testing (biopsy).
2. Follow Chapter 1 guidelines for **safe, effective, informed posttest care**.

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**Clinical Alert**

1. Scans cannot be done over open wounds or through dressings.
2. This examination must be performed before radiographic studies involving barium. If such scheduling is not possible, at least 24 hours must elapse between the barium procedure and the sonogram.
3. The gallbladder's ability to contract may be tested by administering a fatty substance and rescanning.

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**Abdominal Aorta Sonogram**

The abdominal aorta sonogram is a noninvasive examination used to evaluate the abdominal aorta and its major tributaries for structural abnormalities such as aneurysms and the presence of thrombus. Many laboratories include Doppler evaluations to characterize blood flow through vessels. Typically, the path of the abdominal aorta is traced from its most proximal portion to the region of its bifurcation into the iliac arteries.

**Reference Values**

*Normal* Normal pattern image showing regular contour and diameter of the aorta. The walls strongly reflecting echoes, whereas the blood-filled lumen is echo free.

**Procedure**

1. Ask the patient to lie quietly on an examination table. Scans are generally performed with the patient in the supine and decubitus positions.
2. Cover the skin with a layer of coupling gel, oil, or lotion.
3. Ask the patient to regulate breathing patterns as instructed during the examination.
4. Tell the patient that the total examination time is about 30 minutes.
5. See Chapter 1 guidelines for **intra-test care**.

**Clinical Implications**

1. The typical abnormal pattern reveals aortic aneurysms with or without thrombus. Intimal dissections and leaks also may be detected.
2. Barium from recent radiographic studies compromises the study.
3. Obesity adversely affects tissue visualization.

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose, benefits, and procedure of the test.
2. Instruct the patient to remain NPO for at least 8 hours before the examination to fully dilate the gallbladder and to
improve anatomic visualization of all structures.

3. Assure the patient that there is no pain involved. However, the patient may feel uncomfortable lying quietly for a long period.

4. Explain that a liberal coating of coupling agent must be applied to the skin so that there is no air between the skin and the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.

5. Explain that the patient will be instructed to control breathing patterns while the images are being made.

6. See Chapter 1 guidelines for safe, effective, informed pretest care.

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### Clinical Alert

1. Scans cannot be done over open wounds or through dressings.
2. This examination must be performed before radiographic studies involving barium. If such scheduling is not possible, at least 24 hours must elapse between the barium procedure and the sonogram.

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### Posttest Patient Aftercare

1. The patient may resume normal diet and fluids.
2. Interpret test outcomes and counsel appropriately about possible further testing (arteriogram) and treatment (surgery).
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

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### Abdominal Ultrasound

This noninvasive procedure visualizes all solid organs of the upper abdomen, including the liver, gallbladder, bile ducts, pancreas, kidneys, spleen, and large abdominal blood vessels. Some diagnostic laboratories may perform organ-specific studies, such as renal or hepatobiliary ultrasound, together with abdominal ultrasound. This study is valuable in detecting a variety of pathologies, including fluid collections, masses, infections, and obstructions.

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### Reference Values

**Normal** Normal size, position, and appearance of the liver, gallbladder, bile ducts, pancreas, kidneys, spleen, as well as the abdominal aorta and inferior vena cava and their major tributaries

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### Procedure

1. Ask the patient to lie quietly on the examination table. Scans are generally performed with the patient in the supine and decubitus positions.
2. Cover the skin with a layer of coupling gel, oil, or lotion.
3. Explain that the patient will be asked to regulate breathing patterns as instructed during the examination.
4. Tell the patient that total examination time is about 30 to 60 minutes.
5. See Chapter 1 guidelines for intratest care.

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### Clinical Implications

1. **Liver** abnormalities reveal:
   a. Cysts, abscesses, tumors, and metastases
   b. Parenchymal disease (eg, cirrhosis)
   c. Variations in portal venous flow
   d. Hepatic arterial and venous flow patterns
2. **Gallbladder and bile duct** abnormalities reveal:
   a. Duct dilation or obstruction
   b. Gallstones
   c. Cholecystitis
   d. Tumors
3. **Pancreas** abnormalities reveal:
   a. Pancreatitis
   b. Pseudocyst
   c. Cysts and tumors, including adenocarcinoma
4. **Kidney** abnormalities reveal:
   a. Hydronephrosis
   b. Cysts, tumors, abscesses
   c. Abnormal size, number, location of kidneys
   d. Calculi
   e. Perirenal fluid collections
   f. Patency and flow through renal artery; patency of renal vein
5. **Adrenal** abnormalities reveal:
   a. Pheochromocytoma
   b. Adrenal hemorrhage
   c. Metastases
6. **Spleen** abnormalities reveal:
   a. Splenomegaly
   b. Evidence of lymphatic disease, lymph node enlargement
   c. Evidence of trauma
7. **Vascular** abnormalities in the upper abdomen reveal:
   a. Aneurysm
   b. Thrombi
c. Abnormal blood flow patterns

8. Miscellaneous pathologies include:
   a. Ascites
   b. Mesenteric or omental cysts or tumors
   c. Congenital absence or malplacement of organs
   d. Retroperitoneal tumors
   e. Hematomas

Clinical Alert
The results of sonograms alone cannot differentiate malignant from benign conditions.

Interfering Factors
1. Intestinal gas overlying the area of interest interferes with sonographic visualization.
2. Barium from recent radiology studies compromises the study.
3. Obesity adversely affects tissue visualization.

Interventions
Pretest Patient Preparation
1. Explain test purpose, benefits, and procedure.
2. Instruct patient to remain NPO for a minimum of 8 hours before the examination to fully dilate the gallbladder and to improve anatomic visualization of all structures. Some laboratories prefer that the last meal before the study contain low quantities of fat.
3. Assure the patient that there is no pain involved. However, the patient may feel uncomfortable lying quietly for a long period.
4. Explain that a liberal coating of coupling agent must be applied to the skin so that there is no air between the skin and the transducer to allow for easy movement of the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
5. Explain that the patient will be instructed to control breathing patterns while the images are being made.
6. See Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert
1. Scans cannot be done over open wounds or through dressings.
2. This examination must be performed before radiographic studies involving barium. If such scheduling is not possible, at least 24 hours must elapse between the barium procedure and the sonogram.

Posttest Patient Aftercare
1. Normal diet and fluids are resumed.
2. Interpret test outcomes and counsel appropriately.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

OTHER BODY STRUCTURE SONOGRAMS

Breast Sonogram (Sonomammography)

Ultrasound mammography is useful for differentiating cystic, solid, and complex lesions; in the diagnosis of disease in women with very dense breasts; and in the follow-up care of women with fibrocystic breast disease. It is recommended as the initial method of examination in a young woman with palpable mass and in a pregnant woman with a newly palpable mass. The pregnant patient presents a dilemma because malignancies in pregnancy grow rapidly and the increased glandular tissue causes difficulties in mammography. Ultrasound may be used to evaluate women who have silicone prostheses in their breasts. The prosthesis is readily penetrated by the ultrasound beam, and tissues behind the prosthesis can be examined. Such prostheses are known to obscure masses on physical examination; they also absorb x-ray beams, obscuring portions of the breast parenchyma.

Breast sonography is a valuable guide during breast biopsies and needle localization procedures. Although not optimal, sonographic visualization of the breast is an alternative for women who refuse to have a radiographic mammogram and for those who should not be exposed to radiation.

Reference Values
Normal Symmetric echo pattern in both breasts, including subcutaneous, mammary, and retromammary layers

Procedure
1. Ask the patient to lie on an examination table.
2. Apply a coupling medium, usually a gel, to the exposed breast to promote the transmission of sound.
3. Move a transducer slowly across the breast. In most laboratories, a handheld transducer is used, whereas in some, an automated breast scanner is used. The automated examination requires the patient to assume a position with the breast immersed in a tank of water. The tank contains transducers that are moved by remote control to image the breast.
4. Tell patient that the total examining time is 15 minutes.
Clinical Implications Unusual and distinctive echo patterns may indicate the presence of:

1. Cysts
2. Benign solid growths
3. Malignant tumors
4. Tumor metastasis to muscles and lymph nodes
5. Ductal ectasia
6. Enlarged lymph nodes

Interventions

Pretest Patient Preparation

1. Explain the purpose and procedure of the examination. There is no discomfort involved. Many diagnostic departments show the patient a videotape that explains the test.
2. Ask that the patient wear a 2-piece outfit on the day of examination because the garments on the torso are removed before the examination.
3. Explain that a liberal coating of a coupling agent must be applied to the skin so that there is no air between the skin and the transducer and to allow for easy movement of the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert

1. If the breast sonogram is to be performed on the same day as a radiographic mammogram, advise the patient not to apply any powders, lotions, or other cosmetics to the upper body on the day of the examination.
2. If the breast sonogram is to be used for guidance during a biopsy, make certain that a signed informed consent is secured.
3. Sonomammography is not an appropriate method for visualizing microcalcifications.

Posttest Patient Aftercare

1. The breasts are cleaned and dried, and the patient is advised to contact her referring clinician for outcomes.
2. Answer the patient's questions regarding procedures and explain need for possible further testing (biopsy) and treatment: medical, diet elimination of caffeine, or surgical (mastectomy or lumpectomy).
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Prostate Sonogram (Transurethral Ultrasound [TRUS])

The prostate sonogram is used to visualize the prostate gland, typically in response to an elevated concentration of prostate-specific antigen (PSA) on a blood test or as a complement to a digital rectal examination. Ultrasound of the prostate is also used as a guidance mechanism for biopsy procedures and to assist in placement of radiation “seeds.” Carcinoma of the prostate is the second most common cause of cancer-related death in American men.

The patient typically is instructed to prepare by administering a Fleet enema before the procedure. The patient usually is examined with the use of a small endorectal transducer that is inserted while the patient is in the left lateral decubitus position with the knees flexed toward the chest. Multiple images of the prostate, rectal walls, prostate urethra, and ejaculatory ducts are taken. Prostatic volumes are calculated from two-dimensional (2-D) measurements. Doppler evaluation is used to assess blood flow through the prostate or any mass that might be detected.

Reference Values

Normal Normal size, volume, shape, location, and echo texture of prostate and adjacent structures

Procedure

1. Ask the patient to void and to remove clothing from the waist down.
2. Position the patient on an examination table in the left lateral decubitus position, with his knees flexed toward the chest. The patient is draped.
3. Perform a digital rectal examination before inserting the rectal transducer.
4. Carefully insert a slim endorectal transducer, lubricated and sheathed with a condom, a few centimeters into the rectum.
5. Perform scans by using a slight rotation of the probe handle. Total examination time is about 15 to 20 minutes.
6. See Chapter 1 guidelines for intratest care.

Clinical Implications Abnormalities that may be detected include:

1. Prostatic enlargement—increased volume measurements may indicate:
   a. Benign prostatic hypertrophy (BPH)
   b. Space-occupying lesion (tumor, cyst, abscess)
2. Prostatic calcifications
3. Prostatitis
4. Prostate cancer, classically seen as a low-level echo structure within the outer gland (peripheral and central zones)

Interfering Factors Excess fecal matter in the rectum compromises the study.
**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose and procedure of the test.
2. Assure the patient that no pain is involved. However, a sensation of fullness within the rectum is to be expected. Because the transducer is typically draped within a condom, check for latex sensitivities.
3. Many laboratories require administration of a Fleet enema about 1 hour before the study.
4. Advise the patient to empty the bladder immediately before the study.
5. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Clinical Alert**

1. If the prostate examination is performed in conjunction with a prostatic biopsy, be certain to obtain a signed informed consent.
2. If the patient is latex sensitive, contact the laboratory.

**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel about any identified prostatic abnormalities and need for possible further testing (tissue biopsy with cytologic or histologic exam) and treatment.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Scrotal Sonogram**

This noninvasive ultrasound study is useful in diagnosing testicular masses, varicoceles, hydroceles, spermatoceles, and diffuse processes. Doppler ultrasound or color-flow Doppler evaluation is helpful in demonstrating the presence of torsion of the testes. Testicular ultrasound is used to evaluate scrotal pain and to demonstrate the scope of scrotal trauma.

**Reference Values**

*Normal*
Normal scrotal structures, testicles, epididymis, and spermatic cord
Normal scrotal blood flow

**Procedure**

1. Ask the patient to lie on his back. The penis is gently retracted, and the scrotum is supported on a rolled towel.
2. Apply an acoustic gel to the skin, then pass the transducer repeatedly over the scrotum. Sonographic images are generated.
3. Tell patient that total examination time is about 30 minutes.
4. Use color Doppler studies to assess presence, absence (as in torsion), or increase (as in infection and certain neoplasms) of blood flow in the testicle.
5. See Chapter 1 guidelines for intratest care.

**Clinical Implications**
Abnormal results are associated with:

1. Abscess
2. Infarcted testes (torsion)
3. Tumor (primary and metastatic)
4. Hydrocele
5. Spermatocele
6. Adherent scrotal hernia
7. Cryptorchism
8. Epididymitis (chronic or acute), orchitis
9. Hematoma (associated with trauma)
10. Tuberculosis infection (associated with AIDS)
11. Testicular microlithiasis

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose, benefits, and procedure of the test.
2. Assure the patient that there is no pain involved.
3. Explain that a liberal coating of coupling media must be applied to the scrotum. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
4. See Chapter 1 guidelines regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel appropriately about further testing (nuclear scan), treatment, medications for infection (TB, HIV) and/or surgery.
2. Follow Chapter 1 guidelines regarding safe, effective, informed posttest care.

**Eye and Orbit Sonograms**

Ultrasound can be used to describe both normal and abnormal tissues of the eye when no alternative visualization is possible because of opacities caused by inflammation or hemorrhage. This information is valuable in the management of
eyes for keratoprosthesis. Orbital lesions can be detected and distinguished from inflammatory and congestive causes of exophthalmos with a high degree of reliability. An extensive preoperative evaluation before vitrectomy or surgery for vitreous hemorrhages is also done. In this case, the vitreous cavity is examined to rule out retinal and choroidal detachments and to detect and localize vitreoretinal adhesions and intraocular foreign bodies. Also, patients who are to have intraocular lens implants after removal of cataracts must be measured for the exact length of the eye (within 0.1 mm).

Reference Values

Normal Pattern image indicating normal soft tissue of eye and retrobulbar orbital areas, retina, choroid, and orbital fat

Procedure

1. Place a small, very-high-frequency transducer on the eye directly, or position it over a water standoff pad placed onto the eye surface. Multiple images and measurements are taken.
2. Anesthetize the eye area by instilling eye drops.
3. Ask the patient to fix the gaze and hold very still.
4. Place a probe gently on the corneal surface.
5. Tell the patient that if a lesion in the eye is detected, as much as 30 minutes may be required to differentiate the pathologic process accurately. Otherwise, orbital examinations can be done in 8 to 10 minutes.
6. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. Abnormal patterns are seen in:
   a. Alkali burns with corneal flattening and loss of anterior chamber
   b. Detached retina
   c. Keratoprosthesis
   d. Extraocular thickening in thyroid eye disease
   e. Pupillary membranes
   f. Cyclotic membranes
   g. Vitreous opacities
   h. Orbital mass lesions
   i. Inflammatory conditions
   j. Vascular malformations
   k. Foreign bodies
2. Abnormal patterns are also seen in tumors of various types based on specific ultrasonic patterns:
   a. Solid tumors (eg, meningioma, glioma, neurofibroma)
   b. Cystic tumors (eg, mucocele, dermoid, cavernous hemangioma)
   c. Angiomatous tumors (eg, diffuse hemangioma)
   d. Lymphangioma
   e. Infiltrative tumors (eg, metastatic lymphoma, pseudotumor)

Interfering Factors If at some time the vitreous humor in a particular patient has been replaced by a gas, no result can be obtained.

Interventions

Pretest Patient Preparation

1. Explain the purpose, benefits, and procedure of the test.
2. Instill topical anesthetic drops into the eyes before the examination is performed; this usually is done in the examining department.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Instruct the patient to refrain from rubbing the eyes until the effects of anesthetic have disappeared. This type of friction could cause corneal abrasions.
2. Advise the patient that minor discomfort and blurred vision may be experienced for a short time. Counsel regarding possible further testing and/or treatment for infection (medical or surgical for detached retina).
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

When a ruptured globe is suspected, ophthalmic ultrasound should not be performed. Excessive pressure applied to the globe may cause expulsion of the contents and increases the risk for introduction of bacteria.

Thyroid Sonogram (Neck Ultrasound)

This ultrasound study is used to evaluate a neck mass or to determine the size of the thyroid and reveal the depth and dimension of thyroid goiters and nodules. The response of a mass in the thyroid to suppressive therapy can be monitored by successive examinations. Theoretically, this technique offers the possibility of a good estimation of thyroid weight—information that is important in radioiodine therapy for Graves' disease.

The examination is easy to do, is often done before surgery, and gives 85% accuracy. Often, these studies are done in conjunction with radioactive iodine uptake tests. With pregnant patients, ultrasound studies are the method of choice
because radioactive iodine is harmful to the developing fetus.

Reference Values

**Normal**
Normal, homogenous pattern of thyroid and adjacent structures, including strap muscles and blood vessels

Procedure

1. Have the patient lie on the back on the examining table, with the neck hyperextended.
2. Place a pillow under the shoulders for comfort and to bring the transducer into better contact with the thyroid.
3. Apply an acoustic couplant (gel, lotion, or oil) to the patient's neck. This affords good contact between the transducer and the patient's skin and allows the transducer to be moved easily across the neck's surface. An alternate procedure involves separation of the neck surface from the transducer by a gel-filled pad that permits proper transmission of the ultrasound waves through the thyroid.
4. Tell the patient that the examination time is about 30 minutes.
5. See Chapter 1 guidelines for *intratest* care.

Clinical Implications

1. An abnormal pattern may consist of a cystic, complex, or solid echo pattern.
2. Solitary "cold" nodules identified on radioisotope scans may appear as echo-free cysts on ultrasound. Most often, cysts are benign. Solid-appearing lesions may represent benign adenomas or malignant tumors. A biopsy is the only definitive method to determine the nature of such tumors.
3. Overall gland enlargement is indicative of goiter or thyroiditis.
4. Sonographic studies of the neck may also reveal parathyroid lesions or evidence of changed lymph nodes.
5. Certain congenital deformities related to the embryologic development of neck structures may be detected, most commonly thyroglossal duct cyst, brachial cleft cyst, or cystic hygroma.

Interfering Factors

1. Nodules < 1 cm in diameter may escape detection.
2. Cysts not originating in the thyroid may show the same ultrasound characteristics as thyroid cysts.
3. Lesions > 4 cm in diameter frequently contain areas of cystic or hemorrhagic degeneration and give a mixed echogram that is difficult to correlate with specific disease.

Interventions

**Pretest Patient Preparation**

1. Explain the purpose and procedure of the test.
2. Assure the patient that there is no pain involved. However, the patient may feel uncomfortable maintaining the neck position during the examination.
3. Explain that a liberal coating of coupling agent must be applied to the skin so that there is no air between the skin and the transducer and to allow for easy movement of the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
4. Advise the patient to refrain from wearing necklaces to the laboratory.
5. See Chapter 1 guidelines for *safe, effective, informed pretest care*.

**Clinical Alert**

Thyroid or neck biopsies are often performed with ultrasound guidance. If a biopsy is performed, a witnessed, informed consent must be signed in advance by the patient.

**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel about follow-up testing (thyroid nuclear scans) or treatment for thyroid (surgical removal) or neck abnormalities.
2. Follow Chapter 1 guidelines regarding *safe, effective, informed posttest care*.

**VASCULAR ULTRASOUND STUDIES (DUPLEX SCANS)**

**Overview of Duplex Scans**

The combination of anatomic imaging of blood vessels and hemodynamic information provided by Doppler ultrasound results in duplex scans. These noninvasive studies can be performed on literally any area of human anatomy. Blood velocity is detected by positioning the Doppler sample gate within the lumen of the desired vessel. The resultant *spectral trace* ([Fig. 13.1](#)) also provides information as to the direction, phase, pulsatile rhythm, and resistivity of flow. *Antegrade* flow is demonstrated above the baseline. *Retrograde* flow (ie, flow in the direction opposite then expected) is demonstrated by a spectral trace below the baseline. Flow that is antegrade through all phases (systole as well as diastole) demonstrates a *low-resistive* profile, which is normally associated with many visceral blood vessels (eg, renal artery, internal carotid artery). High-resistance, or *triphasic*, flow is typically associated with peripheral arteries (eg, femoral artery, brachial artery) and shows a forward-backward-forward pattern in each cycle. *Spectral broadening* occurs when the sample contains blood cells moving at many velocities; this is generally associated with a flow disturbance. Mathematical ratios that contrast peak or mean velocities at various stages in the cycle can give further clues to the integrity of the vascular system examined. Color Doppler ultrasound generally is used to code flow velocities and direction with color and can readily differentiate the patency of vessels. B-flow is a technique that images the blood itself...
and has an enhanced ability to display plaque margins.

**FIGURE 13.1** Blood velocity tracings show direction, phasicity, pulsatility, and resistivity of flow.

### Cerebrovascular Ultrasound (Carotid and Vertebral Arteries) Duplex Scans

Carotid duplex scans examine the major extracranial arteries supplying the brain to gain information about cerebrovascular blood flow. Carotid scans are used in the evaluation of ischemia, headache, dizziness, hemiparesis, paresthesias, and speech and visual disturbances. Testing is commonly performed before major cardiovascular surgery and as a follow-up to many surgeries.

#### Reference Values

**Normal**

- Normal vascular anatomy and course of common carotid artery, internal and external carotids, and vertebral arteries
- No evidence of stenosis or occlusion; normal flow patterns

#### Procedure

1. Ask the patient to lie on the examining table with the neck slightly extended. The head typically is turned away from the side being examined.
2. Apply an acoustic coupling gel to the neck area to enhance the transmission of sound. During Doppler evaluation, an audible signal, representing blood flow, can be heard.
3. Move a handheld transducer gently up and down the neck while images of appropriate blood vessels are made. Examine both sides of the neck.
4. Tell patient that the examination time is 30 to 60 minutes.
5. See Chapter 1 guidelines for *inratest* care.

#### Clinical Implications

Abnormal images and Doppler signals may provide evidence of the following:

1. Plaque
2. Stenosis
3. Occlusion
4. Dissection
5. Aneurysm
6. Carotid body tumor
7. Arteritis

#### Interfering Factors

1. Severe obesity and patient movement compromise examination quality.
2. Cardiac arrhythmias and disease may cause changes in hemodynamic patterns.

#### Interventions

**Pretest Patient Preparation**

1. Explain the test purpose, benefits, and procedure. Patient should refrain from smoking or consuming caffeine for at least 2 hours before the study. Assure the patient that no radiation is employed, typically no contrast medium is injected, and no pain is involved. Some slight discomfort may be experienced from lying with head extended.
2. Advise the patient that a liberal coating of coupling gel must be applied to the skin to promote sound transmission. A sensation of warmth or wetness may be felt during application. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination. Necklaces and earrings must be removed before the study.
3. See Chapter 1 guidelines for *safe, effective, informed pretest care*.

**Posttest Patient Aftercare**

1. Remind the patient to remove any residual gel from the skin.
2. Interpret test outcomes, provide support, and counsel appropriately should an abnormality be detected. Monitor and counsel for arterial disease and possible further testing (angiogram) and treatment (surgery).
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Peripheral Arterial Doppler Studies; Lower Extremity Arterial (LEA) and Upper Extremity Arterial (UEA) Duplex Scans

Peripheral arterial studies visualize and document the arterial blood flow in the extremities. Duplex ultrasound scans can determine the presence, amount, and location of plaques and are helpful in assessing the cause of claudication. Graft patency and condition may also be evaluated. Ultrasound analysis is used to evaluate the site of a prior surgical or percutaneous intervention. Some institutions also incorporate segmental blood pressure readings into these examinations. Flow characteristics of upper versus lower extremities can be contrasted by calculating a mathematical ratio between pressures (see Ankle-Brachial Index and Segmental Pressures, page 892).

Reference Values
Normal Normal arterial anatomy of the extremity Normal triphasic blood flow and flow velocities No evidence of plaques or other pathologic processes

Procedure
1. Ask the patient to lie on the examining table with the leg or arm turned out slightly and the knee or elbow partially bent.
2. Apply an acoustic coupling gel to the leg from groin down or to the arm from shoulder down to enhance the transmission of sound. During Doppler evaluation, an audible signal, representing blood flow, can be heard.
3. Move a handheld transducer gently up and down the limb while images of appropriate blood vessels are made. Examine both sides.
4. Tell the patient that the examination time is about 60 minutes.
5. For all procedures, see Chapter 1 guidelines for intratest.

Clinical Implications Abnormal tracings (see Fig. 13-1) and Doppler signals may provide evidence of the following:
1. Plaque or calcification (particularly in the diabetic patient)
2. Stenosis (hemodynamically significant lesions produce > 50% stenosis)
3. Occlusion
4. Arteritis
5. Aneurysm
6. Pseudoaneurysm
7. Graft diameter reduction
8. Abnormal communication between artery and vein

Interfering Factors
1. Severe obesity compromises examination quality.
2. Cardiac arrhythmias and disease may cause changes in hemodynamic patterns.

Interventions
Pretest Patient Preparation
1. Explain the test purpose, benefits, and procedure. Instruct the patient to refrain from smoking or consuming caffeine for at least 2 hours before the test. Assure the patient that no radiation is employed, typically no contrast medium is injected, and no pain is involved. Some slight discomfort may be experienced from lying with the extremity extended or if segmental blood pressures are taken.
2. Advise the patient that a liberal coating of coupling gel must be applied to the skin to promote sound transmission. A sensation of warmth or wetness may be felt during application. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Remind the patient to remove any residual gel from the skin.
2. Interpret test outcomes, provide support, and counsel appropriately should an abnormality be detected. Monitor and counsel for arterial disease and possible further testing (arteriogram or venogram) and treatment (surgery).
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Ankle-Brachial Index (ABI) and Segmental Pressures

In some laboratories, as an adjunct to duplex scanning, blood pressures throughout the extremities are measured and contrasted. In the typical four-cuff technique, pneumatic cuffs are applied to the upper thigh, the lower thigh, the upper calf, and the area just above the ankle. Additionally, cuffs are applied to the upper arms to determine brachial pressures. Segmental pressures provide physiologic information that can confirm a vascular cause for ischemic rest pain and claudication. The ankle-brachial index (ABI) is calculated by dividing the ankle pressure (in mm Hg) by the brachial pressure. Many laboratories perform a resting ABI followed by an exercise ABI.

Reference Values
Normal ABI > 1.0 when a normal multiphasic waveform is present A difference of 20 mm Hg (or 20 torr) between the right and left brachial pressures may indicate proximal arterial obstruction on the side with reduced pressure. The gradual pressure drop, as measured from upper thigh or arm to ankle or wrist, should not exceed 20 mm Hg (or 20 torr) between any two segments
Procedure

1. Ask the patient to lie on the table with the extremity extended.
2. Place pneumatic cuffs (usually four) at intervals along the extremity.
3. Place a flow-sensing device (often a continuous-wave Doppler device) distal to a cuff. Inflate the cuff (this is often done automatically) to suprasystolic values and then slowly deflate until flow resumes. Record the pressure at which flow resumes.
4. Repeat this technique, distal to each cuff, until the entire extremity has been evaluated. Measure brachial pressures as well.
5. Examine both extremities.
6. Tell the patient that the total examination time (for pressures only) is generally < 15 minutes. If an exercise/stress ABI is ordered, the at-rest study will be followed by exercise. After walking for 5 minutes on a treadmill, the ABI procedure is repeated.
7. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. Asymmetry in brachial pressure > 10 mm Hg (>10 torr) is suspicious for arterial disease.
2. ABI < 1.0 is suspicious for disease. The lower the numeric value for this index, the more severe the disease may be (eg, ABI < 0, associated with impending tissue loss).
3. Generally speaking, pressure gradients between successive segments on the same extremity should vary by < 20 mm Hg (<20 torr). Variations that exceed this value suggest significant disease (occlusion or stenosis).
4. A difference of > 20 mm Hg (>20 torr) between similar segments on opposite sides may suggest obstructive vascular disease.

Clinical Alert

1. Segmental pressures are a screening tool that cannot distinguish stenosis from total occlusion and cannot be specific in determining the exact location of disease.
2. Vessel calcifications (commonly seen in the diabetic patient) can falsely elevate systolic pressures.

Interfering Factors

1. Severe obesity compromises examination quality.
2. Cardiac arrhythmias and disease may cause changes in hemodynamic patterns.

Interventions

Pretest Patient Preparation

1. Explain the test purpose, benefits, and procedure. Instruct the patient to refrain from smoking or consuming caffeine for at least 2 hours before the study. Assure patient that no radiation is employed, typically no contrast medium is injected, and no pain is involved. Some discomfort may be experienced from lying with the extremity extended or when pneumatic cuffs are inflated.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes, provide support, and counsel appropriately should an abnormality be detected. Monitor and counsel for arterial disease and explain need for possible further testing (arteriogram) and treatment (medical or surgical).
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Peripheral Venous Doppler Studies; Lower Extremity Venous (LEV) and Upper Extremity Venous (UEV) Duplex Scans

This procedure examines venous blood flow in the selected extremity (upper or lower). It is most commonly used to assess deep venous thrombosis and can also be used to "map" veins to be harvested and used for grafts. Peripheral ultrasound is also used to locate veins for venous access and to assess dialysis access grafts. This examination has replaced contrast venography in many institutions.

Reference Values

Normal Duplex Scan
Normal venous anatomy of the extremity Spontaneous phasic flow pattern (rises and falls with respiration) Normal venous augmentation (exhibits increased flow proximal to the site of venous compression) Competent, intact valves, with no evidence of thrombi

Procedure

1. Ask the patient to lie on the examining table with the leg or arm turned out slightly and the knee or elbow partially bent.
2. Apply an acoustic coupling gel to the leg from the groin down or to the arm from the shoulder area down to enhance the transmission of sound. During Doppler evaluation, an audible signal, representing blood flow, can be heard.
3. Move a handheld transducer gently up and down the limb while images of appropriate blood vessels are made. At intervals, apply gentle compression to the vessel. Examine both sides.
4. Tell the patient that the examination time is about 30 minutes.
5. See Chapter 1 guidelines for intratest care.
Clinical Implications Abnormal images and Doppler signals may provide evidence of the following:

1. Acute or chronic deep venous thrombosis
2. Occlusive venous disease
3. Valvular incompetence
4. Clotted grafts

Interfering Factors

1. Severe obesity compromises examination quality.
2. Cardiac arrhythmias and disease may cause changes in hemodynamic patterns.

Interventions

Pretest Patient Preparation

1. Explain the test purpose, benefits, and procedure. Instruct the patient to refrain from smoking for at least 2 hours before the study. Assure the patient that no radiation is employed, typically no contrast medium is injected, and no pain is involved. Some slight discomfort may be experienced from lying with the extremity extended or when compression is applied.
2. Advise the patient that a liberal coating of coupling gel must be applied to the skin to promote sound transmission. A sensation of warmth or wetness may be felt during application. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Remind the patient to remove any residual gel from the skin.
2. Interpret test outcomes, provide support, and counsel appropriately should an abnormality be detected. Monitor and counsel for venous disease and need for possible further testing and/or treatment (medical, drugs, or surgical).
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

HEART ULTRASOUND STUDIES

Heart Sonogram (Echocardiogram; Doppler Echocardiography)

This noninvasive technique for examining the heart can provide information about its position and size, movements of the valves and chamber, and velocity of blood flow. Echoes from pulsed high-frequency sound waves are used to locate and study the movements and dimensions of cardiac structures. Because the heart is a blood-filled organ, sound can be transmitted through it readily to the opposite wall and to the heart-lung interface. This test is commonly used to determine biologic and prosthetic valve dysfunction, to evaluate a pericardial effusion, to evaluate the velocity and direction of blood flow, to furnish direction for further diagnostic study, and to monitor cardiac patients over an extended period. Echocardiography is also used to monitor heart failure patients relying on a left ventricular assist device (LVAD). One of the advantages of this diagnostic technique is that it can be performed at the bedside with mobile equipment or can be done in the laboratory.

The various modes of echocardiography are capable of providing a great range of information concerning cardiac structure and function. The following are common types of echocardiograms:

Two-dimensional (2-D): used to produce gray-scale, cross-sectional images of the heart's anatomy

M-mode: used to generate depictions of rapidly moving structures such as valves and for standardized dimensional measurements

Continuous-wave Doppler and pulsed-wave Doppler: used to determine velocity of blood flow

Color 2-D: used for identifying areas of disturbed or eccentric blood flow

Color M-mode: used for evaluating movement of cardiac structures

Specialized types of echocardiography include:

Stress echocardiography: used to provide information relating to the function of heart structures during high cardiac output states. A treadmill or upright bicycle may be used, or the heart can be stressed by an infusion of dobutamine.

Transesophageal echocardiography (TEE): A miniature ultrasound transducer is placed at the end of a tube inserted into the esophagus to provide a closer view of cardiac structures without interference from superficial chest tissues (see page 897).

Fetal echocardiography: performed through the pregnant woman's abdomen when there is a question of congenital cardiac defect (see page 870).

Contrast echocardiography: A liquid containing nontoxic microbubbles is injected into a vein to opacify cardiac structures.
These special techniques may require a signed, informed consent before performance and involve more complicated procedures. Check with the individual laboratory for specific guides and protocols.

Reference Values

Normal position, size, and movement of heart valves and chamber walls as visualized in 2-D, M-mode, and Doppler mode Color M-mode and color Doppler assessments of heart structures within normal limits

Procedures

1. Ensure that a specific diagnosis accompanies the request for the test (eg, “rule out pericardial effusion,” “determine severity of mitral stenosis”). If a stress echocardiogram is ordered, the patient’s ability to perform exercise must be indicated.
2. Ask the patient to lie on the examining table in a slight side-lying position.
3. Apply an acoustic gel to the skin surface over the chest to permit maximum penetration of the ultrasound beam. Hold the transducer over various regions of the chest and upper abdomen to obtain the appropriate views of the heart.
4. Tell that patient that there should be no pain or discomfort involved. Leads may be attached for a simultaneous electrocardiogram reading during the ultrasound procedure.
5. Tell the patient that the examination time is 30 to 45 minutes.
6. See Chapter 1 guidelines for intratest care.

Clinical Implications Abnormal values help to diagnose:

1. Acquired cardiac disease
   a. Valvular disease, stenosis insufficiency, prolapse, and regurgitation
   b. Cardiomyopathies
   c. Evidence of coronary artery disease
   d. Pericardial disease, including effusion, tamponade, and pericarditis
   e. Endocarditis
   f. Cardiac neoplasm
   g. Intracardiac thrombi
2. Prosthetic valve function
3. Congenital heart disease

Interfering Factors

1. Dysrhythmias interfere with the test.
2. Hyperinflation of the lungs with mechanical ventilation, especially with positive end-expiratory pressure (PEEP) > 10 cm H₂O, precludes adequate ultrasound imaging of the heart.
3. False-negative and false-positive diagnoses have been identified (especially in M-mode echocardiograms), including diagnoses of pleural effusion, dilated descending aorta, pericardial fat pad, tumors encasing the heart, clotted blood, and loculated effusions.
4. Doppler study results can vary greatly if the transducer position does not provide satisfactory angles for the beam.

Interventions

Pretest Patient Preparation

1. Explain the purpose, benefits, and procedure of the test.
2. Assure the patient that no pain is involved. However, some discomfort may be felt from lying quietly for a long period.
3. Explain that a liberal coating of coupling agent must be applied to the skin so that there is no air between the skin and the transducer and to permit easy movement of the transducer over the skin. A sensation of warmth or wetness may be felt. Although the acoustic couplant does not stain, advise the patient not to wear good clothing for the examination.
4. See Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert

Certain specialized echocardiographic procedures, such as stress echocardiography and TEE, may require individualized patient preparation. Check with the laboratory to determine specific protocols and preparation.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately about cardiac disorders and explain need for possible further testing and/or treatment (medical, drugs, or surgical).
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Transesophageal Echocardiogram (TEE)

TEE permits optimal ultrasonic visualization of the heart when traditional transthoracic (noninvasive) echocardiography fails or proves inconclusive. A miniaturized high-frequency ultrasound transducer is mounted on an endoscope and coupled with an ultrasound instrument to display and record ultrasound images from the heart. Endoscope controls allow remote manipulation of the transducer tip. Various images of heart anatomy can be displayed by rotating the tip of the
instrument and by varying the depth of insertion into the esophagus.

Indications for TEE include the following:

1. To assess function of prosthetic valves, diagnose endocarditis, evaluate valvular regurgitation and congenital abnormalities, and examine the aorta for dissecting aneurysms
2. To monitor left ventricular wall motion intraoperatively
3. To measure ejection fraction in selected patients
4. Situations in which a transthoracic echocardiogram has not been satisfactory (eg, obesity, chest wall trauma, chronic obstructive pulmonary disease)
5. When results of traditional transthoracic echocardiography do not agree or correlate with other clinical findings

**Reference Values**

**Normal** Normal position, size, and function of heart valves and heart chambers

**Procedure**

1. Apply a topical anesthetic to the pharynx. Insert a bite block into the mouth. This reduces the risk for damage to the patient's teeth and oral structures and accidental damage to the endoscope.
2. Ask the patient to assume a left lateral decubitus position while the lubricated endoscopic instrument is inserted to a depth of 30 to 50 cm. Ask the patient to swallow to facilitate advancement of the device.
3. Manipulate the ultrasound transducer to provide a number of image planes.
4. For all procedures, see **Chapter 1** guidelines for **intranest** care.

**Procedural Alert**

A variety of medications may be used during this procedure. Generally, these drugs are intended to sedate, anesthetize, reduce secretions, and serve as contrast agents for the ultrasound. (See **Appendix A**.)

**Clinical Implications**

**Abnormal TEE findings include:**

1. Heart valve disease: stenosis, insufficiency, prolapse, and regurgitation
2. Pericardial effusion, pericarditis, tamponade
3. Congenital heart disease
4. Aortic dissection
5. Left ventricular dysfunction
6. Endocarditis
7. Intracardiac tumors or thrombi

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose, procedure, and the benefits and risks of the test.
2. The patient must remain NPO for at least 4 to 8 hours before the procedure to reduce the risk for aspiration. Pretest medication such as analgesics or sedatives may be ordered. Check with the laboratory or physician for specific instructions.
3. Obtain baselines vital signs.
4. Establish an intravenous access line to administer medications or contrast agents.
5. Remove dentures and any loose objects from patient's mouth.
6. See Chapter 1 guidelines for **safe, effective, informed pretest care**.

**Patient Posttest Aftercare**

1. Interpret test results; monitor vital signs and level of consciousness (if the patient is sedated). Ensure patent airway. Explain need for possible further testing and/or treatment: medical (drugs) or surgical (eg, cardiac catheterization).
2. Position the patient on the side, if sedated, to prevent risk for aspiration.
3. Ascertain return of swallowing, coughing, and gag reflexes before allowing patient to take oral food or fluids. Generally, the patient should remain NPO for at least 1 hour after the test.
4. Follow Chapter 1 guidelines for **safe, effective, informed posttest care**.

**Clinical Alert**

Swallowing reflexes may be diminished for several hours because of the effects of the topical anesthetic.

**BIBLIOGRAPHY**


D’Cruz IA: Echocardiographic Anatomy. Stamford, CT, Appleton & Lange, 1996


Goroll AH, May LA, Mully AG: Primary Care Medicine, Office Evaluation and Management of Adult Patient, 4th ed. Philadelphia, Lippincott Williams & Wilkins, 2001


Sanders RC: Ultrasound. Philadelphia, Lippincott Williams & Wilkins, 2001

INTERNET SITES

www.acr.org
www.auntminnie.com
www.intelihealth.com
www.webmd.com
Diagnostic evaluations of body fluid balance, electrolytes, lung ventilatory function, blood gas exchange in the lungs, oxygen tissue saturation by pulse oximetry, and acid-base balance are important determinants of normal body function (homeostasis). Homeostatic mechanisms are affected by a variety of exogenous (originating from without, eg, stress) and endogenous (originating from within, eg, immune system) factors. Evidence supports a strong association between stress and altered immune function, which can subsequently lead to abnormal pathophysiology observable by a wide array of diagnostic tests. Abnormal test outcomes in hospitalized patients, as well as when complications of treatment occur (as in kidney and respiratory diseases, diabetes, anemia), gastric fluid loss, medication diuretics, sepsis, and fever, for this reason are discussed in this chapter. Other factors that need to be assessed include respiratory rate, fluid intake, urine output, amount of water diarrhea, emesis, weight gain or loss, presence of burned or excoriated skin, food intake, and evidence of dehydration edema.

PULMONARY FUNCTION TESTS

Pulmonary Physiology

There are three aspects of pulmonary function: perfusion, diffusion, and ventilation. Perfusion relates to blood flow through pulmonary vessels; diffusion refers to movement of oxygen and carbon dioxide across alveolar capillary membranes; and ventilation relates to air exchange between alveolar spaces and the atmosphere.

During breathing, the lung-thorax system acts as a bellows to provide air to the alveoli for adequate gas exchange to take place. Like a spring or rubber band, the lung tissue also possesses the property of elasticity. When the inspiratory muscles contract, the thorax and lungs expand; when the same muscles relax and the force is removed, the thorax and lungs return to their resting position. Also, when the thorax and lungs expand, the alveolar pressure is lowered below atmospheric pressure. This permits air to flow into the trachea, bronchi, bronchioles, and alveoli. Expiration is mainly passive. It occurs because the thorax and lungs recoil to their resting position: the alveolar pressure increases above atmospheric pressure, and air flows out through the respiratory tract. The major function of the lung is to provide adequate ventilation to meet the metabolic demands of the body during rest and during exercise. The primary purpose of pulmonary blood flow is to conduct mixed venous blood through the capillaries of the alveoli so that oxygen (O₂) can be taken up by the blood and carbon dioxide (CO₂) can be removed from the blood.

Purpose of Tests

Pulmonary function tests determine the presence, nature, and extent of pulmonary dysfunction caused by obstruction, restriction, or both. When ventilation is disturbed by an increase in airway resistance, the ventilatory defect is called an obstructive ventilatory impairment. When ventilation is disturbed by a limitation in chest wall excursion, the defect is referred to as a restrictive ventilatory impairment. When ventilation is altered by both increased airway resistance and limited chest wall excursion, the defect is termed a combined or mixed defect. Table 14.1 presents the conditions that affect ventilation.

Table 14.1 Conditions That Affect Ventilation

<table>
<thead>
<tr>
<th>Examples</th>
<th>Causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>RESTRICTIVE VENTILATORY</td>
<td></td>
</tr>
</tbody>
</table>
**IMPAIREDS**

<table>
<thead>
<tr>
<th>Condition</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chest wall disease</td>
<td>Injury, kyphoscoliosis, spondylitis, muscular dystrophy, other neuromuscular diseases</td>
</tr>
<tr>
<td>Extrathoracic conditions</td>
<td>Obesity, peritonitis, ascites, pregnancy</td>
</tr>
<tr>
<td>Interstitial lung disease</td>
<td>Interstitial pneumonitis, fibrosis, pneumoconioses (eg, asbestosis, silicosis), granulomatosis, edema, sarcoidosis</td>
</tr>
<tr>
<td>Pleural disease</td>
<td>Pneumothorax, hemothorax, pleural effusion, fibrothorax</td>
</tr>
<tr>
<td>Space-occupying lesions</td>
<td>Tumors, cysts abscesses</td>
</tr>
</tbody>
</table>

**OBSTRUCTIVE VENTILATORY IMPAIRMENTS**

<table>
<thead>
<tr>
<th>Condition</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peripheral airway disease</td>
<td>Bronchitis, bronchiectasis, bronchiolitis, bronchial asthma, cystic fibrosis</td>
</tr>
<tr>
<td>Pulmonary parenchymal disease</td>
<td>Emphysema</td>
</tr>
<tr>
<td>Upper airway disease</td>
<td>Pharyngeal, tracheal or laryngeal tumors, edema, infections, foreign bodies, collapsed airway, stenosis</td>
</tr>
</tbody>
</table>

**MIXED-DEFECT VENTILATORY IMPAIRMENTS**

<table>
<thead>
<tr>
<th>Condition</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pulmonary congestion</td>
<td>Both increased airway resistance and limited expansion of chest cavity and/or chest wall; obstruction caused by bronchial edema, compression of respiratory airway owing to increased interstitial (and intravenous fluid) pressure; restriction caused by impaired elasticity, anatomic deformity (eg, kyphosis, lordosis, scoliosis)</td>
</tr>
</tbody>
</table>

*Characterized by interference with chest wall or lung movement, "stiff lung," and an actual reduction in the volume of air that can be inspired.*

†Characterized by the need for increased effort to produce airflow; respiratory muscles must work harder to overcome obstructive forces during breathing; prolonged and impaired airflow during expiration; airway resistance increases and lungs become very compliant.

‡Combined or mixed; exhibits components of both obstructive and restrictive ventilatory impairments.

Pulmonary function tests may reveal locations of abnormalities in the airways, alveoli, and pulmonary vascular bed early in the course of a disease, when the physical examination and radiographic studies still appear normal.

**Indications for Tests**

1. Early detection of pulmonary or cardiogenic pulmonary disease (see Table 14.1)
2. Differential diagnosis of dyspnea
3. Presurgical assessment (eg, ability to tolerate intraoperative anesthetics, especially during thoracic procedures)
4. Evaluation of risk factors for other diagnostic procedures
5. Detection of early respiratory failure
6. Monitoring progress of bronchopulmonary disease
7. Periodic evaluation of workers exposed to materials harmful to the respiratory system
8. Epidemiologic studies of selected populations to determine risks for or causes of pulmonary diseases
9. Workers' compensation claims
10. Monitoring after pharmacologic or surgical intervention

**Classification of Tests**

Pulmonary function tests evaluate the ventilatory system and alveoli in an indirect, overlapping way. The patient's age, height, weight, ethnicity, and gender are recorded before testing because they are the basis for calculating predicted values.

Pulmonary function tests are generally divided into three categories:

1. **Airway flow rates** typically include measurements of instantaneous or average airflow rates during a maximal forced exhalation to assess airway patency and resistance. These tests also assess responses to inhaled bronchodilator or bronchial provocations.
2. **Lung volumes and capacities** measure the various air-containing compartments of the lung to assess air-trapping (hyperinflation, overdistention) or reduction in volume. These measurements also help to differentiate obstructive from restrictive ventilatory impairments.
3. **Gas exchange (diffusion capacity)** measures the rate of gas transfer across the alveolar capillary membranes to assess the diffusion process. It can also monitor for side effects of drugs, such as bleomycin (antineoplastic) or amiodarone (antiarrhythmic), which can cause interstitial pneumonitis or pulmonary fibrosis. Diffusion capacity in the absence of lung disease (eg, anemia) can also be evaluated.
Symbols and Abbreviations

Pulmonary function studies and blood gas analyses measure quantities of gas mixtures and their components, blood and its constituents, and various factors affecting these quantities. The symbols and abbreviations given here are based on standards developed by American physiologists. Familiarity with the major and secondary symbols facilitates interpretation of any combination of these symbols (see Chart 14.1, Chart 14.2, Chart 14.3 and Chart 14.4).

Chart 14.1 Gas Volumes: Symbols and Abbreviations

Large capital letters denote primary symbols for gases:

Symbols and Abbreviations

V: Gas volume
\( \dot{V} \): Gas volume per unit time (the dot over the symbol indicates the factor per unit time, as in flow)

P: Gas pressure or partial pressure of a gas in a gas mixture (exhaled air) or in a liquid (blood)
F: Fractional concentration of a gas

Small capital letters indicate the type of gas measured in relation to respiratory tract location or function:

Symbols and Abbreviations

A: Alveolar gas
D: Dead space gas
E: Expired gas
I: Inspired gas
T: Tidal gas

Chemical symbols for gases may be placed after the small capital letters:

Symbols and Abbreviations

O2: Oxygen
CO: Carbon monoxide
CO2: Carbon dioxide
N2: Nitrogen

Combinations of Symbols

The following are some examples of the ways these symbols may be combined:

Symbols and Abbreviations

FICO2: Fractional concentration of inspired oxygen
Vt: Tidal volume
Ve: Volume of expired gas
PACO: Partial pressure of carbon dioxide in alveolar gas

Blood Gas Symbols

Large capital letters are used as primary symbols for blood determinations:

Symbols and Abbreviations

C: Concentration of a gas in blood
S: Percent saturation of hemoglobin
Q: Volume of blood
\( \dot{Q} \): Volume of blood per unit time (blood flow)

To indicate whether blood is capillary, venous, or arterial, lowercase letters are used:

Symbols and Abbreviations

v: Venous blood
a: Arterial blood
c: Capillary blood
s: Shunted blood
Chart 14.2 Combinations of Symbols and Abbreviations

Blood gas symbols may be combined in the following ways:

Symbols and Abbreviations

PO2: Oxygen tension or partial pressure of oxygen
PaO2: Arterial oxygen tension or partial pressure of oxygen in arterial blood
PAO2: Alveolar oxygen tension or partial pressure of oxygen in the alveoli
PCO2: Carbon dioxide tension or partial pressure of carbon dioxide
PaCO2: Partial pressure of carbon dioxide in arterial blood
PvCO2: Partial pressure of carbon dioxide in venous blood
pH: Hydronium ion concentration
pHa: Hydronium ion concentration in arterial blood
SO2: Oxygen saturation
SaO2: Percent saturation of oxygen in arterial blood as measured by hemoximetry (direct method)
SpO2: Percent saturation of oxygen in arterial blood as determined by pulse oximetry (indirect method)
SvO2: Percent saturation of oxygen in venous blood
TCO2: Total carbon dioxide content

Chart 14.3 Lung Volume Symbols: Pulmonary Function Terminology

This list indicates terms used in measuring lung volumes and the units that express these measurements.

FVC = Forced vital capacity: maximum amount of air that can be exhaled forcibly and completely after a maximal inspiration (liters)

FEV\_1 = Forced expiratory volume at specific time intervals (eg, 1, 2, and/or 3 seconds): volume of air expired during the first, second, third, etc., seconds of FVC maneuver (liters)

FEV\_t = Forced expiratory volume at specific time intervals (eg, 1, 2, and/or 3 seconds): volume of air expired during the first, second, third, etc., seconds of FVC maneuver (liters)

FEV\_t/FVC = Ratio of a timed forced expiratory volume to the forced vital capacity (eg, FEV\_1/FVC) (percent)

FEF\_200–1200 = Forced expiratory flow between 200 mL and 1200 mL: average flow of expired air measured after the first 200 mL and average during the next 1000 mL of the FVC maneuver (liters/second)

FEF\_25–75 = Forced expiratory flow between 25% and 75%: average flow of expired air measured between 25% and 75% of the FVC maneuver (liters/second)

PEFR = Peak expiratory flow rate: maximum flow of expired air attained during an FVC maneuver (liters/second or liters/minute)

PIFR = Peak inspiratory flow rate: maximum flow of inspired air achieved during a forced maximal inspiration (liters/second or liters/minute)

FIVC = Forced inspiratory vital capacity: maximum amount of air that can be inhaled forcibly and completely after a maximal expiration (liters)

FRC = Functional residual capacity: volume of air remaining in the lung at the end of a normal expiration (ie, end-tidal expiration) (liters)

IC = Inspiratory capacity: maximum amount of air that can be inspired from end-tidal expiration (liters)

IRV = Inspiratory reserve volume: maximum amount of air that can be inspired from end-tidal inspiration (liters)

ERV = Expiratory reserve volume: maximum amount of air that can be expired from end-tidal expiration (liters)

RV = Residual volume: volume of gas left in the lung after a maximal expiration (liters)

VC = Vital capacity: maximum volume of air that can be expired after a maximal inspiration (liters)

TLC = Total lung capacity: volume of gas contained in the lungs after a maximal inspiration (liters)

DLCO = Carbon monoxide diffusing capacity of the lung: rate of diffusion of carbon monoxide across the alveolar capillary membrane (ie, rate of gas transfer across the alveolar capillary membrane) (milliliters/minute per millimeter of mercury)

DL/V\_A = Carbon monoxide diffusing capacity per liter of alveolar volume (milliliters/minute per millimeter of mercury per liter of alveolar volume)

CV = Closing volume: volume at which the lower lung zones cease to ventilate, presumably as a result of airway closure (percent of vital capacity)

MVV = Maximum voluntary ventilation: maximum number of liters of air a patient can breathe per minute by a voluntary effort (liters/minute)

VISO\_V = Volume of isoflow: volume for which flow is the same with air and with helium during an FVC maneuver (percent)
Chart 14.4 Miscellaneous Symbols

This list shows some of the other symbols found in this chapter.

\[
f = \text{Frequency (of breathing)}
\]

\[
\text{CL} = \text{Compliance of the lung}
\]

\[
D = \text{Diffusing capacity}
\]

\[
\text{COHb} = \text{Carboxyhemoglobin}
\]

\[
\text{DLO}_2 = \text{Oxygen diffusing capacity of the lung}
\]

\[
\text{A-aDO}_2 = \text{Alveolar-to-arterial oxygen gradient}
\]

\[
\text{BSA} = \text{Body surface area (square meters)}
\]

\[
\text{H}_2\text{CO}_3 = \text{Carbonic acid}
\]

\[
\text{HCO}_3^- = \text{Bicarbonate ion}
\]

\[
\text{TGV} = \text{Thoracic gas volume (also expressed as V}_{\text{TG}}
\]

\[
\text{Raw} = \text{Airway resistance}
\]

\[
\text{G}_{\text{aw}} = \text{Airway conductance}
\]

\[
\text{sG}_{\text{aw}} = \text{Specific airway conductance}
\]

\[
\text{F-V} = \text{Flow-volume}
\]

\[
\text{V-T} = \text{Volume-time}
\]

Airway Flow Rates

Airway flow rates provide information about the severity of airway obstruction and serve as an index of dynamic function. The lung volume at which the flow rates are measured is useful for identifying a central or peripheral location of airway obstruction.

Spirometry, Forced Expiratory Maneuver Volume-Time Spirogram (V-T Tracing); Flow-Volume Spirogram (F-V Loop) Lung capacities, volumes, and flow rates are clinically measured by a mechanical device called a *spirometer*. The mechanical signal is converted to an electrical signal, which records the amounts of gas breathed in and out and produces a spirogram. Spirometers can be grouped into two major categories: (1) the mechanical or volume-displacement types (water-filled, dry-rolling seal, wedge, or bellows), and (2) the electronic or flow-sensing types (pneumotachometer or hot-wire anemometer) (Fig. 14.1). Spirometry determines the effectiveness of the various mechanical forces involved in lung and chest wall movement. The values obtained provide quantitative information about the degree of obstruction to airflow or the degree of restriction of inspired air. The forced expiratory maneuver (spirometry) is useful to quantify the extent and severity of airway obstruction. It measures the maximum amount of air that can be exhaled rapidly and forcibly, after a maximal deep inspiration. The results are a measure of airway function and the patency of the airway.

**FIGURE 14.1** MedicAir Plus electronic spirometer. (Courtesy of QRS Diagnostic, LLC, Plymouth, MN)

The forced expiratory volumes exhaled within 1, 2, or 3 seconds are sometimes referred to as *timed vital capacities* (FEV\(_1\), FEV\(_2\), and FEV\(_3\), respectively). These measurements are useful for evaluating a patient’s response to bronchodilators. Generally, if the FEV\(_1\) is < 80% (>0.80) of predicted and/or the FEF\(_{25-75}\) is < 60% (>0.60) of predicted, bronchodilators are administered with a handheld nebulizer, and the spirometry is repeated. Recently, combination bronchodilators (albuterol and ipratropium) have been introduced. Studies have shown a better bronchodilator response with the combined drugs than either alone. An increase in these values of 20% or more (>0.20) above the prebronchodilator level suggests a significant response to the bronchodilator and is consistent with a diagnosis of reversible obstructive airway disease (eg, asthma). Persons with emphysema typically do not demonstrate this type of response to bronchodilator. Measured (actual) spirometry values are compared with predicted values by means of regression equations using age, height, weight, ethnicity, and gender and are expressed as a percentage of the predicted value. Typically, a value >80% (>0.80) of predicted is considered within normal limits.

**Reference Values**

**Normal**

FVC: >80% (>0.80) of predicted value FEV\(_1\): FEV\(_1\), FEV\(_2\), FEV\(_3\), >80% (>0.80) of predicted value FEV\(_1\)/FVC: FEV\(_1\), 80%–85% (0.80–0.85) of FVC FEV\(_2\), 90%–94% (0.90–0.94) of FVC FEV\(_3\), 95%–97% (0.95–0.97) of FVC

Predicted values are based on the patient’s age, height, ethnicity, and gender.

**Procedure**

1. Have the patient either sit or stand. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Ask the patient to take a maximal inspiration and then forcibly and completely exhale into the spirometer.
3. Have the patient repeat this maneuver a minimum of three times. The two best tracings should compare within 5%
(0.05) of one another, or additional forced expiratory efforts will be needed.
4. Administer bronchodilators with a handheld nebulizer, and repeat spirometry if indicated.
5. See Chapter 1 guidelines for intratest care.

### Clinical Alert

Spirometry is a patient effort–dependent test; as such, if the patient does not provide his or her best effort, the results may be inconclusive.

### Clinical Alert

1. Before testing, assess the patient's ability to comply with breathing requirements.
2. The patient may experience lightheadedness, shortness of breath, or other slight discomforts. These symptoms are generally transitory. An appropriate rest period is usually all that is needed. If symptoms persist, testing is terminated.
3. Rarely, momentary loss of consciousness (caused by anoxia during forced expiration) may occur. Follow established protocols for testing this.
4. Assess for contraindications such as pain or altered mental status.

### Clinical Implications

1. With obstructive ventilatory impairments such as asthma, airway collapse occurs during forced expiratory effort. This leads to decreases in airway flow rates and also, in the more severe forms, to apparent loss of volumes.
2. Decreased values occur in chronic lung diseases that cause trapping of air such as emphysema, chronic bronchitis, cystic fibrosis, or asthma.
3. With restrictive ventilatory impairments, the FVC is reduced; however, flow rates can be normal or elevated.

### Interfering Factors

1. Bronchodilators (eg, albuterol) should be withheld for at least 4 hours if tolerated.
2. Respiratory infections may decrease airflow during the maneuver.

### Interventions

#### Pretest Patient Care

1. Explain the purpose and procedure of the spirometry test. Explain that the patient will be asked to perform a maximal forced inspiration in addition to the forced expirations.
2. Remind patient that a light meal may be eaten before the test. However, no caffeine should be taken before testing. Specific instructions will be given regarding the use of bronchodilators or inhaler medications before the test.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

#### Posttest Patient Aftercare

1. Evaluate for dizziness, shortness of breath, or chest discomfort. Usually these symptoms are transitory and subside after a short rest. If symptoms persist, use established follow-up protocols.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

### Peak Inspiratory Flow Rate (PIFR)

The peak inspiratory flow rate (PIFR) measures the function of the airways, identifies reduced breathing on inspiration, and is totally dependent on the effort the patient makes to inspire. The PIFR is the maximum flow of air achieved during a forced maximal inspiration.

#### Reference Values

**Normal** Approximately 300 L/min or 5 L/sec Predicted values are based on age, sex, and height.

#### Procedure

1. Have the patient either sit or stand. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Ask the patient to take a maximal inspiration, forcibly and completely exhale into the spirometer, and then inspire forcibly and completely again.
3. Have the patient repeat this maneuver a minimum of three times. Report the highest value.

### Clinical Implications

1. PIFR is reduced in neuromuscular disorders, with weakness or poor effort, and in extrathoracic airway obstruction (ie, substernal thyroid, tracheal stenosis, and laryngeal paralysis).
2. The PIFR is decreased in upper airway obstruction.

### Interfering Factors

Poor patient effort compromises the test. Inability to maintain an airtight seal around the mouthpiece

#### Interventions

**Pretest Patient Care**

1. Explain the purpose and procedure of the test. Assess the patient's ability to comply.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

#### Posttest Patient Aftercare
1. See Chapter 1 guidelines for safe, effective informed posttest care.
2. See aftercare guidelines for volume-time spirogram on page 908.

**Peak Expiratory Flow Rate (PEFR)** The peak expiratory flow rate (PEFR) measurement is used as an index of large airway function. It is the maximum flow of expired air attained during a forced expiratory maneuver.

**Reference Values**

**Normal** Approximately 450 L/min or 7.5 L/sec Predicted values are based on age, sex, and height.

**Procedure**

1. Have the patient either sit or stand. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Ask the patient to take a maximal inspiration, forcibly and completely exhale into the spirometer, and then inspire forcibly and completely again.
3. Have the patient repeat this maneuver a minimum of three times. Report the highest value.
4. Be aware that PEFR can also be measured with a handheld peak flow meter.

**Clinical Implications**

1. The PEFR usually is decreased in obstructive disease (eg, emphysema), during acute exacerbations of asthma, and in upper airway obstruction (eg, tracheal stenosis).
2. The PEFR usually is normal in restrictive lung disease but is reduced in severe restrictive situations.

**Interfering Factors** Poor patient effort compromises the test. Inability to maintain an airtight seal around the mouthpiece

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the test. Assess the patient's ability to comply.
2. See Chapter 1 guidelines for safe, effective, informed pretest care guidelines.

**Posttest Patient Aftercare**

1. Monitor patient for dizziness, lightheadedness, or chest pain following the test. Generally, these symptoms are transient and will subside quickly. If not, follow established protocols.
2. See aftercare for volume-time spiromgrams on page 908.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Lung Volumes and Capacities**

Lung volumes can be considered as basic subdivisions of the lung (not actual anatomic subdivisions). They may be subdivided as follows:

1. Total lung capacity (TLC)
2. Tidal volume (V₇)
3. Inspiratory capacity (IC)
4. Inspiratory reserve volume (IRV)
5. Residual volume (RV)
6. Functional residual capacity (FRC)
7. Expiratory reserve volume (ERV)
8. Vital capacity (VC)

Combinations of two or more volumes are termed capacities. These volumes and capacities are shown graphically in Figure 14.2. Also shown are the values found in normal adult men. Measurement of these values can provide information about the degree of air-trapping or hyperinflation.


**Functional Residual Capacity (FRC)** Functional residual capacity (FRC) is used to evaluate both restrictive and obstructive lung defects. Changes in the elastic properties of the lungs are reflected in the FRC. The FRC is the volume of gas contained in the lungs at the end of a normal quiet expiration (see Fig. 14.2).

**Reference Values**

**Normal** Approximately 2.50–3.50 L Predicted values are based on age, height, weight, ethnicity, and gender. The observed value should be 75%–125% (0.75–1.25) of the predicted value.

**Procedure**
1. Fit the patient with nose clips, then instruct the patient to breathe through the mouthpiece/filter (bacterial/viral) combination that is attached to the lung volume apparatus. The patient is generally in the seated position.

2. Be aware that there are two methods, depending on the instrument used:
   a. Nitrogen washout or open-circuit technique
   b. Helium dilution or closed-circuit technique

3. Have the patient breathe normally for about 3–7 minutes.

4. Perform the test a second time. Remember that results for FRC should vary by not more than 5% to 10% (0.05 to 0.10). Report the average of the test values.

5. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. A value < 75% (<0.75) of the predicted is consistent with restrictive ventilatory impairment.
2. A value > 125% (>1.25) of predicted demonstrates air-trapping (hyperinflation), consistent with obstructive airway disease (eg, emphysema, asthma, bronchiolar obstruction).

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test. Explain that this is a noninvasive test requiring patient cooperation. Assess the patient's ability to comply.
2. Record the patient's age, gender, weight, and height.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Explain test outcomes; allow the patient to rest if necessary.
2. See Chapter 1 for safe, effective, informed posttest guidelines.

Residual Volume (RV)

Residual volume (RV) can help to distinguish between restrictive and obstructive ventilatory defects. It is the volume of gas remaining in the lungs after a maximal exhalation. Because the lungs cannot be completely emptied (ie, a maximal expiratory effort cannot expel all of the gas), RV is the only lung volume that cannot be measured directly from the spirometer. It is calculated mathematically by subtracting the expiratory reserve volume (ERV) from the FRC (see Fig. 14.2).

Reference Values

Normal Approximately 1200–1500 mL Predicted values are based on age, gender, and height.

Procedure

1. Remember that the RV is determined indirectly from other tests, ie, it is mathematically derived by subtracting the measured expiratory reserve volume (ERV) from the FRC.
2. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. An increase in the RV (>125% >1.25] of predicted) indicates that, despite a maximal expiratory effort, the lungs still contain an abnormally large amount of gas (air-trapping). This type of change occurs in young asthmatic patients and usually is reversible. In emphysema, the condition is permanent.
2. Increased RV is characteristic of emphysema, chronic air-trapping, and chronic bronchial obstruction.
3. The RV and the FRC usually increase together, but not always.
4. The RV sometimes decreases in diseases that occlude many alveoli.
5. An RV < 75% (<0.75) of predicted is consistent with restrictive disorders (eg, interstitial pulmonary fibrosis).

Interfering Factors Residual volume normally increases with age.

Interventions

Pretest Patient Care

1. Explain the purpose of the test and how the results are calculated.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and monitor as necessary.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Expiratory Reserve Volume (ERV)

Expiratory reserve volume (ERV) is the largest volume of gas that can be exhaled from end-tidal expiration. This measurement identifies lung or chest wall restriction. The ERV can be estimated mathematically by subtracting the inspiratory capacity (IC) from the vital capacity (VC). The ERV accounts for approximately 25% of the VC and can vary greatly in patients of comparable age and height (see Fig. 14.2).

Reference Values

Normal Approximately 1200–1500 mL (1.20–1.50 L) Predicted values are based on age, height, and gender.

Procedure

1. Have the patient either sit or stand. Place nose clips on the nose and instruct the patient to breathe normally
through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Ask the patient to exhale completely and resume normal breathing. Record results on graph paper.
3. Have the patient repeat this maneuver a minimum of at least twice. The measured volumes should be within 5%–10% (0.05–0.10) of one another. Report the average value.

Clinical Implications
1. A decreased ERV indicates a chest wall restriction resulting from nonpulmonary causes.
2. Decreased values are associated with an elevated diaphragm (eg, massive obesity, ascites, pregnancy).
   Decreased values also occur with massive enlargement of the heart, pleural effusion, kyphoscoliosis, or thoracoplasty.
3. Decreases in ERV also are seen in obstruction resulting from an increase in the RV impinging on the ERV.

Interventions
Pretest Patient Care
1. Explain the purpose and procedure of the spirometry test. Inform the patient that the test is noninvasive. Assess the patient's ability to comply with test procedures.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcomes and counsel about respiratory abnormalities.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Inspiratory Capacity (IC) Inspiratory capacity (IC) measures the largest volume of air that can be inhaled from the end-tidal expiratory level. This measurement is used to identify lung or chest wall restrictions. Mathematically, the IC is the sum of the tidal volume (VT) and the inspiratory reserve volume (IRV) (see Fig. 14.2).

Reference Values
Normal Approximately 3000–3300 mL (3.00–3.30 L) Predicted values are based on age, height, and gender.

Procedure
1. Have the patient either sit or stand. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. After several breaths, ask the patient to inhale maximally, expanding the lungs as much as possible from end-tidal expiration. Have the patient the resume normal breathing. Record the results on graph paper.
3. Repeat step 2 two or more times until the two best values are within 5% of each other. Select the largest inspired volume value.

Clinical Implications
1. Changes in the IC usually parallel increases or decreases in the vital capacity (VC).
2. Decreases in IC can be related to either restrictive or obstructive ventilatory impairments.

Interventions
Pretest Patient Care
1. Instruct the patient about the purpose and procedure of the test and the need for patient cooperation.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test outcomes and monitor patient.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Vital Capacity (VC) Measurement of the vital capacity (VC) identifies defects of lung or chest wall restriction. The VC is the largest volume of gas that can be expelled from the lungs after the lungs are first filled to the maximum extent and then slowly emptied to the maximum extent. Mathematically, it is the sum of the IC and the ERV (see Fig. 14.2).

Reference Values
Normal Approximately 4.50–5.00 L Predicted values are based on age, gender, height, and ethnicity.

Procedure
1. Have the patient either sit or stand. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Instruct the patient then to inhale as deeply as possible and then to exhale completely, with no forced or rapid effort.
3. Record results on graph paper.
4. Repeat the procedure until the measurements are within about 5% of each other.

Clinical Alert
Inadequate patient effort causes lower VC values.

Clinical Implications
1. A reduced VC is defined as a value < 80% (<0.80) of predicted.
2. The VC can be lower than expected in either a restrictive or an obstructive disorder.
3. A decreased VC can be related to depression of the respiratory center in the brain, neuromuscular diseases, pleural effusion, pneumothorax, pregnancy, ascites, limitations of thoracic movement, scleroderma, kyphoscoliosis, or tumors.
4. The VC increases with physical fitness and greater height.
5. The VC decreases with age (after age 30 years).
6. The VC is generally less in women than in men of the same age and height.
7. The VC is decreased by approximately 15% in African Americans and by 20% to 25% in Asians, compared with Caucasians of the same age, height, and gender.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test and need for patient cooperation. Assess for interfering factors.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret outcomes, monitor patient signs and symptoms, and follow up if necessary.
2. See Chapter 1 for safe, effective, informed posttest care guidelines.

Total Lung Capacity (TLC)

Total lung capacity (TLC) is used mainly to evaluate obstructive defects and to differentiate restrictive from obstructive pulmonary disease. It measures the volume of gas contained in the lungs at the end of a maximal inspiration. Mathematically, it is the sum of the VC and the RV, or the sum of the primary lung volumes (see Fig. 14.2). This value is calculated indirectly from other tests.

Reference Values

Normal
Approximately 5.70–6.20 L Predicted values are based on age, height, gender, and ethnicity. All pulmonary volumes and capacities are about 20%–25% less in women than in men.

Procedure

1. Have the patient either sit or stand. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Ask the patient to inspire maximally and exhale maximally. The total amount of air exhaled is the VC.
3. Use the following formula to derive the TLC mathematically: TLC = VC + RV.

Clinical Implications

1. An obstructive impairment is characterized by an increased TLC. However, a normal or increased TLC does not mean that ventilation or the surface area for diffusion is normal. The TLC may be normal or increased in bronchiolar obstruction with hyperinflation and in emphysema.
2. The TLC is decreased in edema, atelectasis, neoplasms, pulmonary congestion, pneumothorax, and thoracic restriction.
3. A decreased TLC is the hallmark of a restrictive ventilatory impairment.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test. Even though it is noninvasive, it does require patient effort and cooperation.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret outcomes (see Fig. 14.3), monitor patient signs and symptoms, and follow up if necessary.

FIGURE 14.3 Pulmonary function report of a 47-year-old female whose chief complaint is shortness of breath. The report includes spirometry, lung volumes, diffusion capacity, maximum voluntary ventilation, and maximal respiratory pressures. Note: The shape or configuration of the flow-volume loop (lower left corner of report) is
significant for airflow obstruction (ie, obstructive ventilatory impairment). The current flow-volume loop is essentially normal in appearance. (Courtesy of Pulmonary Diagnostic Laboratory, Froedtert Hospital, Milwaukee, WI)

2. See Chapter 1 for safe, effective, informed posttest care guidelines.

Gas Exchange (Diffusing Capacity)

Gas exchange in the lungs is referred to as respiration, whereas the movement of gas in and out of the lung is ventilation. In Europe, the diffusing capacity is termed the transfer factor. Gas exchange involves the movement of oxygen (O$_2$) from the alveolus (gas exchange units in the lung) to the blood (ie, diffusion across the alveolar/capillary membrane) and movement of carbon dioxide (CO$_2$) from the blood into the alveolus for subsequent removal.

Carbon Monoxide Diffusing Capacity (DLCO) The diffusing capacity measurement determines the rate of gas transfer across the alveolar capillary membranes. Carbon monoxide (CO) combines with hemoglobin about 210 times more readily than does O$_2$. If there is a normal amount of hemoglobin in the blood, the only other significant limiting factor to CO uptake is the state of the alveolar capillary membranes. Normally, the amount of CO in the blood is insufficient to affect the test. Two categories of factors (ie, physical and chemical) determine the rate of gas (CO) transfer across the lung. The physical determinants are CO driving pressure, surface area, thickness of capillary walls, and diffusion coefficient for CO. The chemical determinants are red blood cell volume and reaction rate with hemoglobin. This test is used to diagnose pulmonary vascular disease, emphysema, and pulmonary fibrosis and to evaluate the extent of functional pulmonary capillary bed in contact with functional alveoli. The alveolar volume (VA) can also be determined. The DLCO measures the diffusing capacity of the lungs for CO. The DLO$_2$ is obtained by multiplying the DLCO by 1.23.

Reference Values

Normal Approximately 25 mL/min/mmHg (8.4 mmol/min/kPa) Predicted values are based on the patient’s height, age, and gender.

Procedure

1. Have the patient either sit or stand. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the diffusion instrument.
2. Ask the patient to expire maximally and then inspire maximally (a diffusion gas mixture), hold breath for 10 seconds, and then exhale, at which time a sample of exhaled gas is obtained.
3. Two techniques are used by laboratories:
   a. Single-breath or breath-holding technique
   b. Steady-state technique
4. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. Decreased values are associated with:
   a. Multiple pulmonary emboli
   b. Emphysema
   c. Lung resection
   d. Pulmonary fibroses:
      1. Sarcoidosis
      2. Systemic lupus erythematosus (SLE)
      3. Asbestosis
      4. Pneumonia
   e. Anemia
   f. Increased levels of carboxyhemoglobin (COHb)
   g. Pulmonary resection
   h. Scleroderma
2. Increased values are observed in polycythemia, left-to-right shunts, pulmonary hemorrhage, and exercise.
3. The value is relatively normal in chronic bronchitis.

Interfering Factors Exercise (with an increased cardiac output) and polycythemia increase the value. Because increased levels of COHb (as seen in smokers) and anemia decrease the value, the DLCO is corrected for COHb levels > 10% (>0.10) and hemoglobin (Hb) values < 8 g/dL (<80 g/L).

Interventions

Pretest Patient Care

1. Explain the purpose and procedure. Assess for interfering factors and explain that this noninvasive test requires patient cooperation. Assess the patient’s ability to comply.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Explain test outcomes (see Fig. 14.3) and possible need for follow-up testing to monitor course of therapy (eg, anti-inflammatory drugs, bronchodilators, and some antiarrhythmics and antineoplastics).
2. See Chapter 1 for safe, effective, informed posttest care guidelines.

Maximum Voluntary Ventilation (MVV) Maximum voluntary ventilation (MVV) measures several physiologic phenomena occurring at the same time, including thoracic cage compliance, lung compliance, airway resistance, and available muscle force. It is the number of liters of air that the patient can breathe per minute with maximal voluntary effort.

Reference Values

Normal Approximately 160–180 L/min Predicted values are based on the patient’s age, height, and gender. A healthy
person may vary by as much as 25%–35% from mean group values.

**Procedure**

1. Have the patient either sit or stand. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Instruct the patient to breathe into the spirometer as deeply and rapidly as possible for 10 to 15 seconds. Usually, the frequency reaches 40 to 70 breaths per minute, and the tidal volumes are about 50% of VC.
3. Extrapolate actual values from the 10- to 15-second time interval to a 1-minute time period.
4. Be aware that typically, the maneuver is performed twice. Report the largest value.

**Interfering Factors** Poor patient effort can be ruled out by using the following formula to predict the MVV of the patient:

$$\text{Predicted MVV} = 35 \times \text{FEV}_1$$

This is a useful check to determine whether the recorded MVV is indicative of adequate patient effort. Low values can be related to patient effort and not to pathophysiology.

**Clinical Implications**

1. Obstructive ventilatory impairments of moderate to severe degree, abnormal neuromuscular control, and poor patient effort are causes of low values.
2. In restrictive disease, the value is usually normal; however, in more severe forms, MVV may be decreased.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the test. Explain that it is a noninvasive test that requires patient cooperation. Assess the patient's ability to comply.
2. Record the patient's age, height, and gender.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Explain test outcome (see Fig. 14.3) and possible need for follow-up testing and treatment.
2. See Chapter 1 regarding safe, effective, informed posttest care guidelines.

**Maximal Respiratory Pressure (MRP)**

The maximal respiratory pressure (MRP) measurements assess ventilatory muscle strength in persons with neuromuscular disorders such as poliomyelitis, emphysema, and pulmonary fibroses. The maximal expiratory pressure (MEP) is the greatest pressure that can be generated at or near total lung capacity after a maximal inspiration, whereas the maximal inspiratory pressure (MIP) is measured at or near the residual volume after a maximal expiration.

**Reference Values**

**Normal**

- Maximal expiratory pressure (MEP): approximately 100–250 cm H₂O
- Maximal inspiratory pressure (MIP): approximately 40–125 cm H₂O

Predicted values are based on the patient's age and gender.

**Procedure**

1. Instruct the patient, who should be in a seated position and wearing a nose clip, to inspire maximally. Place the mouthpiece of the handheld pressure manometer into the mouth and have the patient perform a forced expiration. Record this maximal sustained (1 to 3 seconds) pressure against the internal occlusion of the manometer as the MEP.
2. Repeat this same procedure to obtain the MIP, except that this time the patient fully exhales before placing the mouthpiece of the manometer in the mouth. Have the patient then inspire forcefully, and record the maximal sustained (1 to 3 seconds) pressure.
3. Repeat each procedure, and record the best of three measurements for each.
4. See Chapter 1 guidelines for intratest care.

**Interfering Factors** The MIP and MEP measurements depend on patient effort; low values may be caused by poor effort rather than loss of respiratory muscle strength. If the patient does not inspire or expire maximally before performing the pressure measurement, the value may be low. Also, sustained efforts longer than 3 seconds should be avoided because they can cause a decrease in cardiac output as a result of increased intrathoracic pressures.

**Clinical Implications**

1. Decreases in both MEP and MIP are seen in neuromuscular disorders (eg, myasthenia gravis, poliomyelitis).
2. Decreased MEP is common in both severe obstructive disease (eg, emphysema) and severe restrictive ventilatory impairment (eg, interstitial pulmonary fibrosis).
3. Decreased MIP is observed in patients with chest wall abnormalities (eg, kyphoscoliosis) and in hyperinflation (eg, emphysema).

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the test. Explain that it is a noninvasive, effort-dependent maneuver that requires patient cooperation.
2. Record the patient's age and sex.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
1. Explain test outcomes (see Fig. 14.3) and possible need for follow-up testing and treatment.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Closing Volume (CV) In a healthy person, the concentration of alveolar nitrogen, after a single breath of 100% $O_2$, rapidly increases near the end of expiration. This rise is caused by closure of the small airways in the bases of the lung. The point at which this closure occurs is called the closing volume (CV). CV is used as an index of pathologic changes occurring within the small airways (those < 2 mm in diameter). The conventional pulmonary function tests are not sensitive enough to make this determination. This test relies on the fact that the upper lung zones contain a proportionately larger residual volume of gas than the lower lung zones do; there is a gradient of intrapleural pressure from the top to the bottom of the lung. Additionally, the uniformity of gas distribution within the lungs can be measured.

Reference Values
Normal Average is 10% to 20% (0.10 to 0.20) of the patient's vital capacity (VC) Predicted values are derived from mathematical regression equations and are based on the patient's age and gender.

Procedure
1. Have the patient assume a seated position. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Ask the patient to exhale completely, to inhale 100% $O_2$, and then to exhale completely at the rate of approximately 0.5 L/second.
3. During exhalation, monitor simultaneously both the expired volume and percentage of alveolar nitrogen on an X-Y recorder. Remember that a sudden increase in nitrogen represents the closing volume.

Clinical Implications
1. Values are increased for those conditions in which the airways are narrowed (eg, bronchitis, early airway obstruction, chronic smokers, old age).
2. A change in the slope of the nitrogen curve of > 2% is indicative of maldistribution of inspired air (ie, uneven alveolar ventilation).
3. Congestive heart failure, with subsequent edema, may also contribute to decreasing patency of the small airways leading to an increase in the CV.

Interfering Factors
1. The CV increases with age.
2. Patients in congestive heart failure may show an increased CV.

Interventions
Pretest Patient Care
1. Explain the purpose and procedure of the test. Explain that this is a noninvasive test that requires patient cooperation. Assess the patient's ability to comply with breathing requirement and instructions. Assess for interfering factors.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Explain the meaning of test outcomes and possible need for follow-up testing and treatment of early small airway disease.
2. See Chapter 1 for safe, effective, informed posttest care guidelines.

Volume of Isoflow (VISO)
This test is designed to detect pathologic changes occurring in the small airways and may be more sensitive than conventional pulmonary function tests. Helium has the unique property of lowering gas density. Therefore, after the patient breathes a helium-oxygen gas mixture, the effects of convective acceleration and turbulence are negated. Any abnormality observed in the F-V loop, then, results from an increase in resistance to laminar (nonturbulent) flow, which indicates small airway abnormalities or lung disease.

Reference Values
Normal Average is 10%–25% of VC. Predicted values are based on age.

Procedure
1. Have the patient assume a seated position. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Have the patient perform a baseline F-V loop, which is recorded by a spirometer on an X-Y recorder.
3. Have the patient next breathe a mixture of 80% He and 20% $O_2$ for several breaths and then perform another F-V loop maneuver; this is the HeliOx F-V loop.
4. Superimpose the F-V loop tracings, and measure the volume of isoflow at the point at which the two loops intersect.

Clinical Implications An increased volume of isoflow is consistent with early small airway obstruction (eg, asthma).

Interventions
Pretest Patient Care
1. Explain the purpose and procedure of the test.
2. Follow guidelines in Chapter 1 regarding safe, effective, pretest care.
Posttest Patient Aftercare

1. Interpret test outcomes and possible need for follow-up testing and treatment.
2. See Chapter 1 guideline for safe, effective, informed posttest care.

Body Plethysmography: Thoracic Gas Volume (VTG), Compliance (CL), Airway Resistance (R_{aw})

This test measures several parameters. Thoracic gas volume (\( V_{TG} \)) comprises all the air contained within the thorax, whether or not it is in ventilatory communication with the rest of the lung. Compliance of the lung (CL) is an indication of its elasticity, and airway resistance (\( R_{aw} \)) is a measurement of the resistance to airflow in the tracheobronchial tree. The measurement of VTG via body plethysmography is an application of Boyle's law, which states that, for a gas at constant temperature, pressure and volume vary inversely (\( P_1V_1 = P_2V_2 \)). Airway resistance (\( R_{aw} \)) increases with decreased lung volumes and decreases with higher lung volumes in a nonlinear, hyperbolic fashion. Compliance (CL) increases in obstructive diseases (eg, emphysema) and decreases in restrictive processes (eg, interstitial lung disease).

Reference Values

**Normal** Thoracic gas volume (VTG): approximately 2.50–3.50 L
Compliance (CL): 0.2 L/cm H\(_2\)O (2.04 L/kPa)
Airway resistance (\( R_{aw} \)): 0.6–2.4 L/s/cm H\(_2\)O

Predicted values are based on the patient's age, height, weight, and gender.

Procedure

1. Have the patient sit in the plethysmograph (body box). Fit with nose clips, and have the patient breathe through a mouthpiece/filter (bacterial/viral) combination connected to a transducer (Fig. 14.4).

   ![Body plethysmograph](https://via.placeholder.com/150)

   **FIGURE 14.4** Body plethysmograph. (Source: SensorMedics Corp., a subsidiary of VIASYS Healthcare, Yorba Linda, CA, USA)

2. Ensure that the body box door is secured. Delay the test for a few minutes to allow the box pressure to stabilize.
3. Instruct the patient to perform a panting maneuver while holding the cheeks rigid and the glottis open against a closed shutter located within the transducer assembly. Record box and mouth pressures on the oscilloscope to provide data for VTG.
4. Next, ask the patient to breathe rapidly and shallowly. Record box pressure changes versus flow on the oscilloscope to provide data for \( R_{aw} \).
5. To determine CL, pass a balloon catheter through the nose into the patient's esophagus. Ensure that the inflated balloon is connected to a transducer, and instruct the patient to breathe normally. Record the changes in intraesophageal pressure during normal respiration (which mimic changes in intrapleural pressure) to provide data for CL.

Clinical Implications

1. An increased VTG demonstrates air-trapping, consistent with obstructive pulmonary disease.
2. An increased \( R_{aw} \) demonstrates increased resistance to airflow through the tracheobronchial tree; this is seen in asthma, emphysema, bronchitis, and other forms of obstruction. The \( R_{aw} \) distinguishes between restrictive and obstructive ventilatory defects.
3. An increase in CL (ie, lung is more distensible) is seen in obstructive diseases.
4. A decrease in CL (ie, lung is more stiff) is seen in fibrotic diseases, restrictive diseases, pneumonia, congestion, and atelectasis.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test.
2. Assure the patient that although the chamber is airtight, the test only takes a few minutes. A technician will be in constant attendance to open the door should that be necessary. Assess for ability to comply with test requirements and instructions. Tactfully assess for predisposition to claustrophobia, panic attacks, or other similar responses.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Allow the patient time to rest quietly if necessary.
2. Explain the meaning of test outcomes.
3. See Chapter 1 for safe, effective, informed posttest care guidelines.

Bronchial Provocation: Methacholine, Histamine Challenge

Bronchial provocation challenge testing is performed in patients with normal pulmonary function tests who have suspected underlying bronchial hyperreactivity. Additionally, the asthmatic patient is more sensitive to the bronchoconstrictive effects of cholinergic agents (eg, methacholine chloride) than is the healthy person as observed on a spirometry test. Airway resistance (\( R_{aw} \)) tests are also sensitive monitors of...
response to bronchoconstrictive agents.

Reference Values

**Normal**
Positive response: >20% (or >0.20) decrease in FEV<sub>1</sub> from baseline or >35% (>0.35) increase in R<sub>aw</sub>
Negative response: <20% (or <0.20) decrease in FEV<sub>1</sub> from baseline or <35% (<0.35) increase in R<sub>aw</sub>

**Procedure**

1. Have the patient assume the seated position. Place nose clips on the nose and instruct the patient to breathe normally through a mouthpiece/filter (bacterial/viral) combination into the spirometer.
2. Have the patient perform a forced expiratory maneuver, and measure and record the baseline FEV<sub>1</sub> (or an R<sub>aw</sub> measurement).
3. Have the patient inhale increasing concentrations of methacholine chloride (0.062–16.00 mg/mL) or histamine by nebulizer. Repeat the FVC or R<sub>aw</sub> maneuver after each successive concentration is inhaled. Be aware that a 20% reduction in the FEV<sub>1</sub> or 35% increase in R<sub>aw</sub> is considered a positive response.
4. Administer an inhaled bronchodilator when or if a decrease of >20% from baseline is reached.
5. Be aware that if a patient goes through all dilution ratios and a 20% reduction in the FEV<sub>1</sub> or >35% increase in R<sub>aw</sub> is not reached, the test is considered negative.
6. Remember that if the methacholine causes no change, histamine testing may be ordered.
7. See Chapter 1 for guidelines for intratest care.

**Clinical Implications** A positive response to methacholine or histamine is consistent with bronchial hyperreactivity. Approximately 5% to 10% of asthmatic persons do not respond to the methacholine challenge test.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the test and the need for patient cooperation. Assess the patient's ability to comply.
2. Withhold bronchodilators for 8 hours and antihistamines for 48 hours before testing, if tolerated.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Clinical Alert**

1. Inhalation of methacholine can cause bronchospasm, chest pain, shortness of breath, and general discomfort.
2. These effects can be reversed with a bronchodilator.

**Posttest Patient Aftercare**

1. Explain the meaning of test outcomes.
2. If the test is positive, advise the patient to avoid antigens that may be causing hypersensitivity reaction and bronchospasms.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Carbon Dioxide (CO<sub>2</sub>) Response** This test evaluates the respiratory response to increasing concentrations of inspired CO<sub>2</sub>. As alveolar levels of CO<sub>2</sub> increase, so does arterial CO<sub>2</sub>. The central chemoreceptors respond by initiating impulses to the respiratory control centers. In the healthy person, this causes the rate and depth of breathing to increase. The act of breathing successively greater concentrations of CO<sub>2</sub> should result in an increase in minute volume (VE), when compared with the VE during breathing of room air alone. (Room air contains 0.03% CO<sub>2</sub>.)

Reference Values

**Normal**
Increase in minute ventilation of 3 L/min/mmHg increase of CO<sub>2</sub> (3 L/min/0.133 kPa)

**Procedure**

1. Remember that \( \text{VE} \) is determined while the patient breathes room air for several minutes into an instrument (eg, spirometer) that records the frequency of breathing (f) and the tidal volume (VT). Use the following formula to calculate the minute volume: \( \text{VE} = f \times VT \).
2. Have the patient breathe a gas mixture of 2% CO<sub>2</sub> in room air for 5 minutes. During the last 2 minutes, record f and VT and calculate the \( \text{VE} \).
3. Have the patient breathe gas mixtures of 4% CO<sub>2</sub> and 6% in room air. Mixtures can be increased to as much as 8% CO<sub>2</sub>. Repeat the entire process with each successive concentration.
4. Construct a graph to plot the changes in VE against the concentration of inspired CO<sub>2</sub> (FICO<sub>2</sub>).
5. See Chapter 1 guidelines for intratest care.

**Clinical Implications** Lack of response to increasing inspired CO<sub>2</sub> concentrations suggests a disturbance in the normal physiologic pathway of ventilatory changes to hypercapnia. This may result from ingestion of central nervous system depressants (eg, anesthetics, barbiturates, narcotics) or from airflow obstruction (eg, chronic obstructive pulmonary disease [COPD]).

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure of the test and need for patient cooperation. Assess the patient's ability to comply.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
1. Interpret the test outcome and advise that pharmacologic intervention may be necessary to sensitize the chemoreceptors.

2. See Chapter 1 guidelines regarding safe, effective, posttest care.

Exercise Stress Testing, Maximum Oxygen Consumption (\(\text{ VO}_2\) max) Test

Respiratory disease reduces the ability to perform exercise. Dynamic exercise that involves large muscle groups produces increases in metabolic \(\text{ O}_2\) consumption (\(\text{ VO}_2\)) and \(\text{ CO}_2\) production (\(\text{ VCO}_2\)). This increase in metabolic demand leads to stresses on other mechanisms taking part in \(\text{ O}_2\) and \(\text{ CO}_2\) transport. Exercise testing measures the functional reserves of these mechanisms by testing under load. Analysis of bronchogenic and cardiovascular disorders includes procedures that measure respiratory outcomes, blood gas values, and cardiovascular responses during exercise. Ventilation and gas exchange are altered during exercise in healthy persons; however, specific abnormalities are noted in the presence of cardiovascular or respiratory impairment. Exercise tests are valuable for assessing the severity and type of impairment in existing or undiagnosed conditions. The normal response to graded exercise is an increase in ventilation and cardiac output such that alveolar and arterial gases are maintained at optimal levels to meet metabolic demands. Measurement of the patient's ventilatory and alveolar-arterial gas responses during exercise is the primary objective of a pulmonary exercise stress test. No significant or abnormal changes in the electrocardiographic (ECG) complex, blood pressure, airflow patterns during inspiration and expiration, arterial blood gases (ABGs) and chemistry, or hemodynamic pressures should occur. Exercise testing is done to evaluate fitness, functional capacity, and other limiting factors in persons with obstructive or restrictive diseases. The efficiency of the cardiopulmonary system may be altered during exercise; exercise testing assesses ventilation, gas exchange, and cardiovascular function during increased demands. Dyspnea on exertion due to cardiovascular causes can be differentiated from that due to respiratory causes. Precise information about mechanisms that influence \(\text{ O}_2\) and \(\text{ CO}_2\) transport during exercise can be obtained by using a staged approach. An exercise test can detect or exclude many conditions, even though the response may be nonspecific. For example, if the patient complains of severe shortness of breath despite a normal exercise response, a psychogenic cause is likely. However, a few conditions exhibit diagnostic responses, eg, exercise-induced asthma or myocardial ischemia. These tests can also reveal the degree of impairment in conditions affecting the respiratory and circulatory systems and may uncover unsuspected abnormalities (Table 14.2).

### Table 14.2 Normal Ventilatory and Arterial Blood Gas Responses to Graded Exercise

<table>
<thead>
<tr>
<th>Value</th>
<th>Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\text{ O}_2) consumption ((\text{ VO}_2))</td>
<td>Increase</td>
</tr>
<tr>
<td>(\text{ CO}_2) production ((\text{ VCO}_2))</td>
<td>Increase</td>
</tr>
<tr>
<td>Ventilatory equivalents for (\text{ O}_2) and (\text{ CO}_2)</td>
<td>No change</td>
</tr>
<tr>
<td>Respiratory exchange ratio (RER)</td>
<td>Increase</td>
</tr>
<tr>
<td>Minute ventilation (VE)</td>
<td>Increase</td>
</tr>
<tr>
<td>Blood lactate</td>
<td>Increase</td>
</tr>
<tr>
<td>(\text{ VD/VT}) ratio</td>
<td>Decrease</td>
</tr>
<tr>
<td>(\text{ A-aDO}_2)</td>
<td>Slight increase</td>
</tr>
<tr>
<td>Arterial blood gas tensions (eg, (\text{ PaO}_2, \text{ PaCO}_2))</td>
<td>No change</td>
</tr>
<tr>
<td>Bicarbonate concentration ((\text{ HCO}_3^-))</td>
<td>Decrease</td>
</tr>
<tr>
<td>Oxygen saturation ((\text{ SaO}_2))</td>
<td>No change</td>
</tr>
</tbody>
</table>

The majority of clinical problems can be assessed during the simple procedures included in stage 1 (see Procedure section for description) and should be done before more complex tests. Abnormal results indicate that more precise information is required through stage 2 protocols. If stage 3 protocols are implemented, arterial blood analysis is necessary. In 75% of cases, stage 1 is sufficient. Oxygen titration can be done during graded exercise to determine the oxygen needs for improving exercise tolerance and increasing functional capacity.

### Clinical Alert

1. **Absolute contraindications** to exercise testing include: Acute febrile illness
   - Pulmonary edema
   - Systolic blood pressure >250 mm Hg (>33 kPa)
   - Diastolic blood pressure >120 mm Hg (>16 kPa)
   - Uncontrolled hypertension
   - Uncontrolled asthma
   - Unstable angina

2. **Relative contraindications** to exercise testing include: Recent myocardial infarction (<4 weeks)
   - Resting tachycardia (>120 bpm)
   - Epilepsy
   - Respiratory failure
   - Resting ECG abnormalities

### Reference Values

**Normal** Increase in ventilation, heart rate, and blood pressure appropriate to the level of exercise No abnormal changes in the ECG (no arrhythmias), ABGs, or hemodynamic pressures

### Procedure
1. **Stage 1**
   a. Record blood pressure readings, ECG analysis, and ventilation during incremental cycle ergometry or treadmill walking.
   b. Take measurements at the end of each minute. Remember that the test continues until maximum-allowed symptoms occur (ie, to a symptom-limited maximum). Measure O\(_2\) uptake (\(\text{VO}_2\)) and CO\(_2\) output (\(\text{VCO}_2\)) if possible.
   c. Alert patient that total examination time is approximately 30 minutes.

2. **Stage 2**
   a. Be aware that more complex analytic methods are required.
   b. Have exercise build to a steady state, usually 3 to 5 minutes for each workload.
   c. In addition to stage 1 measurements, determine mixed venous CO\(_2\) tension by means of rebreathing techniques.

3. **Stage 3**
   a. Be aware that blood gas sampling and analysis are required.
   b. Insert an indwelling catheter into the brachial or radial artery.
   c. In addition to stage 2 tests, determine measurements for cardiac output, alveolar ventilation, ratio of dead space to tidal volume (\(\text{VD}/\text{VT}\)), alveolar-arterial O\(_2\) tension difference (\(\text{A-aDO}_2\)), venous admixture ratio, and lactate concentration.

4. See Chapter 1 guidelines for intratest care.

**Clinical Implications**

Altered values may reveal:

1. Cardiac dysrhythmias or ischemia
2. Degree of functional impairment caused by obstructive or restrictive ventilatory disease
3. Hypoventilation
4. Workload level at which metabolic acidosis (lactic acidosis) occurs

**Interfering Factors**

1. The exercise tolerance of any person is affected by the degree of impairment related to:
   a. Mechanical factors
   b. Ventilatory efficiency
   c. Gas exchange factors
   d. Cardiac status
   e. Physical condition
   f. Sensitivity of the respiratory control mechanism
2. Obese persons have a higher-than-normal oxygen consumption at any given work rate, even though muscular and work efficiency values are normal.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose and procedure for exercise stress testing and assess for contraindications, interfering factors, and ability to comply.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Explain the meaning of test outcomes and possible need for lifestyle changes.
2. See Chapter 1 for safe, effective, informed posttest care guidelines.

**Arterial Blood Gases (ABGs)**

**Overview of Arterial Blood Gas Tests**

Measurements of arterial blood gases (ABGs) are obtained to assess adequacy of oxygenation and ventilation, to evaluate acid-base status by measuring the respiratory and nonrespiratory components, and to monitor effectiveness of therapy. They are also used to monitor critically ill patients, to establish baseline values in the perioperative and postoperative period, to detect and treat electrolyte imbalances, to titrate appropriate oxygen flow rates, to qualify a patient for use of oxygen at home, and in conjunction with pulmonary function testing.

Reasons for using arterial rather than venous blood to measure blood gases include the following:

1. Arterial blood provides a better way to sample a mixture of blood from various parts of the body.
   a. Venous blood from an extremity gives information mostly about that extremity. The metabolism in the extremity can differ from the metabolism in the body as a whole. This difference is accentuated in the following instances:
      1. In shock states, when the extremity is cold or underperfused
      2. During local exercise of the extremity, as in opening and closing a fist
      3. If the extremity is infected
   b. Blood from a central venous catheter usually is an incomplete mix of venous blood from various parts of the body. For a sample to be completely mixed, the blood would have to be obtained from the right ventricle or pulmonary artery.
2. Arterial blood measurements indicate how well the lungs are oxygenating blood.
a. If it is known that the arterial $\text{O}_2$ concentration is normal (indicating that the lungs are functioning normally) but the mixed venous $\text{O}_2$ concentration is low, it can be inferred that the heart and circulation are failing.

b. Oxygen measurements of central venous catheter blood reveal tissue oxygenation but do not separate contributions of the heart from those of the lungs. If central venous catheter blood has a low $\text{O}_2$ concentration, it means either that the lungs have not oxygenated the arterial blood well or that the heart is not circulating the blood effectively. In the latter case, the body tissues must take on more than the normal amount of $\text{O}_2$ from each cardiac cycle because the blood is flowing slowly and permits this to occur; this produces a low venous $\text{O}_2$ concentration.

3. Arterial samples provide information about the ability of the lungs to regulate acid-base balance through retention or release of $\text{CO}_2$. Effectiveness of the kidneys in maintaining appropriate bicarbonate levels also can be gauged.

**NOTE**

Arterial puncture sites must satisfy the following requirements: (1) available collateral blood flow; (2) superficial or easily accessible location; and (3) relatively nonsensitive periarterial tissues.

The radial artery is usually the site of choice, but brachial and femoral arteries can also be used. Samples can be drawn from direct arterial sticks or from indwelling arterial lines.

**Clinical Alert**

1. Before obtaining an arterial blood sample, assess for the following contraindications to an arterial stick or indwelling line: Absent palpable radial artery pulse
   - Negative modified Allen's test, indicating obstruction in the ulnar artery (i.e., compromised collateral circulation)—do not attempt to use radial artery for blood sample
   - Cellulitis or infection in the area
   - Severe thrombocytopenia
   - Prolonged prothrombin or partial thromboplastin time (relative contraindication)

2. A Doppler probe or finger-pulse transducer may be used to assess circulation. This may be especially helpful with dark-skinned or uncooperative patients.

3. Before obtaining an arterial blood sample, record the most recent hemoglobin (Hb) concentration, the mode and flow of oxygen therapy, and the temperature. If the patient has recently undergone suctioning or been placed on mechanical ventilation, or if the inspired oxygen concentration has been changed, wait at least 15 minutes before drawing the sample. This waiting period allows circulating blood levels to return to baseline. Hyperthermia and hypothermia also influence oxygen release from hemoglobin at the tissue level.

**Reference Values**

*Normal* See Table 14.3.

**Table 14.3 Normal Values for Commonly Ordered Arterial Blood Gas Studies**

<table>
<thead>
<tr>
<th></th>
<th>Adults</th>
<th>Pediatrics</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>7.35–7.45</td>
<td>7.32–7.42</td>
</tr>
<tr>
<td>Pa $\text{CO}_2$</td>
<td>35–45 mmHg (4.6–5.9 kPa)</td>
<td>30–40 mmHg (4.0–5.3 kPa)</td>
</tr>
<tr>
<td>Pa $\text{O}_2$</td>
<td>$&gt;$80 mmHg (&gt;10.6 kPa)</td>
<td>80–100 mmHg (10.6–13.3 kPa)</td>
</tr>
<tr>
<td>Sa $\text{O}_2$</td>
<td>$&gt;$94% (&gt;0.94)</td>
<td></td>
</tr>
<tr>
<td>$\text{CO}_2$ content</td>
<td>45–51 vol% (19.3–22.4 mmol/L)</td>
<td></td>
</tr>
<tr>
<td>$\text{O}_2$ content</td>
<td>15–22 vol % (6.6–9.7 mmol/L)</td>
<td></td>
</tr>
<tr>
<td>Base excess</td>
<td>$&gt;$2 mEq/L (&gt;2 mmol/L)</td>
<td></td>
</tr>
<tr>
<td>Base deficit</td>
<td>$&lt;$–2 mEq/L (&lt;–2 mmol/L)</td>
<td></td>
</tr>
<tr>
<td>HCO$_3^−$</td>
<td>22–26 mEq/L (22–26 mmol/L)</td>
<td></td>
</tr>
<tr>
<td>Hb</td>
<td>12–16 g/dL or 120–160 g/L (women); 13.5–17.5 g/dL or 135–175 g/L (men)</td>
<td></td>
</tr>
<tr>
<td>Hct</td>
<td>37%–47% (women); 40%–54% (men)</td>
<td></td>
</tr>
<tr>
<td>COHb</td>
<td>$&lt;$2% (&lt;0.02)</td>
<td></td>
</tr>
<tr>
<td>[NA $^+$]</td>
<td>135–148 mEq/L (135–148 mmol/L)</td>
<td></td>
</tr>
<tr>
<td>[K $^+$]</td>
<td>3.6–5.2 mEq/L (3.6–5.2 mmol/L)</td>
<td></td>
</tr>
<tr>
<td>[Ca $^{++}$]</td>
<td>4.2–5.1 mEq/L (2.1–2.5 mmol/L)</td>
<td></td>
</tr>
<tr>
<td>[Cl $^-$]</td>
<td>98–106 mEq/L (98–106 mmol/L)</td>
<td></td>
</tr>
</tbody>
</table>

**Procedure**

1. Observe standard precautions and follow agency protocols.
2. Have the patient assume a sitting or supine position.
3. Perform the modified Allen’s test to assess collateral circulation before performing a radial puncture, as follows:
   - a. Use pressure to obliterate both radial and ulnar pulses.
   - b. Make the hand blanch, then release pressure over only the ulnar artery. In a positive test, note flushing
immediately; the radial artery may then be used for puncture.

c. If collateral circulation from the ulnar artery is inadequate (negative test), choose another site.

4. Elevate the patient's wrist with a small pillow, and ask the patient to extend the fingers downward (this flexes the wrist and positions the radial artery closer to the surface).

5. Palpate the artery and maneuver the patient's hand back and forth until a satisfactory pulse is felt.

6. Swab the area liberally with an antiseptic agent (eg, an agent with an iodine base).

7. Optional: After assessing for allergy, inject the area with a small amount (<0.25 mL) of 1% plain lidocaine (Xylocaine) if necessary to anesthetize site. This allows for a second attempt without undue pain.

8. Prepare a 20- or 21-gauge needle on a preheparinized self-filling syringe, puncture the artery, and collect a 3- to 5-mL sample. During the procedure, if the patient feels a dull or sharp pain radiating up the arm, withdraw the needle slightly and reposition it. If repositioning does not alleviate the pain, the needle should be withdrawn completely.

9. Withdraw the needle and place a 4- × 4-inch absorbent bandage over the puncture site. Maintain pressure over the site with two fingers for a minimum of 2 minutes or until no bleeding is evident; it may be necessary to use a pressure dressing, secured to the site with elastic tape, for several hours.

10. Meanwhile, ensure that all air bubbles in the blood sample are expelled as quickly as possible. Air in the sample changes ABG values. Cap the syringe and gently rotate to mix heparin with the blood.

11. Label the sample with patient's name, identification number, date, time, mode of O₂ therapy, and flow rate.

12. Place the sample on ice and transfer it to the laboratory. This prevents alterations in gas tensions resulting from metabolic processes that continue after blood is drawn.

13. See Chapter 1 guidelines for intratest care.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure for obtaining an arterial blood sample.
2. If the patient is apprehensive, explain that a local anesthetic can be used.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Evaluate color, motion, sensation, degree of warmth, capillary refill time, and quality of pulse in the affected extremity or at the puncture site.
2. Monitor puncture site and dressing for arterial bleeding for several hours. No vigorous activity of the extremity should be undertaken for 24 hours.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

1. Some patients experience lightheadedness, nausea, or vasovagal syncope during arterial puncture. Respond according to established protocols.
2. ABG measurements do not indicate the degree of an abnormality. For this reason, the vital signs and mental function of the patient must be used as guides to determine adequacy of tissue oxygenation.
3. Pressure must be applied to the arterial puncture site, and the site must be watched carefully for bleeding for several hours. Instruct the patient to report any bleeding from the site.
4. Information for the laboratory should include the fraction of inspired oxygen (FIO₂), which is 0.21 (21%) for room air, and the time when the sample was obtained. Do not use blood for ABG measurements if sample is >3 hours old.
5. In the clinical setting (eg, perioperative or intensive care environment), ABG studies usually include the following: pH, PaCO₂, SaO₂, CO₂ content, O₂ content, PaO₂, base excess or deficit, HCO₃⁻, hemoglobin, hematocrit, CO, Na⁺, and K⁺ (see Table 14.3).

Alveolar-to-Arterial Oxygen Gradient (A-aDO₂); Arterial-to-Alveolar Oxygen Ratio (a/A Ratio)

This test gives an approximation of the difference in the partial pressure of O₂ between the alveoli and arteries. The alveolar to arterial oxygen gradient assesses oxygen delivery by comparing the arterial oxygen level to the theoretical maximum alveolar oxygen level. It identifies the cause of hypoxemia and intrapulmonary shunting as either (1) ventilated alveoli but no perfusion, (2) unventilated alveoli with perfusion, or (3) collapse of both alveoli and capillaries.

Reference Values

Normal A-aDO₂: <10 mm Hg ( <1.33 kPa) at rest (room air) A-aDO₂: 20–30 mm Hg (2.7–4.0 kPa) at maximum exercise (room air) a/A ratio: 75%

Procedure

1. Obtain and analyze an arterial blood sample. This gives the arterial partial pressures of oxygen (PaO₂) and of carbon dioxide (PaCO₂). The barometric pressure (PB) and water vapor pressure (PH₂O) are also known, as is the fractional concentration of inspired oxygen (FIO₂), which is 0.21 (21%) for room air.
2. From these, derive the alveolar oxygen tension (PAO₂), the arterial-to-alveolar oxygen ratio (a/A ratio), and the alveolar-to-arterial difference for PO₂ (A-aDO₂) by use of formulas.

Clinical Implications

1. Increased values may be caused by:
   a. Mucus plugs
   b. Bronchospasm
c. Airway collapse, as seen in:
   1. Asthma
   2. Bronchitis
   3. Emphysema

2. Hypoxemia (increased A-a\textsubscript{DO$_2$}) is caused by:
   a. Atrial septal defects
   b. Pneumothorax
   c. Atelectasis
   d. Emboli
   e. Edema

**Interfering Factors** Values increase with age (age in years +10 divided by 4 gives an estimate of a normal gradient) and increasing O\textsubscript{2} concentration (gradient increases by 5–7 mm Hg [0.6–0.9 kPa] for every 10% increase in oxygen).

**Interventions**

**Pretest Patient Care**

1. Explain the purpose, benefits, and risks of arterial blood sampling (see page 928).
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome and assess, monitor, and intervene appropriately for hypoxemia and ventilatory disturbances.
2. Frequently observe the puncture site for bleeding (see page 928).
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Partial Pressure of Carbon Dioxide (PCO\textsubscript{2})** This test measures the pressure or tension exerted by dissolved CO\textsubscript{2} in the blood (10% of CO\textsubscript{2} is carried in plasma and 90% in red blood cells) and is proportional to the partial pressure of CO\textsubscript{2} in the alveolar air. The test is commonly used to detect a respiratory abnormality and to determine the alkalinity or acidity of the blood. To maintain CO\textsubscript{2} within normal limits, the rate and depth of respiration vary automatically with changes in metabolism. This test is an index of the effectiveness of alveolar ventilation; it is the most physiologically reflective blood gas measurement. An arterial sample directly reflects how well air is exchanged with blood in the lungs. CO\textsubscript{2} tension in the blood and in cerebrospinal fluid is the major chemical factor regulating alveolar ventilation. When the CO\textsubscript{2} tension in arterial blood (PaCO\textsubscript{2}) rises from 40 to 45 mmHg (5.3 to 6.0 kPa), it causes a 3-fold increase in alveolar ventilation. A PaCO\textsubscript{2} of 63 mmHg (8.4 kPa) increases alveolar ventilation 10-fold. When the FICO\textsubscript{2} is > 0.05 (5%), the lungs can no longer be ventilated fast enough to prevent a dangerous rise of CO\textsubscript{2} concentration in tissue fluids. Any further increase in CO\textsubscript{2} begins to depress the respiratory center, causing a progressive decline in respiratory activity rather than an increase.

**Reference Values**

**Normal** PaCO\textsubscript{2} (arterial blood): 35–45 mmHg (4.7–6.0 kPa) P\textsubscript{v}CO\textsubscript{2} (venous blood): 41–57 mmHg (5.4–7.6 kPa)

**Procedure**

1. Obtain an arterial blood sample (or venous sample if requested) according to protocols. See page 929 for arterial blood sample specimen collection and Chapter 2 for venous blood sample specimen collection.
2. Introduce a small amount of this blood into a blood gas analyzing instrument (Fig. 14.5), and measure the CO\textsubscript{2} tension by a silver–silver chloride electrode.

![FIGURE 14.5 Blood gas and electrolyte analyzer. (Source: Radiometer Copenhagen, Radiometer America, Inc., Westlake, OH, USA)](source)

**Clinical Implications**

1. A rise in PaCO\textsubscript{2} (hypercapnia) usually is associated with hypoventilation (CO\textsubscript{2} retention); a decrease is associated with hyperventilation (“blowing off” CO\textsubscript{2}). A reduction in PaCO\textsubscript{2}, through its effect on the plasma bicarbonate concentration, decreases renal bicarbonate reabsorption. For each 1-mm Hg (0.133-kPa) decrease in the PaCO\textsubscript{2}, the plasma bicarbonate will decrease by approximately 1 mEq/L (1 mmol/L). Because HCO\textsubscript{3}~ and PaCO\textsubscript{2} bear this close mathematical relationship, and this ratio, in turn, defends the hydrogen ion concentration, the outcome is that the steady-state PaCO\textsubscript{2} in simple metabolic acidosis is equal to the last two digits of the arterial pH (pHa). Also, addition of 15 to the bicarbonate level equals the last two digits of the pHa. Failure of the PaCO\textsubscript{2} to achieve predicted levels defines the presence of superimposed respiratory acidosis on alkalosis.

2. Causes of decreased PaCO\textsubscript{2} include:
   a. Hypoxia
   b. Nervousness
   c. Anxiety
   d. Pulmonary emboli
e. Pregnancy  
f. Pain  
g. Other cause of hyperventilation

3. Causes of increased PaCO₂ include:

a. Obstructive lung disease
   1. Chronic bronchitis  
   2. Emphysema  

b. Reduced function of respiratory center
   1. Overreaction  
   2. Head trauma  
   3. Anesthesia  

c. Other, less common causes of hypoventilation (eg, pickwickian syndrome)

Clinical Alert

Increased PaCO₂ may occur, even with normal lungs, if the respiratory center is depressed. Always check laboratory reports for abnormal values. When interpreting laboratory reports, remember that PaCO₂ is a gas and is regulated by the lungs, not the kidneys.

Interventions

Pretest Patient Care

1. Explain the purpose, benefits, and risks of the invasive arterial blood sampling procedure. Assess the patient's ability to cooperate.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret the test outcome (see Fig. 14.7). Assess, monitor, and intervene appropriately for hypoxemia and ventilatory disturbances.

   FIGURE 14.7 Arterial blood gas report of a 39-year-old male with amyotrophic lateral sclerosis (ALS). (Courtesy of Pulmonary Diagnostic Laboratory, Froedtert Hospital, Milwaukee, WI)

2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Oxygen Saturation (SO₂) This measurement is a ratio between the actual O₂ content of the hemoglobin and the potential maximum O₂ carrying capacity of the hemoglobin. The SO₂ is a percentage indicating the relationship between O₂ and hemoglobin; it does not indicate the O₂ content. The maximum amount of O₂ that can be combined with hemoglobin is called the oxygen capacity. The combined measurements of SO₂, PO₂, and hemoglobin (Hb) indicate the amount of O₂ available to tissues (tissue oxygenation). Pulse oximetry (SpO₂) (Fig. 14.6) is a noninvasive technique that permits continuous real-time monitoring and trending of arterial oxygen saturation. However, it cannot differentiate carboxyhemoglobin (COHb). As a result, the SpO₂ is generally higher than the actual arterial oxygen saturation (SaO₂) by the amount of COHb and a more direct measurement involves taking an arterial blood sample and measuring with a blood gas analyzer.

   FIGURE 14.6 Pulse oximeter. (Source: Nonin Medical, Inc, Plymouth, MN, USA)

Reference Values

Normal SaO₂ (arterial blood): >95% (>0.95) SvO₂ (mixed venous blood): 70%–75% (0.70–0.75) SaO₂ (arterial) in newborns: 40%–90% (0.40–0.90)

Procedure
1. Obtain an arterial blood sample (see page 929 for arterial and Chapter 2 for venous). Two methods are used for determining SO$_2$:
   a. **Direct method:** Introduce the blood sample into hemoximeter, a spectrophotometric device for direct determination of SO$_2$.
   b. **Calculated method:** Calculate SO$_2$ from oxygen content (the volume of O$_2$ actually combined with hemoglobin) and oxygen capacity (the volume of O$_2$ to which hemoglobin could combine). Both of these values are expressed as volume percentages (vol%), or milliliters per deciliter of blood. Use the following formula:
   \[
   \text{SO}_2 = 100 \times \frac{\text{O}_2 \text{ content}}{\text{O}_2 \text{ capacity}}
   \]

2. **Pulse oximetry:** A small, clip-like sensor is placed on a digit over the fingernail (or toenail or earlobe, if necessary). The instrument, using transmitted light waves (in the infrared spectrum and sensors, determines SO$_2$ noninvasively and is referred to as the SpO$_2$.

**Limitations**

1. SO$_2$ measures only the percentage of oxygen being carried by hemoglobin; it does not reveal the actual amount of oxygen available to the tissues (oxygen content).
2. Pulse oximetry equipment evaluates pulsatile blood flow. Many factors can interfere with the ability to measure flow:
   a. Digit motion
   b. A decrease in blood flow to the digit (e.g., cool extremity, decreased peripheral pulses, vasoconstriction, nail bed thickening, ambient light, digit malformation, vasoconstrictive drugs, localized obstruction)
   c. Decreased hemoglobin (anemia) or abnormal hemoglobin (COHb)
   d. Pulse rate and rhythm

**Interfering Factors** Recent smoking or exposure to close second-hand smoke or to CO can increase the level of COHb, as can use of certain paint and varnish-type stripping agents, especially when they are applied in closed or poorly ventilated areas. The effect is to decrease the SaO$_2$ with little or no affect on the PaO$_2$.

**Clinical Implications**

1. Abnormal results occur in pulmonary diseases involving cyanosis and erythrocytosis.
2. Abnormal results occur with venous-to-arterial shunts.
3. Values are abnormal in Rh incompatibility caused by blocking antibodies.
4. Values usually are normal in polycythemic vera.
5. Values are decreased in ventilation-perfusion mismatching.
6. Values decrease with age.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose, benefits, and risks of invasive arterial blood sampling. Assess the patient's ability to comply with the procedure.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Care**

1. Interpret test outcomes. Assess, monitor, and intervene appropriately for bleeding at puncture site and for hypoxemia or other respiratory dysfunctions.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Oxygen Content (CO$_2$)**

The actual amount of O$_2$ in the blood is termed the oxygen content (CO$_2$). Blood can contain less O$_2$ than it is capable of carrying. About 98% of all O$_2$ delivered to the tissues is transported in chemical combination with hemoglobin. One gram of hemoglobin is capable of combining with 1.34 mL of O$_2$, whereas 100 mL of blood plasma can carry a maximum of only 0.3 mL of O$_2$ (under normoxic conditions or atmospheric conditions). The CO$_2$ measurement is determined mathematically.

**Reference Values**

- **Normal** CaO$_2$ (arterial blood): 15–22 vol% or 15–22 mL/dL of blood (6.6–9.7 mmol/L) CvO$_2$ (venous blood): 11–16 vol% or 11–16 mL/dL of blood (4.9–7.1 mmol/L)

**Procedure**

1. Obtain an arterial or venous blood sample.
2. Measure the SO$_2$, PO$_2$, and hemoglobin concentration (Hb).
3. Use the following formulas for calculating O$_2$ content:
   \[
   \text{CaO}_2 = 1.34 (\text{SaO}_2 \times \text{Hb}) + 0.003 (\text{PaO}_2)
   \]
   \[
   \text{CvO}_2 = 1.34 (\text{SvO}_2 \times \text{Hb}) + 0.003 (\text{PvO}_2)
   \]

**NOTE**

0.003 = Bunsen solubility for oxygen in the blood.
Clinical Implications: Decreased CaO₂ is associated with:

1. COPD
2. Postoperative respiratory complications
3. Flail chest
4. Kyphoscoliosis
5. Neuromuscular impairment
6. Obesity-caused hypoventilation
7. Anemia

Interventions
Pretest Patient Care

1. Explain the purpose, benefits, and risks of invasive arterial blood sampling (see page 928).
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome. Assess, monitor, and intervene appropriately for bleeding at the puncture site and for hypoxemia or ventilatory disturbances.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Partial Pressure of Oxygen (PO₂): Oxygen is carried in the blood in two forms: dissolved in plasma (<2%) and combined with hemoglobin (98%). The partial pressure of a gas determines the force it exerts in attempting to diffuse through the pulmonary membrane. The PO₂ reflects the amount of O₂ passing from the pulmonary alveoli into the blood; it is directly influenced by the fraction of inspired oxygen (FIO₂). This test measures the pressure exerted by the O₂ dissolved in the plasma. It evaluates the ability of the lungs to oxygenate the blood and is used to assess the effectiveness of oxygen therapy. The PO₂ indicates the ability of the lungs to diffuse O₂ across the alveolar membrane into the circulating blood.

Reference Values
Normal PaO₂ (arterial blood): >80 mm Hg (>10.6 kPa) PvO₂ (venous blood): 30–40 mm Hg (4.0–5.3 kPa)

Procedure

1. Obtain an arterial (or venous, if requested) blood sample (see page 929 for arterial and Chapter 2 for venous).
2. Introduce a small amount of this blood into a blood gas analyzing instrument (see Fig. 14.5), and measure the O₂ tension with a polarographic electrode (developed by Leland Clark, sometimes referred to as the Clark electrode).

Clinical Implications

1. Increased PaO₂ is associated with:
   a. Polycythemia
   b. Increased FIO₂
   c. Hyperventilation
2. Decreased PaO₂ is associated with:
   a. Anemias
   b. Cardiac decompensation
   c. Insufficient atmospheric O₂
   d. Intracardiac shunts
   e. COPD
   f. Restrictive pulmonary disease
   g. Hypoventilation caused by neuromuscular disease
3. Decreased PaO₂ with normal or decreased PACO₂ is associated with:
   a. Diffuse interstitial pulmonary infiltration
   b. Pulmonary edema
   c. Pulmonary embolism
   d. Postoperative extracorporeal circulation

Clinical Alert

In persons with COPD, ventilatory efforts are stimulated by the hypoxic state, whereas for a healthy person, the respiratory stimulus is the buildup of CO₂. Although supplemental oxygen increases the PaO₂ in such patients, it can also result in less effective breathing because ventilatory efforts are no longer stimulated. The administration of oxygen “knocks out” this hypoxic drive, and CO₂ retention results.

Interventions
Pretest Patient Care

1. Explain the purpose, benefits, and risks of arterial blood sampling. Assess the patient’s level of cooperation and understanding.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret the test outcome (see Fig. 14.7). Assess, monitor, and intervene appropriately for bleeding at the puncture site and for respiratory or ventilatory disturbances.

2. See Chapter 1 guidelines for safe, effective, informed posttest care.

Carbon Dioxide (CO$_2$) Content; Total Carbon Dioxide (TCO$_2$) In normal blood plasma, >95% of the total CO$_2$ content (TCO$_2$) is contributed by bicarbonate ion (HCO$_3^-$), which is regulated by the kidneys. The other 5% is contributed by the dissolved CO$_2$ gas and by carbonic acid (H$_2$CO$_3$). Dissolved CO$_2$ gas, which is regulated by the lungs, therefore contributes little to the TCO$_2$, and the TCO$_2$ gives little information about the lungs. The HCO$_3^-$ in the extracellular spaces exists first as CO$_2$, then as H$_2$CO$_3$; later, much of it is changed to sodium bicarbonate (NaHCO$_3$) by the buffers in the plasma and erythrocytes. This test is a general measure of the alkalinity or acidity of venous, arterial, or capillary blood. This test measures the CO$_2$ contributions from dissolved CO$_2$ gas, total H$_2$CO$_3$, HCO$_3^-$, and carbaminohemoglobin (CO$_2$HHb).

**Reference Values**

**Normal** 23–30 mEq/L or 23–30 mmol/L

**Procedure**

1. Collect a venous or arterial blood sample of 5 mL in a heparinized syringe.
2. Measure the sample by a blood gas analyzer. If the collected blood sample cannot be studied immediately, place the syringe in an iced container.
3. Use the following formula: TCO$_2$ = HCO$_3^-$ + H$_2$CO$_3$

**Clinical Implications**

1. **Increased** TCO$_2$ occurs in:
   a. Severe vomiting
   b. Emphysema
   c. Aldosteronism
   d. Use of mercurial diuretics
2. **Decreased** TCO$_2$ occurs in:
   a. Severe diarrhea
   b. Starvation
   c. Acute renal failure
   d. Salicylate toxicity
   e. Diabetic acidosis
   f. Use of chlorothiazide diuretics

**NOTE**

In diabetic acidosis, the supply of ketoacids exceeds the demands of the cell. Blood plasma acids rise. Blood plasma HCO$_3^-$ decreases because it is used to neutralize these excess acids.

**Table 14.4** presents the changes in pH, HCO$_3^-$, and PaO$_2$ that occur in various ventilatory disturbances and acid-base imbalances.

**Table 14.4 Summary of Ventilatory and Acid-Base Changes in Four Underlying Conditions of Acid-Base Imbalance**

<table>
<thead>
<tr>
<th>Form of Disturbance</th>
<th>pHa</th>
<th>Bicarbonate (HCO PaCO$_2$)</th>
<th>Occurrence</th>
</tr>
</thead>
<tbody>
<tr>
<td>RESPIRATORY ACIDOSIS</td>
<td>Decrease Normal</td>
<td>Increase Depression of respiratory centers Drug overdose Barbiturate toxicity Use of anesthetics Interference with mechanical function of the thoracic cage Deformity of thoracic cage Kyphoscoliosis Airway obstruction Extrathoracic tumors Asthma Bronchitis Emphysema</td>
<td></td>
</tr>
</tbody>
</table>
Circulatory disorders
Congestive heart failure
Shock

Chronic: compensated via renal reabsorption of the bicarbonate ion

RESPIRATORY ALKALOSIS
Acute: caused by increased alveolar ventilation and excessive blowing off of CO$_2$ and water

Normal Increase Increase

Hyperventilation

Hysteria
Lack of oxygen
Toxic stimulation of the respiratory centers
High fever
Cerebral hemorrhage
Excessive artificial respiration
Salicylates

Chronic: compensated via glomerular filtration of the bicarbonate ion

NONRESPIRATORY OR METABOLIC ACIDOSIS
Acute: caused by accumulation of fixed body acids or loss of bicarbonate from the extracellular fluid

Decrease Decrease Normal Acid gain

Renal failure
Diabetic ketoacidosis
Lactic acidosis
Anaerobic metabolism
Hypoxia
Base loss
Diarrhea
Renal tubular acidosis

Chronic: compensated via hyperventilation through stimulation of central chemoreceptors

NONRESPIRATORY OR METABOLIC ALKALOSIS
Acute: caused by loss of fixed body acids or gain in bicarbonate in extracellular fluid

Increase Increase Normal Acid loss

Loss of gastric juice
Vomiting
Potassium or chloride depletion
Base gain
Excessive bicarbonate or lactate administration

Chronic: compensated via hypoventilation

*Although these four basic imbalances occur individually, a combination of two or more is observed more frequently. These disturbances may have an antagonistic or a synergistic effect on each other.

†Uncompensated disturbances are referred to as acute and compensated ones as chronic. Compensation occurs via the mechanism not involved. Compensation is most efficient in respiratory acidosis.

‡Acid-base disturbances force kidney and lungs to compensate for changes in pH. Hyperventilation or hypoventilation can restore pH to normal within 15 minutes; the kidney, however, can take 2 to 3 days to compensate.

§The degree of hypoventilation is precisely related to the degree of hypobicarbonatemia. For each 1 mEq/L fall in bicarbonate, P CO$_2$ falls by 1 to 1.3 mm Hg. A close mathematical relationship prevails between bicarbonate and P CO$_2$; their ratio defines the prevailing hydrogen on concentration. For this reason, the steady-state P CO$_2$ in simple metabolic acidosis is equal to the last two digits of the pH. Failure of the P CO$_2$ to reach predicted levels defines the presence of superimposed respiratory acidosis or alkalosis.

||Decreases in Pa O$_2$ are interpreted separately and are referred to as hypoxemia.
Clinical Alert

1. A double use of the term CO$_2$ is one of the main reasons why understanding of acid-base problems may be difficult. Use the terms CO$_2$ content and CO$_2$ gas to avoid confusion. Remember the following: CO$_2$ content (ie, TCO$_2$) is mainly bicarbonate and a base. It is a solution and is regulated by the kidneys. CO$_2$ gas is mainly acid. It is regulated by the lungs.

2. The panic value for CO$_2$ content is <6.0 mEq/L (<6.0 mmol/L); it usually is associated with severe metabolic acidosis, with the pH often <7.1. This is a life-threatening situation.

Interfering Factors A number of drugs can either increase or decrease TCO$_2$.

Interventions

Pretest Patient Care

1. Explain the purpose, benefits, and risks of arterial blood sampling. Assess the patient’s ability to comply with the procedure.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes. Assess, monitor, and intervene appropriately for acid-base imbalances.
2. Monitor and intervene for bleeding at the puncture site and for respiratory or ventilatory disturbances.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Blood pH

The pH is the negative logarithm of the hydrogen ion concentration in the blood. The sources of hydrogen ions are volatile acids, which can vary between a liquid and a gaseous state, and nonvolatile acids, which cannot be volatilized but remain fixed (eg, dietary acids, lactic acids, ketoacids).

NOTE

A pH value of 7 is neutral; acidity increases as the pH falls from 7 to 1, and alkalinity increases as the pH rises from 7 to 14. Limits of pH compatible with life fall within the range of 6.9 to 7.8.

Blood pH measures the body’s chemical balance and represents a ratio of acids to bases. It is also an indicator of the degree to which the body is adjusting to dysfunctions by means of its buffering systems. It is one of the best ways to determine whether the body is too acidic or too alkaline and is an indicator of the patient’s metabolic and respiratory status. The acid-base balance in the extracellular fluid is extremely delicate and intricate and must be kept within the very narrow range of 7.35 to 7.45 (slightly alkaline). Values < 7.35 indicate an acid state, whereas pH values > 7.45 indicate an alkaline state.

Reference Values

Normal pH$_{a}$ (arterial blood): 7.35–7.45 pH$_{v}$ (venous blood): 7.31–7.41

Procedure

1. Obtain an arterial (or venous if requested) blood sample.
2. Use one of the following two methods to determine the pH:
   a. Direct method: Analyze a small amount of blood by a blood gas machine (see Fig. 14.5); measure the pH by a modified Severinghaus electrode.
   b. Indirect method: Solve the Henderson-Hasselbalch equation for the pH of a buffer system. In this equation, pK is the negative logarithm of the acid dissociation constant (the pH at which the associated and unassociated forms of an acid exist in equal concentrations). [A$^-$] is the concentration of the ionized form (in this case HCO$_3^-$, the major blood base), and [HA] is the concentration of the free acid (in this case H$_2$CO$_3$, the major blood acid), in milliequivalents per liter.

   \[
   \text{pH} = \text{pK} + \log \frac{[A^-]}{[HA]}
   \]

   \[
   \text{pH} = \text{pK} + \log \frac{[\text{HCO}_3^-]}{[\text{H}_2\text{CO}_3]}
   \]

   \[
   \text{pH} = 6.1 = \log \frac{[\text{HCO}_3^-]}{0.03(\text{PaCO}_3)}
   \]

Clinical Implications

1. Generally speaking, the pH is decreased in acidemia (acidosis) because of increased formation of acids, and pH is increased in alkalemia (alkalosis) because of a loss of acids.
2. When interpreting an acid-base abnormality, certain steps should be followed:
   a. Check the pH to determine whether an acid or an alkaline state exists.
   b. Check the PCO$_2$ to determine whether a respiratory acidosis or alkalosis is present. (PCO$_2$ is the breathing component.)
   c. Check the HCO$_3^-$ concentration to determine whether a metabolic acidosis or alkalosis is present. (HCO$_3^-$ is the renal component.)
3. See Table 14.4 for a more complete explanation of the changes occurring in acute and chronic respiratory and
metabolic acidosis and alkalosis.

4. Metabolic acidemia (acidosis) occurs in:
   a. Renal failure
   b. Ketoacidosis in diabetes and starvation
   c. Lactic acidosis
   d. Strenuous exercise
   e. Severe diarrhea

5. Metabolic alkalemia (alkalosis) occurs in:
   a. Hypokalemia
   b. Hypochloremia
   c. Gastric suction or vomiting
   d. Massive doses of steroids
   e. Sodium bicarbonate administration
   f. Aspirin intoxication

6. Respiratory alkalemia (alkalosis) occurs in:
   a. Acute pulmonary disease
   b. Myocardial infarction
   c. Chronic and acute heart failure
   d. Adult cystic fibrosis
   e. Third trimester of pregnancy and during labor and delivery
   f. Anxiety, neuroses, psychoses
   g. Pain
   h. Central nervous system diseases
   i. Anemia
   j. Carbon monoxide poisoning
   k. Acute pulmonary embolus
   l. Shock

7. Respiratory acidemia (acidosis) occurs in:
   a. Acute or chronic respiratory failure
   b. Ventilatory failure
   c. Neuromuscular depression
   d. Obesity
   e. Pulmonary edema
   f. Cardiopulmonary arrest

**Interfering Factors** A number of drugs may alter the components of acid-base balance. See Appendix J.

**Clinical Alert**

1. Ventilatory failure is a medical emergency. Aggressive and supportive measures must be taken immediately.
2. Rate and depth of respirations may give a clue to blood pH. Acidosis usually increases respirations; this is the body’s way of adjusting once the state is established.
3. Respiratory alkalosis may reflect hyperventilation in response to treatment for hypoxemia; however, correction of hypoxemia is essential.
4. Metabolic alkalosis, which is compensated through hypoventilation, may produce hypoxemia.

**Interventions**

**Pretest Patient Care**

1. Explain the purpose, benefits, and risks of invasive blood sampling.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome (see Fig. 14.7). Assess, monitor, and intervene appropriately for metabolic and respiratory acidosis and alkalosis (see Table 14.4).
2. Frequently observe the arterial puncture site for bleeding (see page 929). Be prepared to initiate proper interventions in the event of life-threatening situations.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Base Excess or Deficit** This test quantifies the patient's total base excess or deficit so that clinical treatment of acid-base disturbances (specifically those that are nonrespiratory in nature) can be initiated. It is also referred to as the whole blood buffer base and is the sum of the concentration of buffer anions (in milliequivalents per liter) contained in whole blood. These buffer anions are the bicarbonate ion (HCO₃⁻) present in plasma erythrocytes, and the hemoglobin, plasma proteins, and phosphates in plasma and red blood cells. The total quantity of buffer anions is 45–50 mEq/L (45–50 mmol/L), or about twice that of HCO₃⁻ alone (22–26 mEq/L or 22–26 mmol/L). Therefore, the quantity of HCO₃⁻ ions accounts for only about half of the total buffering capacity of the blood. The base excess or deficit measurement provides a more complete picture of the buffering that is taking place and is a critical index of nonrespiratory versus respiratory changes in acid-base balance.

**Reference Values**

**Normal** Normal values are between ± 2 mEq/L (±2 mmol/L). A positive value indicates a base excess (ie, nonvolatile acid deficit). A negative value indicates a base deficit (ie, nonvolatile acid excess).

**Procedure**

1. Make calculations from the measurements of pH, PaCO₂, and the hematocrit.
2. Plot these values on a nomogram, and read the base excess or deficit.

Clinical Implications

1. A negative value (less than -2 mEq/L or -2 mmol/L) reflects a nonrespiratory or metabolic disturbance or true base deficit, or a nonvolatile acid accumulation caused by:
   a. Dietary intake of organic and inorganic acids
   b. Lactic acid
   c. Ketoacidosis
2. A positive value (higher than +2 mEq/L or +2 mmol/L) reflects a nonvolatile acid deficit or true base excess.

Anion Gap (AG) This test measures the difference between the sum of the sodium (Na⁺) and potassium (K⁺) ion concentrations (the measured cations) and the sum of the chloride (Cl⁻) and bicarbonate (HCO₃⁻) concentrations (the measured anions). This difference reflects the concentrations of other anions that are present in the extracellular fluid but are not routinely measured, the components of which include phosphates, sulfates, ketone bodies, lactic acid, and proteins. Increased amounts of these unmeasured anions are produced in the acidic state. Primary hypobicarbonatemia is brought about by any combination of three mechanisms: (1) overproduction of acids, which causes replacement of NaHCO₃ by the Na⁺ salt of the offending acid (eg, lactate replaces HCO₃⁻ in lactic acidosis); (2) loss of NaHCO₃ through diarrhea along with renal retention of dietary NaCl, which causes hyperchloremic metabolic acidosis; and (3) generalized renal failure or specific forms of renal tubular acidosis, which cause retention of acids that are normally produced by intermediary metabolism or by urinary excretion of alkali (Table 14.5).

Table 14.5 Subclassification of Anion Gap Metabolic Acidosis (Hypobicarbonatemia) into High- and Low-Potassium Forms.*

<table>
<thead>
<tr>
<th>High-Potassium Form</th>
<th>Low-Potassium Form</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acidifying agents</td>
<td>Diarrhea</td>
</tr>
<tr>
<td>Mineralocorticoid deficiency</td>
<td>Ureteral sigmoidostomy and malfunctioning ileostomy</td>
</tr>
<tr>
<td>Renal diseases such as systemic lupus erythematosus, interstitial nephritis, amyloidosis, hydronephrosis, and sickle cell nephropathy</td>
<td>Renal tubular acidosis, both proximal and distal</td>
</tr>
<tr>
<td>Early nonspecific renal failure</td>
<td></td>
</tr>
</tbody>
</table>

*All metabolic acids can be classified on the basis of how they affect the anion gap.

Hyperbicarbonatemia with sustained increases in HCO₃⁻ levels is brought about by a source of new alkali or by the presence of factors that stimulate renal retention of excess HCO₃⁻. These mechanisms include excessive gastrointestinal loss of acid, exogenous alkali in persons whose kidneys avidly retain NaHCO₃, and renal synthesis of HCO₃⁻ in excess of daily consumption. Other pathophysiologic factors that affect renal reabsorption of >25 mEq/L (>25 mmol/L) of HCO₃⁻ and contribute to sustained hyperbicarbonatemia include extracellular fluid volume contraction, hypercapnia, hypokalemia, hyperaldosteronemia, and hypoparathyroidism (Table 14.6).

Table 14.6 Classification of Anion Gap Metabolic Alkalosis (Hyperbicarbonatemia) on the Basis of Urinary Excretion

<table>
<thead>
<tr>
<th>Saline-Responsive Urinary Chloride Excretion of &lt;10 mEq/day</th>
<th>Saline-Unresponsive Chloride Excretion of &lt;10 mEq/day</th>
</tr>
</thead>
<tbody>
<tr>
<td>EXCESS BODY BICARBONATE CONTENT</td>
<td></td>
</tr>
<tr>
<td>Renal alkalosis</td>
<td>Renal alkalosis—normotensive conditions</td>
</tr>
<tr>
<td>Diuretic therapy</td>
<td>Bartter's syndrome</td>
</tr>
<tr>
<td>Poorly reabsorbable anion therapy, (eg, carbenicillin, penicillin, sulfate, phosphate)</td>
<td>Severe potassium depletion</td>
</tr>
<tr>
<td>Gastrointestinal alkalosis</td>
<td>Refeeding alkalosis</td>
</tr>
<tr>
<td>Gastric alkalosis</td>
<td>Hypercalcemia and hypoparathyroidism</td>
</tr>
<tr>
<td>Intestinal alkalosis (eg, chloride diarrhea)</td>
<td>Hypertensive conditions—endogenous mineralocorticoids</td>
</tr>
<tr>
<td>Exogenous alkali</td>
<td>Primary aldosteronism</td>
</tr>
<tr>
<td>Baking soda</td>
<td>Hyperreninism</td>
</tr>
<tr>
<td>Sodium citrate, lactate, gluconate, acetate</td>
<td>Adrenal enzyme deficiency: 11- and 17-hydroxylase</td>
</tr>
<tr>
<td>Transfusions</td>
<td>Liddle's syndrome</td>
</tr>
<tr>
<td>Antacids</td>
<td>Exogenous mineralocorticoids</td>
</tr>
<tr>
<td></td>
<td>Licorice</td>
</tr>
<tr>
<td></td>
<td>Carbenoxolone</td>
</tr>
<tr>
<td></td>
<td>Chewing tobacco</td>
</tr>
</tbody>
</table>

NORMAL BODY BICARBONATE CONTENT

Contraction alkalosis—urinary loss of NaCl and water without bicarbonate loss causes extracellular fluid contraction around an unchanged body content of alkali, resulting in hyperbicarbonatemia (especially important in persons with edema and persons who have excess body stores of water, sodium, bicarbonate, and chloride).
Reference Values

**Normal** Normal values are between 12 ± 4 mEq/L or 12 ± mmol/L. If potassium concentration is used in the calculation, the normal value is 16 ± 4 mEq/L or 16 ± 4 mmol/L.

**Procedure**

1. Obtain this measurement by calculating the difference between the measured serum cation concentrations (either with or without K⁺) and the measured serum anion concentrations.
2. Use the following formulas:

   \[
   AG = ([Na^+] + [K^+]) - ([Cl^-] + [HCO_3^-])
   \]

   or

   \[
   AG = [Na^+] - ([Cl^-] + [HCO_3^-])
   \]

**Clinical Implications**

1. An anion gap (AG) occurs in acidosis that is caused by excess metabolic acids and excess serum chloride levels. If there is no change in sodium content, anions such as phosphates, sulfates, and organic acids increase the AG because they replace bicarbonate.
2. Increased AG is associated with an increase in metabolic acid when there is excessive production of metabolic acids, as in:
   a. Alcoholic ketoacidosis
   b. Diabetic ketoacidosis
   c. Fasting and starvation
   d. Ketogenic diets
   e. Lactic acidosis
   f. Poisoning by salicylate, ethylene glycol (antifreeze), methanol, or propyl alcohol
3. Increased AG is also associated with decreased loss of metabolic acids, as in renal failure. In the absence of renal failure or intoxication with drugs or toxins, an increase in AG is assumed to be caused by ketoacidosis or lactate accumulation. Anion gap includes the determination of three gaps of toxicology (influence of drugs and heavy metals): (1) anion = type A lactic acidosis due to tissue hypoxia; (2) osmolar gap; and (3) oxygen saturation gap. A list of drugs and toxic substances that cause increased anion gap (>12 mEq/L or >12 mmol/L) including the following: nonacidotic: carbenicillin and sodium salts; metabolic acidosis: acetaminophen (ingestion > 75–100 g), acetazolamide, aluminum phosphate, amiloride, 4-aminopyridine, ammonium chloride, ascorbic acid, benzalkonium chloride, bialaphos, 2-butoxyethanol, carbon monoxide, centrimonium bromide, chloramphenicol, clozapine, cobalt, colchicine, cyanide, dapsone, dimethyl sulfate, dinitrophenol, endosulfan, epinephrine (IV overdose), ethanol, ethylene dibromide, ethylene glycol, fenoprofen, fluoroacetate, formaldehyde, fructose (IV), funnel web spiders, glycol ethers, glyphosate, hydrogen sulfide, ibuprofen (ingestion > 300 mg/kg), inorganic acid, iodine, iron, isoniazid, ketamine, ketoprofen, lime sulfur, margosa oil, metaldehyde, metformin, methanol, methenamine mandelate, misoprostol, monochloracetic acid, naltidixic acid, naproxen, nefopam, niacin, papaverine, paraldehyde, pentyrovaloyl, pentaborane, pentachlorophenol, phenelzine, phenformin (off the market), phenol, phenylbutazone, phosphoric acid, polyethylene glycol (low molecular weight), propofol, propylene glycol, salicylates, sodium azide, sorbitol (IV), strychnine, sublimed sulfur, sulfihame, surfactant herbicide, tetracycline (outdated), tienilic acid, tolueue, tranylcypromine, vacor, valproic acid, verapamil, zidovudine (chronic use > 6 months), and zinc phosphate. Toxins that cause osmolar gap > 10 mOsm from baseline include ethanol, ethylene glycol, glycerol, hypermagnesemia (>9.5 mEq/L or >9.5 mmol/L), isopropanol (acetone), iodide (questionable), mannitol, methanol, and sorbitol. Drugs and toxins that cause decreased anion gap (<6 mEq/L or <6 mmol/L) include the following: acidosis: acetazolamide, amiloride, ammonium chloride, amphotericin B, bromide, fialuridine (FLAU), iodide, kombucha tea, lithium, polymyxin B, spiranolactone, sulindac, tolueue, and tromethamine. Toxins that cause an oxygen saturation gap (>5% difference between measured and calculated value) include carbon monoxide, cyanide (questionable), hydrogen sulfide (possible), methemoglobin, and nitrates.
4. Increased bicarbonate loss with a normal AG is associated with
   a. Decreased renal losses, as in
      1. Renal tubular acidosis
      2. Use of acetozolamide
   b. Increased chloride levels, as in
      1. Altered chloride reabsorption by the kidney
      2. Parenteral hyperalimentation
      3. Administration of sodium chloride and ammonium chloride
   c. Loss of intestinal secretions, as in
      1. Diarrhea
      2. Intestinal suction or fistula
      3. Biliary fistula
5. Low AG is associated with
   a. Multiple myeloma
   b. Hyponatremia caused by viscous serum
   c. Bromide ingestion (hyperchloremia)
1. Interpret test outcomes and assess and monitor appropriately for acid-base disturbances.
2. The AG may provide evidence of a mixed rather than a simple acid-base disturbance.
3. Lactic acidosis should be considered in any metabolic acidosis with increased AG of >15 mEq/L (>15 mmol/L).

**Lactic Acid**

Lactate is a product of carbohydrate metabolism. Lactic acid is produced during periods of anaerobic metabolism when cells do not receive adequate oxygen to allow conversion of fuel sources to CO$_2$ and water. Lactic acid accumulates because of excess production of lactate and decreased removal of lactic acid from blood by the liver. This measurement contributes to the knowledge of acid-base volume and is used to detect lactic acidosis in persons with underlying risk factors such as cardiovascular or renal disease that predispose them to this imbalance. Lactate is elevated in a variety of conditions in which hypoxia occurs and in liver disease. Lactic acidosis can occur in both diabetic and nondiabetic patients. It is often fatal.

**Reference Values**

**Normal** In venous blood: 0.5–2.2 mEq/L (0.5–2.2 mmol/L) In arterial blood: 0.5–1.6 mEq/L (0.5–1.6 mmol/L)

**Procedure**

1. Obtain a venous or arterial blood sample of at least 5 mL.
2. Take the specimen to the laboratory immediately for analysis.

**Clinical Implications**

1. Values are increased in:
   a. Lactic acidosis
   b. Cardiac failure
   c. Pulmonary failure
   d. Hemorrhage
   e. Diabetes
   f. Shock
   g. Liver disease
2. Lactic acidosis can be distinguished from ketoacidosis by the absence of severe ketosis and hyperglycemia in this state.

**Interfering Factors**

Lactic acid levels normally rise during strenuous exercise, when blood flow and oxygen cannot keep pace with the increased needs of exercising muscle.

**Clinical Alert**

An unexplained decrease in pH associated with a hypoxia-producing condition is reason to suspect lactic acidosis.

**Electrolyte Tests**

Electrolytes (ions) are critical for cellular reactions. These electrolytes provide the necessary inorganic chemicals for a variety of cellular functions (eg, nerve impulse transmission, muscular contraction, water balance). Typically, the concentration of cations (positively charged electrolytes), eg, Na$^+$, K$^+$, Ca$^{++}$ and Mg$^+$, is higher in the plasma than in the interstitial fluid owing to the Donnan effect (plasma proteins have a net negative charge), whereas the anions (negatively charged), eg, Cl$^-$, HPO$_4^{--}$, tend to be higher in the interstitial fluid than the plasma.

**Calcium (Ca$^{++}$)**

The bulk of body calcium (99%) is stored in the skeleton and teeth, which act as huge reservoirs for maintaining blood levels of calcium. About 50% of blood calcium is ionized; the rest is protein bound. Only ionized calcium can be used by the body in such vital processes as muscular contraction, cardiac function, transmission of nerve impulses, and blood clotting. The amount of protein in the blood also affects calcium levels because 50% of blood calcium is protein bound. Thus, a decrease in serum albumin will result in a decrease in total serum calcium. The decrease, however, does not alter the concentration of the ionized form of calcium. Measurements of ionized calcium are done during open-heart surgeries, liver transplantations, and other operations in which large volumes of blood anticoagulated with citrate are given. These tests are also used to monitor renal disease, renal transplantation, hemodialysis, hyperparathyroidism, hypoparathyroidism, pancreatitis, and malignancy. Parathyroid hormone (PTH), calcitonin, vitamin D, estrogens, androgens, carbohydrates, and lactose all are factors that influence calcium levels. This test measures the concentration of total and ionized calcium in the blood to reflect parathyroid function, calcium metabolism, and malignancy activity.
Clinical Alert

Hyperparathyroidism and cancer are the most common causes of hypercalcemia. Hypoalbuminemia is the most common cause of decreased total calcium.

Reference Values

Normal See Table 14.7.

Table 14.7 Values for Calcium

<table>
<thead>
<tr>
<th>Age</th>
<th>Total Calcium</th>
<th>Ionized Calcium</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mg/dL</td>
<td>mmol/L</td>
</tr>
<tr>
<td>0–10 d</td>
<td>7.6–10.4</td>
<td>1.90–2.60</td>
</tr>
<tr>
<td>10 d–3 yr</td>
<td>6.7–9.8</td>
<td>2.24–2.75</td>
</tr>
<tr>
<td>3–9 yr</td>
<td>8.8–10.1</td>
<td>2.20–2.70</td>
</tr>
<tr>
<td>4–11 yr</td>
<td>8.9–10.1</td>
<td>2.30–2.70</td>
</tr>
<tr>
<td>11–13 yr</td>
<td>8.8–10.6</td>
<td>2.20–2.65</td>
</tr>
<tr>
<td>13–15 yr</td>
<td>9.2–10.7</td>
<td>2.30–2.55</td>
</tr>
<tr>
<td>15–18 yr</td>
<td>8.4–10.7</td>
<td>2.10–2.67</td>
</tr>
<tr>
<td>Adult</td>
<td>8.8–10.4</td>
<td>2.20–2.60</td>
</tr>
</tbody>
</table>

Procedure

1. Obtain a 5-mL venous blood sample; this will provide sufficient serum for this test.
2. Observe standard/universal precautions. Be aware that heparinized samples are preferred for ionized calcium studies.
3. Place specimens on ice, keep tightly capped, and deliver immediately to the laboratory.

Clinical Alert

Citrated ethylenediaminetetraacetic acid (EDTA) and oxalated blood give falsely low values and should not be used in the syringe.

Clinical Implications

1. Normal levels of total blood calcium, combined with other findings, indicate the following conditions:
   a. Normal calcium levels with overall normal results in other tests indicate no problems with calcium metabolism.
   b. Normal calcium and abnormal phosphorus values indicate impaired calcium absorption owing to alteration of PTH activity or secretion (eg, in rickets, the calcium level may be normal or slightly lowered and the phosphorus level depressed).
   c. Normal calcium and elevated blood urea nitrogen (BUN) levels indicate the following:
      1. Possible secondary hyperparathyroidism: initially, lowered serum calcium results from uremia and acidosis. The reduced calcium level stimulates the parathyroid to release PTH, which acts on bone to release more calcium.
      2. Possible primary hyperparathyroidism: excessive amounts of PTH cause elevation in calcium levels, but secondary kidney disease causes retention of phosphate and concomitant lower calcium levels.
   d. Normal calcium and decreased serum albumin indicates hypercalcemia. Normally, a decrease in calcium is associated with a decrease in albumin.

2. Hypercalcemia (increased total calcium levels [>12 mg/dL or >3 mmol/L]) is caused by or associated with the following conditions:
   a. Hyperparathyroidism due to parathyroid adenoma, hyperplasia of parathyroid glands, or associated hypophosphatemia
   b. Cancer (PTH-producing tumors)
      1. Metastatic bone cancers; cancers of lung, breast, thyroid, kidney, liver, and pancreas
      3. Multiple myeloma with extensive bone destruction, Burkitt’s lymphoma
      4. Primary squamous cell carcinoma of lung, neck, and head
   c. Granulomatous disease (eg, tuberculosis, sarcoidosis)
   d. Thyroid toxicosis
   e. Paget’s disease of bone (also accompanied by high levels of alkaline phosphatase)
   f. Idiopathic hypercalcemia of infancy
   g. Bone fractures combined with bed rest, prolonged immobilization
   h. Excessive intake of vitamin D, milk, antacids
   i. Renal transplantation
   j. Milk-alkali syndrome (Burnett’s syndrome)

3. Hypocalcemia (decreased total calcium levels [<4.0 mg/dL or <1.0 mmol/L]) are commonly caused by or associated with the following conditions:
   a. Pseudohypocalcemia, which reflects reduced albumin levels. The reduced protein is responsible for the low calcium level because 50% of the calcium total is protein bound.
   b. Hypoparathyroidism due to surgical removal of parathyroid glands, irradiation, hypomagnesemia, gastrointestinal (GI) disorders, or renal wasting. The primary form is very rare.
c. Hyperphosphatemia due to renal failure, laxative intake, or cytotoxic drugs
d. Malabsorption due to sprue, celiac disease, or pancreatic dysfunction (fatty acids combine with calcium and are precipitated and excreted in the feces)
e. Acute pancreatitis
f. Alkalosis (calcium ions become bound to protein)
g. Osteomalacia (advanced)
h. Renal failure
i. Vitamin D deficiency, rickets
j. Malnutrition (inadequate nutrition)
k. Alcoholism, hepatic cirrhosis

4. **Increased ionized calcium levels** occur in the following conditions:
   a. Hyperparathyroidism
   b. Ectopic PTH-producing tumors
   c. Increased vitamin D intake
d. Malignancies

5. **Decreased ionized calcium levels** occur in the following conditions:
   a. Hyperventilation to control increased intracranial pressure (total Ca **++** may be normal)
b. Administration of bicarbonate to control metabolic acidosis
c. Acute pancreatitis (e.g., diabetic acidosis, sepsis)
d. Hypoparathyroidism
e. Vitamin D deficiency
f. Magnesium deficiency
g. Multiple organ failure
h. Toxic shock syndrome

---

**Clinical Alert**

1. Severe hypocalcemia occurs when serum calcium is <3 mg/dL (<0.75 mmol/L).
2. Excessive IV fluids decrease albumin levels and thus decrease calcium levels. Total serum protein and albumin should be measured at the same time as calcium for proper interpretation of calcium levels. Ionized calcium is not affected by albumin levels.

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**Panic Values for Total Calcium**

<4.4 mg/dL (<1.1 mmol/L) may produce tetany and convulsions.

>13 mg/dL (>3.25 mmol/L) may cause cardiotoxicity, arrhythmias, and coma.

Rapid treatment of hypercalcemia with calcitonin solution is indicated.

**Panic Values for Ionized Calcium**

<2.0 mg/dL (<0.5 mmol/L) may produce tetany or life-threatening complications.

2.0–3.0 mg/dL (0.5–0.75 mmol/L) in cases of multiple blood transfusions (this is an indication to administer calcium)

>7.0 mg/dL (>1.75 mmol/L) may cause coma.

---

**Interfering Factors**

1. Thiazide diuretics may impair urinary calcium excretion and result in hypercalcemia (most common drug-induced factor).
2. For patients with renal insufficiency undergoing dialysis, a calcium-ion exchange resin is sometimes used for hyperkalemia. This resin may increase calcium levels.
3. Increased magnesium and phosphate uptake and excessive use of laxatives may lower blood calcium level because of increased intestinal calcium loss.
4. When decreased calcium levels are due to magnesium deficiency (as in poor bowel absorption), the administration of magnesium will correct the calcium deficiency.
5. If a patient is known to have or suspected of having a pH abnormality, a concurrent pH test with ionized calcium level should be requested.
6. Many drugs may cause increased or decreased levels of calcium. Calcium supplements taken shortly before specimen collection will cause falsely high values.
7. Elevated serum protein increases calcium; decreased protein decreases calcium.

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**Interventions**

**Pretest Patient Care**

1. Explain purpose and procedure. Encourage relaxation.
2. Be aware that tourniquet application should be as brief as possible when drawing ionized calcium to prevent venous stasis and hemolysis.
3. Ensure that calcium supplements are not taken within 8 to 12 hours before the blood sample is drawn.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Have patient resume normal activities.
2. Interpret test results and monitor appropriately for calcium abnormalities.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Chloride (Cl\(^{-}\)) Chloride, a blood electrolyte, is the major anion that exists predominantly in the extracellular spaces as part of sodium chloride or hydrochloric acid. Chloride maintains cellular integrity through its influence on osmotic pressure and acid-base and water balance. It has the reciprocal power of increasing or decreasing in concentration in response to concentrations of other anions. In metabolic acidosis, there is a reciprocal rise in chloride concentration when the bicarbonate concentration drops. Similarly, when aldosterone directly causes an increase in the reabsorption of sodium (the positive ion), the indirect effect is an increase in the absorption of chloride (the negative ion). Chlorides are excreted with cations (positive ions) during massive diuresis from any cause and are lost from the GI tract when vomiting, diarrhea, or intestinal fistulas occur. Alteration of sodium chloride level is seldom a primary problem. Measurement of chlorides is usually done for inferential value and is helpful in diagnosing disorders of acid-base and water balance. Because of the relatively high chloride concentrations in gastric juices, prolonged vomiting may lead to considerable chloride loss and lowered serum chloride levels. In an emergency, chloride is the least important electrolyte to measure. However, it is especially important in the correction of hypokalemic alkalosis. If potassium is supplied without chloride, hypokalemic alkalosis may persist.

Reference Values

| Normal  | Adults: 96–106 mEq/L or 96–106 mmol/L | Newborns: 96–113 mEq/L or 96–113 mmol/L |

Procedure

1. Obtain a 5-mL venous blood sample in a heparinized Vacutainer tube. Serum can also be used.
2. Observe standard precautions.

Clinical Implications

**NOTE**
Whenever serum chloride levels are much lower than 100 mEq/L (100 mmol/L), urinary excretion of chlorides is also low.

1. **Decreased blood chloride levels** occur in the following conditions:
   a. Severe vomiting
   b. Gastric suction
   c. Chronic respiratory acidosis
   d. Burns
   e. Metabolic alkalosis
   f. Congestive failure
   g. Addison's disease
   h. Salt-losing diseases (syndrome of inappropriate antidiuretic hormone [SIADH])
   i. Overhydration or water intoxication
   j. Acute intermittent porphyria
   k. Salt-losing nephritis
2. **Increased blood chloride levels** occur in the following conditions:
   a. Dehydration
   b. Cushing's syndrome
   c. Hyperventilation, which causes respiratory alkalosis
   d. Metabolic acidosis with prolonged diarrhea
   e. Hyperparathyroidism (primary)
   f. Select kidney disorders (eg, renal tubular acidosis)
   g. Diabetes insipidus
   h. Salicylate intoxication
   i. Head injury with hypothalamic damage
   j. Eclampsia

Interfering Factors

1. The plasma chloride concentration in infants is usually higher than that in children and adults.
2. Certain drugs may alter chloride levels.
3. Increases are associated with excessive IV saline infusions.

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**Clinical Alert**

**Panic Values for Serum Chloride**

<70 or >120 mEq/L (<70 or >120 mmol/L)

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and blood collection procedure.
2. If possible, ensure that the patient fasts at least 8 to 12 hours before the test.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Resume normal activities and diet.
2. Interpret test results and monitor appropriately.
3. If an electrolyte disorder is suspected, daily weight and accurate fluid intake and output should be recorded.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Phosphate (P); Inorganic Phosphorus (PO₄)

Of the human body's total phosphorus content, 85% is combined with calcium in the bone, and the remainder resides within the cells. Most of the phosphorus in the blood exists as phosphates or esters. Phosphate is required for generation of bony tissue and functions in the metabolism of glucose and lipids, in the maintenance of acid-base balance, and in the storage and transfer of energy from one site in the body to another. Phosphorus enters the red blood cells with glucose and therefore is lowered in the plasma after carbohydrate ingestion or infusion. Phosphate levels are always evaluated in relation to calcium levels because there is an inverse relation between the two elements. When calcium levels are decreased, phosphorus levels are increased, and when phosphorus levels are decreased, calcium levels are increased. An excess of one electrolyte in serum causes the kidneys to excrete the other electrolyte. Many of the causes of elevated calcium levels are also causes of decreased phosphorus levels. As with calcium, the controlling factor is PTH.

Reference Values

Normal
- Adults: 2.7–4.5 mg/dL or 0.87–1.45 mmol/L
- Children: 4.5–5.5 mg/dL or 1.45–1.78 mmol/L
- Newborns: 4.5–9.0 mg/dL or 1.45–2.91 mmol/L

Procedure
Obtain a fasting, 5-mL, venous blood sample. Serum is preferred, but heparinized blood is acceptable. Serum should be removed from clot as soon as possible after collection.

Clinical Implications

1. Hyperphosphatemia (increased blood phosphorus levels) is most commonly found in association with kidney dysfunction and uremia. This is because phosphate is so minutely regulated by the kidneys. These conditions include the following:
   a. Renal insufficiency and severe nephritis (accompanied by elevated BUN and creatinine) and renal failure
   b. Hypoparathyroidism (accompanied by elevated phosphorus, decreased calcium, and normal renal function) and pseudohypoparathyroidism
c. Hypocalcemia
d. Milk-alkali syndrome
e. Excessive intake of vitamin D
f. Fractures in the healing stage
g. Bone tumors and metastases
h. Addison's disease
   i. Acromegaly
   j. Liver disease and cirrhosis
   k. Cardiac resuscitation

2. Hypophosphatemia (decreased phosphorus level) occurs in the following conditions:
   a. Hyperparathyroidism
   b. Rickets (childhood) or osteomalacia (adult) and vitamin D deficiency
c. Diabetic coma (increased carbohydrate metabolism)
   d. Hyperinsulinism
e. Continuous administration of IV glucose in a nondiabetic patient (phosphorus follows glucose into the cells)
f. Liver disease and acute alcoholism
g. Vomiting and severe diarrhea
h. Severe malnutrition and malabsorption
   i. Gram-negative sepsis
   j. Hypercalcemia of any cause
   k. Prolonged hypothermia
   l. Respiratory alkalosis due to cellular use of phosphorus for an accelerated glucose metabolism

Interfering Factors

1. Phosphorus levels are normally high in children.
2. Phosphorus levels can be falsely increased by hemolysis of blood; therefore, separate serum from cells as soon as possible.
3. Drugs can be the cause of decreases in phosphorus.
4. The use of laxatives or enemas containing large amounts of sodium phosphate will cause increased phosphorus levels. With oral laxatives, the blood phosphorus level may increase as much as 5 mg/dL (1.6 mmol/L) 2 to 3 hours after intake. This increased level is only temporary (5–6 hours), but this factor should be considered when abnormal levels are seen that cannot otherwise be explained.
5. Seasonal variations exist in phosphorus levels (maximum levels in May and June, lowest levels in winter).

Interventions

Pretest Patient Care

1. Explain test purpose and blood sampling procedures. The patient should fast before the test.
2. Note on test requisition if any catastrophic stressful events have taken place which may cause high phosphorus levels.
3. Note time of day test is drawn; levels are highest in the morning and lowest in the evening.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume normal activities.
2. Interpret test outcomes and monitor as appropriate for calcium imbalances. When phosphorus rises rapidly, calcium drops; watch for arrhythmias and muscle twitching. The signs and symptoms of phosphate depletion may include manifestations in the neuromuscular, neuropsychiatric, GI, skeletal, and cardiopulmonary systems. Manifestations usually are accompanied by serum levels < 1 mg/dL (<0.32 mmol/L).
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

### Clinical Alert

**Panic Value for Phosphate**

<1.0 mg/dL (<0.32 mmol/L)

**Magnesium (Mg^{2+})** Magnesium in the body is concentrated (40%–60%) in the bone, 20% muscle, 30% within the cell itself, and 1% in the serum, and is required for the use of adenosine triphosphate (ADP) as a source of energy. It is therefore necessary for the action of numerous enzyme systems such as carbohydrate metabolism, protein synthesis, nucleic acid synthesis, and contraction of muscular tissue. Along with sodium, potassium, and calcium ions, magnesium also regulates neuromuscular irritability and the clotting mechanism. Magnesium and calcium are intimately linked in their body functions, and deficiency of either one has a significant effect on the metabolism of the other because of magnesium's importance in the absorption of calcium from the intestines and in calcium metabolism. Magnesium deficiency will result in the drift of calcium out of the bones, possible resulting in abnormal calcification in the aorta and the kidney. This condition responds to administration of magnesium salts. Normally, 95% of the magnesium that is filtered through the glomerulus is reabsorbed in the tubule. When there is decreased kidney function, greater amounts of magnesium are retained, resulting in increased blood serum levels. Magnesium measurement is used to evaluate renal function, electrolyte status, and evaluate magnesium metabolism.

**Reference Values**

- **Normal**
  - Adults: 1.8–2.6 mg/dL or 0.74–1.07 mmol/L
  - Children: 1.7–2.1 mg/dL or 0.70–0.86 mmol/L
  - Newborns: 1.5–2.2 mg/dL or 0.62–0.91 mmol/L

**Procedure**

1. Obtain a fasting (4 hours), 5-mL, venous blood sample.
2. Avoid hemolysis, and separate serum from cells as soon as possible. Heparinized blood may be used.

### Clinical Alert

Blood sample should be drawn while the patient is in a prone position, because an upright position increases the magnesium level by 4%.

### Clinical Implications

1. Reduced blood magnesium levels (<1.5 mg/dL or <0.62 mmol/L) occur in the following conditions:
   a. Hypercalcemia of any cause
   b. Diabetic acidosis
   c. Hemodialysis
   d. Chronic renal disease (glomerulonephritis)
   e. Chronic pancreatitis
   f. Hyperaldosteronism
   g. Pregnancy (second and third trimester)
   h. Hypoparathyroidism
   i. Excessive loss of body fluids (eg, sweating, lactation, diuretic abuse, chronic diarrhea)
   j. Malabsorption syndromes
   k. Chronic alcoholism (hepatic cirrhosis)
   l. Long-term hyperalimentation
   m. Syndrome of inappropriate secretion of antidiuretic hormone (SIADH)
2. Increased blood magnesium levels occur in the following conditions:
   a. Renal failure or reduced renal function (acute and chronic)
   b. Dehydration
   c. Hypothyroidism
   d. Addison's disease
   e. Adrenalectomy (adrenocortical insufficiency)
   f. Diabetic acidosis (severe)
   g. Use of antacids containing magnesium (eg, Milk of Magnesia), administration of magnesium salts
   h. Oliguria

### NOTE

In magnesium deficiency states, urinary magnesium decreases before the serum magnesium. Serum magnesium levels may remain normal even when total body stores are depleted up to 20%.

### Interfering Factors
1. Prolonged salicylate therapy, lithium, and magnesium products (eg, antacids, laxatives) will cause falsely increased magnesium levels, particularly if there is renal damage.
2. Calcium gluconate, as well as a number of other drugs, can interfere with testing methods and cause falsely decreased results.
3. Hemolysis will invalidate results because about three fourths of the magnesium in the blood is found intracellularly in the red blood cells.

Interventions

Pretest Patient Care

1. Explain test purpose and blood-drawing procedure.
2. Ensure that patient is fasting for at least 4 hours if possible and is in a prone position when blood is drawn.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and monitor as appropriate. Treatment of diabetic coma often results in low plasma magnesium levels. This change occurs because magnesium moves with potassium into the cells after insulin administration.
2. Measure serum magnesium in persons receiving aminoglycosides and cyclosporine. There is a known association between these therapies and hypermagnesemia. Treatment of hypermagnesemia involves withholding source of magnesium excess, promoting excretion, giving calcium salts, and performing hemodialysis.
3. Be aware that magnesium deficiency may cause apparently unexplained hypocalcemia and hypokalemia. In these instances, patients may have neurologic and/or GI symptoms. Observe for the following signs and symptoms:
   - Muscle tremors, twitching, tetany
   - Hypocalcemia
   - Hyperactive deep tendon reflexes
   - Muscle tremors, twitching, tetany
   - Hypocalcemia
   - Hyperactive deep tendon reflexes
   - Electrocardiogram (ECG): prolonged P-R and Q-T intervals; broad, flat T waves; premature ventricular tachycardia and fibrillation
   - Anorexia, nausea, vomiting
   - Insomnia, delirium convulsions
4. Observe for signs of too much magnesium (which acts as a sedative):
   - Lethargy, flushing, nausea, vomiting, slurred speech
   - Weak or absent deep tendon reflexes
   - Electrocardiogram (ECG): prolonged P-R and Q-T intervals; broad, flat T waves; premature ventricular tachycardia and fibrillation
   - Anorexia, nausea, vomiting
   - Insomnia, delirium convulsions
5. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

Panic Values for Magnesium

1. Hypomagnesemia: <1.2 mg/dL (<0.49 mmol/L), tetany occurs
2. Hypermagnesemia: >5.0 mg/dL (>2.1 mmol/L)
   a. 5.0–10.0 mg/dL (2.1–4.1 mmol/L): CNS depression, nausea, vomiting, fatigue
   b. 10–15 mg/dL (4.1–6.2 mmol/L): coma, ECG changes, respiratory paralysis
   c. 30 mg/dL (12.3 mmol/L): complete heart block
   d. 34–40 mg/dL (14.0–16.0 mmol/L): cardiac arrest

Potassium (K⁺) Potassium is the principal electrolyte (cation) of intracellular fluid and the primary buffer within the cell itself. Ninety percent of potassium is concentrated within the cell; only small amounts are contained in bone and blood. Damaged cells release potassium into the blood. The body is adapted for efficient potassium excretion. Normally, 80% to 90% of the cells' potassium is excreted in the urine by the glomeruli of the kidneys; the remainder is excreted in sweat and in the stool. Even when no potassium is taken into the body (as in a fasting state), 40 to 50 mEq is still excreted daily in the urine. The kidneys do not conserve potassium, and when an adequate amount of potassium is not ingested, a severe deficiency will occur. Potassium balance is maintained in adults on an average dietary intake of 80–200 mEq/day. Normal intake, minimal needs, and maximum tolerance for potassium are almost the same as those for sodium. Potassium plays an important role in nerve conduction, muscle function, acid-base balance, and osmotic pressure. Along with calcium and magnesium, potassium controls the rate and force of contraction of the heart and, thus, the cardiac output. Evidence of a potassium deficit can be noted on an ECG by the presence of a U wave. Potassium and sodium ions are particularly important in the renal regulation of acid-base balance because hydrogen ions are substituted for sodium and potassium ions in the renal tubule. Potassium is more important than sodium because potassium bicarbonate is the primary intracellular inorganic buffer. In potassium deficiency, there is a relative deficiency of intracellular potassium bicarbonate, and the pH is relatively acid. The respiratory center responds to the intracellular acidosis by lowering Pco₂ through the mechanism of hyperventilation. The potassium concentration is greatly affected by the adrenal hormones. Potassium deficiency will cause a significant reduction in protein synthesis. This test evaluates changes in body potassium levels and diagnoses acid-base and water imbalances. The level of potassium is not an absolute value; it varies with circulatory volume and other factors. Because a totally unsuspected potassium imbalance can suddenly prove lethal, its development must be anticipated. Thus, it is important to check the potassium level in severe cases of Addison's disease, uremic coma, intestinal obstruction, acute renal failure, GI loss in the administration of diuretics, steroid therapy, and cardiac patients on digitals. Potassium levels should be monitored during treatment of acidosis, including ketoacidosis of diabetes mellitus.

Reference Values

Normal Adults: 3.5–5.2 mEq/L (3.5–5.2 mmol/L) Children (1–18 years): 3.4–4.7 mEq/L (3.4–4.7 mmol/L) Infants: (7 days–1 year): 4.1–5.3 mEq/L (4.1–5.3 mmol/L) Neonates (0–7 days): 3.7–5.9 mEq/L (3.7–5.9 mmol/L)
Procedure

1. General procedure for potassium (K⁺)
   a. Collect a 5-mL venous blood sample using serum or heparinized Vacutainer tube. Observe standard/universal precautions. Avoid hemolysis in obtaining the sample.
   b. Deliver the sample to the laboratory and centrifuge immediately to separate cells from serum. Potassium leaks out of the cell and levels in the sample will be falsely elevated later than 4 hours after collection.

2. Procedure for hyperkalemia (excess K⁺)
   a. Record fluid intake and output. Check blood volume and venous pressure, which will give clues to dehydration or circulatory overload. Identify ECG changes. In hyperkalemia, these include the following:
      1. Elevated T-wave heart block
      2. Flattened P wave
      3. Cardiac arrest may occur without warning other than ECG changes.
   b. Observe for slow pulse, oliguria, neuromuscular alterations such as muscle irritability and impaired muscle function, flaccid paralysis, tremors, and twitching preceding actual paralysis.
   c. Hyperkalemia can be treated with sodium bicarbonate, glucose, and insulin. Kayexalate, a sodium-potassium exchange resin, can be administered orally, nasogastrically, or rectally.

3. Procedure for hypokalemia (deficiency of K⁺)
   a. Record fluid intake and output. Check blood volume and venous pressure, which will give clues to circulatory overload or dehydration. Identify ECG changes. In hypokalemia, these include the following:
      1. Depressed T waves
      2. Peaking of P waves
   b. Observe for dehydration caused by severe vomiting, hyperventilation, sweating, diuresis, or nasogastric tube with gastric suction. Accurately record state of hydration or dehydration.
   c. Observe for neuromuscular changes such as fatigue, muscle weakness, muscle pain, flabby muscles, paresthesia, hypotension, rapid pulse, respiratory muscle weakness leading to paralysis, cyanosis, respiratory arrest, anorexia, nausea, vomiting, paralytic ileus, apathy, drowsiness, tetany, and coma.
   d. Hypokalemia may be treated with a K⁺-rich diet and K⁺-sparing diuretics. Use salt substitutes containing potassium chloride and administer IV oral potassium chloride supplements.

Clinical Alert

1. Be on the alert for the following arrhythmias, which may occur with hyperkalemia:
   a. Sinus bradycardia
   b. Sinus arrest
   c. First-degree atrioventricular block
   d. Nodal rhythm
   e. Idioventricular rhythm
   f. Ventricular tachycardia
   g. Ventricular fibrillation
   h. Ventricular arrest

2. Be on the alert for the following arrhythmias, which may occur with hypokalemia:
   a. Ventricular premature beats
   b. Atrial tachycardia
   c. Nodal tachycardia
   d. Ventricular tachycardia
   e. Ventricular fibrillation

Clinical Implications

1. Decreased blood potassium (hypokalemia) levels are associated with shifting of K⁺ into cells, K⁺ loss from GI and biliary tracts, renal K⁺ excretion, and reduced K⁺ intake, as can occur in the following conditions:
   a. Diarrhea, vomiting, sweating
   b. Starvation, malabsorption
   c. Bartter's syndrome
   d. Draining wounds
   e. Cystic fibrosis
   f. Severe burns
   g. Primary aldosteronism
   h. Alcoholism, chronic
   i. Osmotic hyperglycemia
   j. Respiratory alkalosis
   k. Renal tubular acidosis
   l. Diuretic, antibiotic, and mineralocorticoid administration
   m. Barium chloride poisoning
   n. Treatment of megaloblastic anemia with vitamin B₁₂ or folic acid

2. Potassium levels of 3.5 mEq/L (3.5 mmol/L) are more commonly associated with deficiency rather than normality. A falling trend (0.1–0.2 mEq/day or 0.1–0.2 mmol/day) is indicative of a developing potassium deficiency. A.
   a. The most frequent cause of potassium deficiency is GI loss.
   b. The most frequent cause of potassium depletion is IV fluid administration without adequate potassium supplements

3. Increased potassium levels (hyperkalemia) occur when K⁺ shifts from cells to intracellular fluid, with inadequate renal excretion and with excessive K⁺ intake, as can occur in the following conditions:
   a. Renal failure, dehydration, obstruction, and trauma
b. Cell damage, as in burns, accidents, surgery, chemotherapy, disseminated intravascular coagulation (damaged cells release potassium into the blood)

c. Metabolic acidosis (drives potassium out of the cells), diabetic ketoacidosis
d. Addison's disease
e. Pseudohypoaldosteronism
f. Uncontrolled diabetes, decreased insulin
g. Primary acquired hyperkalemia, such as systemic lupus erythematosus, sickle cell disease, interstitial nephritis, and tubular disorders
h. Kidney transplant rejection

Interfering Factors

1. Hemolyzed blood may not be used; 
   Potassium levels are elevated to as much as 50% over normal with moderate hemolysis. Opening and closing the fist 10 times with a tourniquet in place results in an increase in potassium level by 10% to 20%. For this reason, it is recommended that the blood sample be obtained without a tourniquet, or that the tourniquet be released after the needle has entered the vein.

2. Drug usage
   a. IV administration of potassium penicillin may cause hyperkalemia; penicillin sodium may cause increased excretion of potassium.
   b. Glucose administered during tolerance testing or the ingestion and administration of large amounts of glucose in patients with heart disease may cause a decrease of as much as 0.4 mEq/L (0.4 mmol/L) in potassium blood levels.
   c. A number of drugs raise potassium levels, especially potassium-sparing diuretics and nonsteroidal anti-inflammatory drugs, especially in the presence of renal disease.
   d. Excessive intake of licorice decreases potassium levels.

3. Leukocytosis, as occurs in leukemia, raises potassium levels.

4. Patients who have thrombocytosis due to polycythemia vera or a myeloproliferative disease may have spuriously high potassium levels. This falsely elevated level is caused by a high number of platelets, which release potassium during coagulation. Therefore, heparinized samples, rather than clotted serum samples, should be used in these patients.

Interventions

Pretest Patient Care

1. Explain test purpose and blood-drawing procedure. Do not have patient open and close fist while drawing blood.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results, monitor changes in body potassium, and intervene as appropriate.
2. Be aware that recognizing signs and symptoms of hypokalemia and hyperkalemia is very important. Many of these originate in the nervous and muscular systems and are usually nonspecific and similar.
3. Remember that the potassium blood level rises 0.6 mEq/L (0.6 mmol/L) for every 0.1 decrease in blood pH.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

1. Panic values for potassium:
   a. < 2.5 mEq/L (<2.5 mmol/L) causes ventricular fibrillation.
   b. > 8.0 mEq/L (>8.0 mmol/L) causes muscle irritability, including myocardial irritability.

2. The most common cause of hypokalemia in patients receiving IV fluids is water and sodium chloride administration without adequate replacement for potassium lost in urine and drainage fluids. A patient receiving IV fluids needs potassium every day. The minimum adult daily dose should be 40 mEq, but the optimum daily dose ranges between 60 and 120 mEq. (Pediatric dose, 0.5–1 mEq/kg of body weight/24 h, not to exceed 40 mEq/d or 10 mEq/h.) Potassium needs are greater in persons with tissue injury, wound infection, and gastric intestinal or biliary drainage. If adequate amounts of potassium (40 mEq/d) are not given in IV solution, hypokalemia will eventually develop. Patients receiving > 10 mEq KC1 in 100 mL of IV solution should be monitored by ECG for potential arrhythmia if the IV rate is = 100/h. Concentrated doses of IV potassium should always be administered via volume-controlled IV infusion devices. A burning sensation may be felt at the site of needle insertion. NS can be infused along with the potassium, or the IV rate can be reduced. Some physicians order a small dose of lidocaine to be added to IV potassium to eliminate the burning sensation some patients experience. Always be sure to check for lidocaine allergies before administration of this local anesthetic.

3. Closely monitor for hypokalemia in patients taking digitalis and diuretics because cardiac arrhythmias can occur. Hypokalemia enhances the effect of digitalis preparations, creating the possibility of digitalis intoxication from even an average maintenance dose. Digitalis, diuretics, and hypokalemia are a potentially lethal combination.

Sodium (Na⁺) Sodium is the most abundant cation (90% of the electrolyte fluid) and the chief base of the blood. Its primary functions in the body are to maintain osmotic pressure and acid-base balance chemically and to transmit nerve impulses. The body has a strong tendency to maintain a total base content, and only slight changes are found even under pathologic conditions. Mechanisms for maintaining a constant sodium level in the plasma and extracellular fluid include renal blood flow, carbonic anhydrase enzyme activity, aldosterone, action of other steroids whose plasma level is controlled by the anterior pituitary gland, renin enzyme secretion, ADH, and vasopressin secretion. Determinations of plasma sodium levels detect changes in water balance rather than sodium balance. Sodium levels are used to determine electrolytes, acid-base balance, water balance, water intoxication, and dehydration.

Reference Values
Normal Adults: 136–145 mEq/L (136–145 mmol/L) Children (1–16 years): 136–145 mEq/L (136–145 mmol/L) Full-term infants: 133–142 mEq/L (133–142 mmol/L) Premature infants: 132–140 mEq/L (132–140 mmol/L)

Procedure

1. Obtain a 5-mL venous blood sample. Heparinized blood can be used. Avoid hemolysis.
2. Observe standard/universal precautions.

Clinical Implications

1. Hyponatremia (decreased sodium levels) reflect a relative excess of body water rather than low total body sodium. Reduced sodium levels (hyponatremia) are associated with the following conditions:
   a. Severe burns
   b. Congestive heart failure (predictor of cardiac mortality)
   c. Excessive fluid loss (eg, severe diarrhea, vomiting, sweating)
   d. Excessive IV induction of nonelectrolyte fluids (eg, glucose)
   e. Addison's disease (impairs sodium reabsorption)
   f. Severe nephritis (nephrotic syndrome)
   g. Pyelocentric obstruction
   h. Malabsorption syndrome
   i. Diabetic acidosis
   j. Drugs such as diuretics
   k. Edema (diuretic hyponatremia)
   l. Large amounts of water by mouth (water intoxication)
   m. Stomach suction accompanied by water or ice chips by mouth
   n. Hypothyroidism
   o. Excessive ADH production

2. Hypernatremia (increased sodium levels) is uncommon, but when it does occur, it is associated with the following conditions:
   a. Dehydration and insufficient water intake
   b. Conn's syndrome
   c. Primary aldosteronism
   d. Coma
   e. Cushing's disease
   f. Diabetes insipidus
   g. Tracheobronchitis

Clinical Alert

Panic Values for Sodium

1. <125 mEq/L (<125 mmol/L) causes weakness, dehydration.
2. 90–105 mEq/L (90–105 mmol/L) causes severe neurologic symptoms, vascular problems.
3. >152 mEq/L (>152 mmol/L) results in cardiovascular and renal symptoms.
4. >160 mEq/L (>160 mmol/L) can cause heart failure.

Interfering Factors

1. Many drugs affect levels of blood sodium.
   a. Anabolic steroids, corticosteroids, calcium, fluorides, and iron can cause increases in sodium level.
   b. Heparin, laxatives, sulfates, and diuretics can cause decreases in sodium level.
2. High triglycerides or low protein causes artificially low sodium values.

Interventions

Pretest Patient Care

1. Explain test purpose and procedure.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and monitor for fluid and sodium imbalances.
2. Remember that IV therapy considerations are as follows:
   a. Sodium balance is maintained in adults with an average dietary intake of 90 to 250 mEq/ day. The maximum daily tolerance to an acute load is 400 mEq/day. A patient who is given 3 L of isotonic saline in 24 hours will receive 465 mEq of sodium. This amount exceeds the average, healthy adult's tolerance level. It will take a healthy person 24 to 48 hours to excrete the excess sodium.
   b. After surgery, trauma, or shock, there is a decrease in extracellular fluid volume. Replacement of extracellular fluid is essential if water and electrolyte balance is to be maintained. The ideal replacement IV solution should have a sodium concentration of 140 mEq/L.
3. Monitor for signs of edema or hypertension, and record and report these if present.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Osmolality and Water-Load Test (Water-Loading Antidiuretic Hormone Suppression Test) Osmolality, which is the measure of the number of dissolved solute particles in solution, increases with dehydration and decreases with overhydration. In general, the same conditions that reduce or increase serum sodium affect osmolality. This test is used
as an evaluation of water and electrolyte balance. It is helpful in assessing hydration status, seizures, liver disease, ADH function, and coma, and it is used in toxicology workups for ethanol, ethylene glycol, isopropanol, and methanol ingestions.

**Clinical Alert**

1. Simultaneous determination of urine and serum osmolalities facilitates interpretation of results. High urinary/serum (U/S) ratio is seen in concentrated urine. Normal ranges for the U/S ratio are 0.2 to 4.7 and may be >3 with overnight dehydration. With poor concentrating ability, the ratio is low but is still >1. In SIADH, sodium and urine osmolalities are high for the serum osmolality.

2. Determination of the urine osmolar gap is used to characterize metabolic acidosis and is described as the sum of urinary concentrations of sodium, potassium, bicarbonate, chloride, glucose, and urea compared with measured urine osmolality.

**Reference Values**

<table>
<thead>
<tr>
<th>Normal Serum osmolality</th>
<th>Adults: 280–303 mOsm/kg H₂O (280–303 mmol/kg H₂O) Newborns: as low as 266 mOsm/kg H₂O (266 mmol/kg H₂O)</th>
<th>Urine osmolality</th>
<th>Adults: 24-hour: 300–900 mOsm/kg H₂O (300–900 mmol/kg H₂O) Random: 50–1400 mOsm/kg H₂O (50–1400 mmol/kg H₂O) After 12-hour fluid restriction: &gt;850 mOsm/kg H₂O (&gt;850 mmol/kg H₂O)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ratio of urine/serum osmolality: 0.2–4.7 (average, 1.0–3.0) Ratio after fluid restriction: 3:1 or a range of 0.2–4.7:1</td>
<td>Osmolar gap Serum: 5–10 mOsm/kg H₂O (5–10 mmol/kg H₂O) Urine: 80–100 mOsm/kg H₂O (80–100 mmol/kg H₂O)</td>
<td>Procedure</td>
<td></td>
</tr>
</tbody>
</table>

**Procedure**

1. Determining osmolality
   a. Obtain a 5-mL venous blood sample. Serum or heparinized plasma is acceptable. Observe standard/universal precautions.
   b. Collect a 24-hour urine specimen concurrently and keep on ice.
   c. Determine osmolality in the laboratory using the freezing point depression methodology for both serum and urine.

2. Determining water-loading antidiuretic hormone suppression
   a. Be aware that the ideal position during the testing period is the recumbent position because the response to water loading is reduced in persons in the upright position.
   b. One hour before testing, the patient is given 300 mL of water to replace fluid lost during the overnight fast. Do not count this water as part of the test load.
   c. Have the patient drink a test load of water (20 mL/kg body weight) within 30 minutes.
   d. After the test load of water is consumed, collect all urine for the next 4 to 5 hours, and check each voiding for volume osmolality and specific gravity. Obtain hourly blood samples for osmolality, and check the entire volume of urine obtained for osmolality.

3. Remember that normal values for water-loading antidiuretic hormone suppression test are excretion of > 90% of water load within 4 hours. Urine osmolality falls to < 100 mOsm/kg (<100 mmol/kg). Specific gravity falls to 1.001.

4. Determine plasma ADH at hourly intervals.

**Clinical Implications of Decreased Renal Function**

1. In decreased renal function, < 80% of fluid is excreted, and urine specific gravity may not fall below 1.010. This phenomenon occurs in the following conditions:
   a. Adrenocortical insufficiency
   b. Malabsorption syndrome
   c. Edema
   d. Ascites
   e. Obesity
   f. Hypothyroidism
   g. Dehydration
   h. Congestive heart failure
   i. Cirrhosis

2. Disorders with increased ADH secretion (SIADH) give an inadequate response; <90% of water is excreted, and urine osmolality remains > 100 mOsm/kg (>100 mmol/kg H₂O). Plasma ADH measured at 90 minutes confirms diagnosis of SIADH.

**Interventions**

**Pretest Patient Care: Decreased Renal Function**

1. Explain the test purpose and procedure. The test takes 5 to 6 hours to complete.
2. Do not allow food, alcohol, medications, or smoking for 8 to 10 hours before testing. No muscular exercise is allowed during the test.
3. Be aware that the patient may experience nausea, abdominal fullness, fatigue, and desire to defecate.
4. Discard first morning urine specimen.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Clinical Alert**

In patients with impaired ability to tolerate the water-loading test, seizures or fatal hyponatremia may occur.

**Posttest Patient Aftercare: Decreased Renal Function**

1. Observe for adverse reactions to water-loading test such as extreme abdominal discomfort, shortness of breath, or chest pain.
2. Remember that if water clearance is impaired, the water load will not induce diuresis, and maximum urinary dilution will not occur.
3. Be aware that accurate results may not be obtained if nausea, vomiting, or diarrhea occur or if a disturbance in bladder emptying is present. Note on chart if any of these effects occur.
4. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Clinical Implications of Hyperosmolality and Hypoosmolality**

1. **Increased values (hyperosmolality)** are associated with the following conditions:
   a. Dehydration
   b. Hypercalcemia
   c. Diabetes mellitus, hyperglycemia, diabetic ketoacidosis
   d. Hypernatremia
   e. Cerebral lesions
   f. Alcohol ingestion (ethanol, methanol, ethylene glycol)
   g. Mannitol therapy
   h. Azotemia
   i. Inadequate water intake
   j. Chronic renal disease

2. **Decreased values (hypoosmolality)** are associated with the following conditions:
   a. Loss of sodium with diuretics and low-salt diet (hyponatremia)
   b. Renal failure
   c. Adrenocortical insufficiency
   d. Inappropriate secretion of ADH, as may occur in trauma and lung cancer
   e. Excessive water replacement (overhydration, water intoxication)
   f. Panhypopituitarism
   g. Diabetes insipidus (central or nephrogenic)
   h. Pyelonephritis

**Clinical Implications of Osmolal Gap**

1. Abnormal levels (>10 mOsm/kg H_2O or >10 mmol/kg H_2O) can occur in the following conditions:
   a. Methanol
   b. Ethanol
   c. Isopropyl alcohol
   d. Mannitol
   e. Severely ill patients, especially those in shock, lactic acidosis, and renal failure

2. Ethanol, glycol, acetone, and paraaldehyde have relatively small osmolal gaps, even at lethal levels.

**Interfering Factors**

1. Decreases in osmolal gap are associated with altitude, diurnal variation with water retention at night, and some drugs.
2. Some drugs also cause increases in osmolal gap.
3. Hypertriglyceridemia and hyperproteinemia cause an elevated osmolal gap.
4. Radiographic contrast medium within 3 days

**Interventions**

**Pretest Patient Care: Hyperosmolality, Hypoosmolality, Osmolar Gap**

1. Explain test purpose and procedure.
2. Ensure that no alcohol is ingested during the 24 hours before the test.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare: Hyperosmolality, Hypoosmolality, Osmolar Gap**

1. Interpret test results and monitor appropriately. A patient receiving IV fluids should have a normal osmolality. If the osmolality increases, the fluids contain relatively more electrolytes than water. If it falls, relatively more water than electrolytes is present.
2. Remember that if the ratio of serum sodium to serum osmolality falls below 0.43, the outlook is guarded. This ratio may be distorted in cases of drug intoxication.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Clinical Alert**

1. Panic serum osmolality values are results <240 or >321 mOsm/kg H_2O (<240 or >321 mmol/kg H_2O). A value of 385 mOsm/kg H_2O (385 mmol/kg H_2O) relates to stupor in hyperglycemia. Values of 400–420 mOsm/kg H_2O (400–420 mmol/kg H_2O) are associated with grand mal seizures. Values >420 mOsm/kg H_2O (>420 mmol/kg H_2O) are fatal.
2. A water-loading ADH suppression test may be ordered to investigate impaired renal excretion of water.

**Sweat Test** This test is done to diagnose cystic fibrosis. Abnormally high concentrations of sodium and chloride appear in the secretions of eccrine sweat glands in persons with cystic fibrosis. This condition is present at birth and persists throughout life. This study uses sweat-inducing techniques (eg, pilocarpine iontophoresis) followed by chemical analysis.
to determine sodium, chloride, and content of collected sweat.

**Reference Values**

**Normal** Sweat sodium: Normal: <70 mEq/L (<70 mmol/L) Cystic fibrosis: >90 mEq/L (>90 mmol/L) Sweat chloride: Normal: <50 mEq/L (<50 mmol/L) Cystic fibrosis: >60 mEq/L (>60 mmol/L)

**Procedure**

1. Be aware that the forearm is the preferred site for stimulation of sweating, but in thin or small babies, the thigh, back, or leg may be used. It may be necessary to stimulate sweating in two places to obtain sufficient sweat for testing, especially in young infants. At least 100 µL of sweat is necessary. In cold weather, or if the testing room is cold, a warm covering should be placed over the arm or other site of sweat collection.
2. Stimulate sweat production by applying gauze pads or filter paper saturated with a measured amount of pilocarpine and attachment of electrodes through which a current of 4 to 5 mAmp is delivered at intervals for a total of 5 minutes (a total of 5–12 minutes, according to NIH).
3. Remove the electrodes and pad, and thoroughly wash the area with distilled water; dry carefully.
4. Remember that successful iontophoresis is indicated by a red area about 2.5 cm in diameter that appears where the electrode was placed.
5. Scrub the skin thoroughly with distilled water and dry carefully. The area for sweat collection must be completely dry, free from contamination by powder or antiseptic, and free of any area that might ooze.
6. Be aware that collection of sweat occurs by applying preweighed filter or sweat collection cups that are taped securely over the red spot. The inside surfaces of the collecting device should never be touched.
7. Leave the paper on for at least 1 hour before removing and then place in a preweighed flask to avoid evaporation. Weigh the flask again. The desired volume of sweat is 200 mg; the minimum volume necessary is 100 mg.
8. If a cup is used, leave in place for 1 hour and then carefully remove by scraping it across the iontophoresed area. This “puddles” the sweat in the cup to reduce evaporation and to redissolve any salts left by the evaporation. Use suction capillary tubes to remove sweat from the collection cups.

**Clinical Implications**

1. Children with cystic fibrosis have sodium and chloride values of >90 mEq/L and >60 mEq/L (>90 mmol/L and >60 mmol/L), respectively.
2. Borderline or gray-zone cases are those with values of 70–90 mEq/L (70–90 mmol/L) for sodium and 50–60 mEq/L (50–60 mmol/L) for chloride. These persons require retesting. Potassium values do not assist in differentiating borderline cases.
3. In adolescence and adulthood, chloride levels > 80 mEq/L (>80 mmol/L) usually indicate cystic fibrosis.
4. Elevated sweat electrolytes also can be associated with the following conditions:
   a. Addison's disease
   b. Congenital adrenal hyperplasia
   c. Vasopressin-resistant diabetes insipidus
   d. Glucose-6-phosphatase deficiency (G6PD)
   e. Hypothyroidism
   f. Familial hypoparathyroidism
   g. Alcoholic pancreatitis

**Interfering Factors**

1. The sweat test is not valuable after puberty because levels may vary over a very wide range among individuals.
2. Dehydration and edema, particularly of areas where sweat is collected, may interfere with test results.
3. A gap of > 30 mEq/L (>30 mmol/L) between sodium and chloride values indicates calculation or analysis error or contamination of the sample.
4. Sweat testing is not considered accurate until the third or fourth week of life because infants < 3 weeks of age may not sweat enough to provide a sufficient sample.
5. Test may be falsely normal in patients with salt depletion, as in periods of hot weather.

**Clinical Alert**

1. The test should always be repeated if the result, the clinical features, or other diagnostic tests do not fit together.
2. The test can be used to exclude the diagnosis of cystic fibrosis in siblings of diagnosed patients.
3. There have been reports of cystic fibrosis patients with normal sweat electrolyte levels.
4. Sweat potassium testing is not diagnostically valuable.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. The sweat test is indicated for the following persons:
   a. Infants who pass initial meconium late; who have intestinal obstruction, failure to thrive, steatorrhea, chronic diarrhea, rapid respiration and retraction with chronic cough, asthma, hypoproteinemia (especially on soybean formula), atelectasis, or hyperpnea on x-ray, hyperprothrombinemia, or rectal prolapse; who taste salty; or who are offspring of a parent with cystic fibrosis (ie, the obligate heterozygote)
   b. Persons suspected of having cystic fibrosis or celiac disease, all siblings of patients with cystic fibrosis, or persons with disaccharide intolerance, recurrent pneumonia, chronic atelectasis, chronic pulmonary disease, bronchioloclastic, chronic cough, nasal polypsis, cirrhosis of liver, and hypertension
   c. Any parents who request a sweat test on their child
2. Inform the patient that a slight stinging sensation is usually experienced, especially in fair-skinned persons.
3. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.
Posttest Patient Aftercare

1. After the cup is removed, carefully wash and dry the skin to prevent irritation caused by collection cups.
2. Have patient resume normal activities.
3. Interpret test results and counsel and monitor patient as appropriate. Provide genetic counseling. Cystic fibrosis is transmitted as an autosomal recessive trait. The Caucasian carrier rate is 1 in 20, and the African American carrier rate is 1 in 60 to 1 in 100.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

BIBLIOGRAPHY

Gale Encyclopedia of Medicine. Farming Hills, MI, Gale, 1999
Martin L: What You Really Need to Know to Interpret Arterial Blood Gases, 2nd ed. Philadelphia, Lippincott Williams & Wilkins, 1999
University of California at San Diego Library.
OVERVIEW OF PRENATAL DIAGNOSIS

MATERNAL TRIPLE SCREEN

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Hormone Testing

Estriol (E3)

Human Chorionic Gonadotropin (HCG)

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AMNIOTIC FLUID STUDIES

Amniocentesis

Amniotic Fluid Alpha1-Fetoprotein (AFP)

Amniotic Fluid Total Volume

Amniotic Fluid Index (AFI)

Amniotic Fluid Creatinine

Amniotic Fluid Lechithin-to-Sphingomyelin (L/S) Ratio (Surfactant Components)

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Amniotic Fluid Color

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Amniotic Fluid and Desaturated Phosphatidylcholine (DSPC)

ADDITIONAL MEASUREMENTS

Group B Streptococcal (GBS) Screening

Fetal Oxygen Saturation Monitoring (FSpO2)

Fetal Nuchal Translucency (FNT)

BIBLIOGRAPHY

OVERVIEW OF PRENATAL DIAGNOSIS

Fetal well-being depends on maternal health. Many routine prenatal tests assess maternal health and well-being. Prenatal testing usually includes a complete blood count or hemoglobin and hematocrit, Rh type and ABO blood group, red cell antibody screening, rubella immunity status, glucose challenge testing (see Chap. 6), urinalysis, maternal serum alpha-fetoprotein (MS-AFP) or maternal triple screen, hepatitis B testing, culture for sexually transmitted diseases, the Venereal Disease Research Laboratory (VDRL) test, and surveillance for group B streptococci. Screening for human immunodeficiency virus (HIV) infection is recommended for all pregnant women to improve the care of HIV-positive women and to identify infants at risk; perinatal transmission is the primary route of HIV infection in children.

Tests in this chapter monitor the status of the maternal-fetal unit, identify the fetus at risk for intrauterine asphyxia, aid in the early diagnosis of infection, and identify genetic and biochemical disorders and major anomalies. (See Chap. 11 for more on genetic disorders.) Tests are also performed to predict normal fetal outcome or to identify the fetus at risk for asphyxia during labor.

MATERNAL TRIPLE SCREEN

The maternal triple screen tests are offered to pregnant women to identify risks for chromosome disorders such as Down's syndrome (trisomy 21); major birth defects, including open neural tube defects such as spina bifida; placental insufficiency; and oligohydramnios. The evaluation consists of three separate blood protein tests done on maternal serum between 14 and 19 weeks of gestation: MS-AFP is decreased in Down's syndrome and neural tube defects, estriol (E3) in Down's syndrome, and beta-human chorionic gonadotropin (hCG) in Down's syndrome. Results are reported as multiples of the median (MoM).

The maternal triple screen is a screening test; therefore, an abnormal (positive) result is not diagnostic, and further testing with ultrasound, amniocentesis, and genetic counseling is indicated. The markers can be positive in normal variations such as multiple births or miscalculated gestational age.

Ultrasound testing is a method of assessing fetal well-being that has become a diagnostic tool for assessment of fetal age, health, growth, and identification of anomalies. Level I ultrasound assesses gestational age, number of fetuses, fetal
death, and the condition of the placenta. Level II ultrasound assesses specific congenital anomalies or abnormalities. In some diagnostic centers, fetal echocardiography is also available. Color-enhanced Doppler sonography is used to measure the velocity and direction of blood flow in fetal and uterine anatomy, to provide information about placental function, and as an especially good predictor of outcome for fetuses that are small for gestational age (see Chap. 13).

Although magnetic resonance imaging (MRI) is used at some prenatal centers, it is still under investigation for diagnostic evaluation in pregnancy, especially in the final trimester (see Chap. 16). Some of the advantages of MRI during pregnancy are that it is a noninvasive technique, it permits easy differentiation between fat and soft tissue, it does not require a full bladder, and it can show the entire fetus in one scan. Currently, MRI confirms fetal abnormalities found by ultrasound and can be used for pelvimetry, placental localization, and determination of size. Fetal MRI is used at medical centers that specialize in fetal diagnosis and treatment (particularly those that perform fetal surgery). Ultrafast MRI is used for evaluation of congenital anomalies that are potentially correctable, such as congenital diaphragmatic hernia, neck masses that result in airway obstruction, and myelomeningocele. MRI is especially useful for definition of maternal anatomy in cases of suspected intraabdominal or retroperitoneal disease. Also under investigation is the combined use of a blood test for pregnancy-associated plasma protein A (PAPPA), which is increased in Down's pregnancy, and ultrasound measurement of neck membrane thickness, which is increased in Down's syndrome.

Maternal Serum Alpha-Fetoprotein (MS-AFP)

AFP, a product of the fetal liver, is normally found in fetal serum, maternal serum, and amniotic fluid. MS-AFP testing is routinely offered between 15 and 18 weeks of gestation to all pregnant women as a screen for neural tube defects; only 5% to 10% of neural tube defects occur in families with previous occurrences.

Reference Values

**Normal** 25 ng/mL (25 µg/L) At 15–18 weeks' gestation: 10–150 ng/mL or 10–150 µg/L

**Procedure**

1. Obtain a 10-mL venous blood sample (red-topped tube). Observe standard precautions. Place specimen in a biohazard bag.
2. Plan the first screening at 15 to 18 weeks. If the result is normal, no further screening is necessary. If MS-AFP is low, consider ultrasound studies to determine exact fetal age. A second screening may be done after an initial elevated MS-AFP. If the result is normal, no further screening is necessary.

**Clinical Implications** Abnormal levels should be followed by ultrasound and amniocentesis.

1. **Elevated** MS-AFP can indicate:
   a. Neural tube defects of spina bifida (a vertebral gap) or anencephaly (>2.5 MoM)
   b. Underestimation of gestational age
   c. Multiple gestation (>4.5 MoM)
   d. Threatened abortion
   e. Other congenital abnormalities
2. **Elevated** MS-AFP early in pregnancy is associated with:
   a. Congenital nephrosis
   b. Duodenal atresia
   c. Umbilical hernia or protrusion
   d. Sacrococcygeal teratoma
3. **Elevated** MS-AFP in the third trimester is associated with:
   a. Esophageal atresia
   b. Fetal teratoma
   c. Hydrocephaly
   d. Rh isomunization
   e. Gastrointestinal tract obstruction
4. **Low** MS-AFP is associated with:
   a. Long-standing fetal death
   b. Down's syndrome (trisomy 21)
   c. Other chromosome abnormalities (trisomy 13, trisomy 18)
   d. Hydatidiform mole
   e. Pseudopregnancy

**Interfering Factors**

1. Obesity causes low MS-AFP.
2. Race is a factor: MS-AFP levels are 10% to 15% higher in blacks and are lower in Asians.
3. Insulin-dependent diabetes results in low MS-AFP.

**Clinical Alert**

1. The incidence of neural tube defect is 1 per 1000 births in the United States, 1 per 5000 in England.
2. Knowledge of the precise gestational age is paramount for the accuracy of this test.
3. If the MS-AFP is elevated and no fetal defect is demonstrated (ie, by ultrasound or amniocentesis), then the pregnancy is at an increased risk (eg, premature birth, low-birth-weight infant, fetal death).

**Interventions**

**Pretest Patient Preparation**
1. Explain the reason for testing the mother's blood.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel appropriately. Explain possible need for further testing (eg, ultrasound, amniocentesis).
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Hormone Testing**

Normally, the amounts of all steroid hormones increase as pregnancy progresses. The maternal unit responds to altered hormone levels even before the growing uterus is apparent. Serial testing may be done to monitor rising levels of a particular hormone over a period of time. Decreasing levels indicate that the maternal-placental-fetal unit is not functioning normally. Biochemical analyses of several hormones can be used to monitor changes in the status of the maternal-fetal unit (See Chap. 3 and Chap. 6).

1. In early pregnancy, hCG in maternal blood provides evidence of a viable pregnancy. The hCG in maternal serum is measured as a sensitive pregnancy test (the hCG level doubles every 48 hours during pregnancy). Also, it is used to monitor the success of in vitro fertilization or insemination, to diagnose trophoblastic tumor, to diagnose ectopic pregnancy (indicated by decrease in hCG over a 48-hour period), and to screen for Down's syndrome in pregnancy. For further discussion of pregnancy tests, see Chapter 6.
2. The hCG, together with prolactin and luteinizing hormone (LH), prolongs the life of the corpus luteum once the ovum is fertilized. hCG stimulates the ovary for the first 6–8 weeks of pregnancy, before placental synthesis of progesterone begins. Its function later in pregnancy (in maternal blood) is unknown.
3. PAPPA, a circulating placental protein, has been shown to increase the stimulatory effects of placental insulin-like growth factors. Decreased serum levels in the maternal circulation in the first 10 weeks after conception are associated with uncomplicated full-term low birth weights. PAPPA levels are detectable within 30 days after conception and slowly increase throughout the first 30 weeks of gestation. Maternal serum levels are 0.43 µg/L (12 pmol/L). Increased PAPPA occurs in Down's pregnancy.
4. Late in pregnancy, the levels of estriol (E₃) and human placental lactogen (hPL) in maternal blood reflect fetal homeostasis. hPL is a protein hormone produced by the placenta. Testing of hPL evaluates only placental functioning. Blood testing of the mother usually begins after the 30th week and may be done weekly thereafter. A concentration of 1 µg/mL (46 nmol/L) hPL may be detected at 6–8 weeks of gestation. The level slowly increases throughout pregnancy and reaches 7 µg/mL (324 nmol/L) at term before abruptly dropping to zero after delivery. hPL functions primarily as a fail-safe mechanism to ensure nutrient supply to the fetus, for example, at times of maternal starvation. However, it does not appear to be required for a successful pregnancy outcome (see Chap. 6).

**Estriol (E₃)**

E₃ is the predominant estrogen in the blood and urine of pregnant women and is of fetal origin. Normal production serves as a measure of the integrity of the maternal-fetal unit and of fetal well-being.

This test is used during pregnancy to evaluate fetal disorders and is part of the maternal triple screen. Declining serial values indicate fetal distress. E₃ is decreased in Down's syndrome and in trisomy 18.

**Reference Values**

**Normal**

<table>
<thead>
<tr>
<th>Weeks of Gestation</th>
<th>E₃ (ng/mL)</th>
<th>SI Units (nmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>28–30</td>
<td>38–140</td>
<td>132–485</td>
</tr>
<tr>
<td>32</td>
<td>35–330</td>
<td>121–1144</td>
</tr>
<tr>
<td>34</td>
<td>45–260</td>
<td>156–901</td>
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<tr>
<td>36</td>
<td>46–350</td>
<td>159–277</td>
</tr>
<tr>
<td>38</td>
<td>59–570</td>
<td>214–1976</td>
</tr>
<tr>
<td>40</td>
<td>90–460</td>
<td>306–1595</td>
</tr>
</tbody>
</table>

Levels peak in the middle or late afternoon. The day-to-day variation is 12%–15%.

**Procedure**

1. Obtain a 5-mL serum sample by venipuncture using a red-topped tube. Draw the specimen at same time of day on each visit. Observe standard precautions. Record weeks of gestation on the requisition or computer screen. Serial measurements may be recommended to establish a trend.
2. Collect 24-hour urine specimens (Estriol: 13–42 mg/24h or 46–164 nmol/d) during the third trimester.

**Clinical Implications**

1. Decreased E₃ is associated with risk for:
   a. Growth retardation
   b. Fetal death
   c. Fetal anomalies (Down's syndrome, fetal encephalopathy)
d. Fetus past maturity
e. Preeclampsia
f. Rh immunization

2. Decreased $E_3$ also occurs in:
a. Anemia
b. Diabetes
c. Malnutrition
d. Liver disease
e. Hemoglobinopathy

Interfering Factors: Administration of radioactive isotopes within the previous 48 hours interferes with this test.

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedures. Serial testing may be required. See Hormone Testing on page 976.
2. No fasting is necessary.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and monitor appropriately. Continuously low $E_3$ values are sometimes seen in normal pregnancy. A decreasing trend is indicative of fetal distress. Provide counseling and support.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Human Placental Lactogen (hPL) (Chorionic Somatomammotropin)

hPL is a growth-promoting hormone of placental origin and is similar to hCG (see Hormone Testing on page 976).

This test is used to evaluate placental function as an index of fetal well-being in at-risk pregnancies. Low hPL levels are associated with intrauterine growth retardation. Falling levels indicate a poor prognosis. The level of hPL correlates best with placental weight, but the clinical significance of this hormone is controversial.

Reference Values

Normal
- Normal maternal serum: <0.5 µg/mL (mg/L or <25 nmol/L)
- Men and nonpregnant women: nondetectable

Procedure

1. Obtain a serum sample of at least 1 mL in two separate vials (red-topped tube) by venipuncture. Observe standard precautions.
2. Record the week of gestation or last menstrual period (LMP) on the test requisition or computer screen. These tests are usually done as serial measurements.

Clinical Implications

1. Normal values are associated with normal intrauterine growth but do not ensure lack of complications.
2. Decreased or falling values are associated with:
   a. Growth retardation
   b. Placental disease
   c. Fetal death
   d. Hypertensive state
3. Low levels are also associated with some normal pregnancies.
4. Increased values are found in trophoblastic tumors.

Interfering Factors: Administration of radiopharmaceuticals 24 hours before venipuncture interferes with this test.

Interventions

Pretest Patient Preparation

1. Explain the reason for testing the mother's blood.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately. Explain possible need for serial blood testing if results are abnormal.
2. Use ultrasound studies to assess any abnormal results.
3. See Chapter 1 guidelines for safe, effective, informed posttest care.

Fetal Fibronectin (fFN)

Fetal fibronectin is abundant in amniotic fluid and may be useful in the diagnosis of ruptured membranes. The detection...
of fFN in vaginal secretions before membrane rupture may be a marker for impending preterm labor within the next 7 to 14 days.

This test helps to predict a preterm delivery when the presenting symptoms are questionable so that early intervention (e.g., tocolytics, corticosteroids, transport to a tertiary center) can be initiated when indicated. This test is for women with intact membranes and cervical dilation of <3 cm. fFN is secreted in early pregnancy to help attach the fertilized egg to the implantation site in the uterus, but it is not secreted after 22 weeks until near term. This test detects preterm labor from 24 until 34 weeks' gestation.

Reference Values
- **Normal** Negative (<0.050 µg/mL): delivery is unlikely to occur within 14 days. Positive (>0.050 µg/mL): delivery within 7–14 days.

Procedure
1. Using a sterile speculum, obtain secretions from the cervix and vagina by rotating a sterile Dacron swab near the outside of the cervix and the posterior fornix of the vagina. Observe standard precautions.
2. Place specimen in a biohazard bag and send the specimen to laboratory. Results may take 24 to 48 hours.

Clinical Implications A level of fFN equal to or greater than a reference value (0.050 µg/mL) is considered positive and means that preterm labor is imminent. Transvaginal cervical ultrasonography may be used along with fFN to assess risk for preterm birth. A short cervix (<25 mm) with a positive fFN is a strong predictor of preterm delivery. The greatest value of this testing is a negative result in order to avoid unnecessary interventions.

Interventions
**Pretest Patient Preparation**
1. Explain test purpose and procedure to the patient.
2. Refer to Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
1. Counsel the patient regarding test results and need for follow-up medication, tocolysis (inhibition of contractions), or preparation for probable delivery.
2. Be sure the mother knows the warning signs of preterm labor.
   - Uterine contractions—a hard feeling over the entire surface of the uterus that lasts 20 seconds or longer. The contractions can be painless. If more than four are felt per hour, notify clinician.
   - Menstrual-like cramps felt low in abdomen; may be constant or come and go
   - Pelvic pressure or fullness in the pelvic area or back of the thighs
   - Backache—a dull pain in the lower back, either constant or rhythmic, that is not relieved by changing positions
   - Persistent diarrhea
   - Intestinal cramps with or without diarrhea
   - Vaginal discharge that is greater than normal or changes in consistency or color (especially if it is pink, bloody, or greenish)
   - A general feeling or sense that something is wrong
3. Explain the possible causes and increased risks associated with preterm labor and birth:
   - Past preterm birth
   - Spontaneous abortion in second trimester
   - Uterine anomaly
   - Diethylstilbestrol exposure
   - Incompetent cervix
   - Hydramnios
   - Bleeding in second and third trimester
   - Preterm labor
   - Premature rupture of membrane
   - Multiple gestation
   - Preterm cervical dilatation > 2 cm (multipara) or > 1 cm (primipara)
   - Pregnancy weight < 115 pounds
   - Mother < 15 years of age
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

TESTS TO PREDICT FETAL OUTCOME AND RISK FOR INTRAUTERINE ASPHYXIA

**Contraction Stress Test (CST)**
This test is done in a hospital or clinic setting to assess fetal heart rate (FHR) in response to uterine contractions via electronic fetal monitoring.

Reference Values
- **Normal** The test result is negative if there are no late decelerations associated with at least three contractions within a 10-minute period. A normal (negative) CST implies that placental support is adequate; that the fetus is probably able to tolerate the stress of labor, should it begin within 1 week; and that there is a low risk for intrauterine death due to hypoxia.

Procedure
1. Obtain the FHR by using an external transducer.

Clinical Implications A positive result indicates increased risk for intrauterine death due to hypoxia. See further discussion of results and contraindications under Oxytocin Challenge Test (discussed below).

Interventions

Pretest Patient Preparation

1. Explain the reason for testing.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately. Explain possible need for follow-up testing.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Oxytocin Challenge Test (OCT); Nipple Stimulation Test; Breast Stimulation Test (BST)

These tests are performed after 28 weeks of gestation, when a nonstress test (NST) is nonreactive or a contraction stress test (CST) is either positive or unsatisfactory. Continuous external fetal monitoring is used. Because uterine contractions are associated with a reduction in uteroplacental blood flow, spontaneous, oxytocin-induced (OCT), or nipple stimulation-induced contractions with a frequency of 3 in 10 minutes may be used clinically as a standard test of fetoplacental respiratory function. Stress of this magnitude has been proven clinically useful in separating fetuses with suboptimal oxygen reserve from those with adequate reserve (the vast majority), and it does not significantly compromise the normal fetus.

Reference Values

Normal The test result is negative if there are no late decelerations associated with at least three contractions within a 10-minute period. A normal (negative) result is reassuring; it implies that placental reserve is sufficient should labor begin within 1 week. There is a false-normal rate of 1 to 2 per 1000 pregnancies. The procedure is usually repeated weekly.

Procedure

1. Be aware that contractions may occur spontaneously, after breast stimulation (BST), or after administration of intravenous oxytocin (OCT) to produce three good-quality contractions of at least 40 seconds’ duration each, within a 10-minute period.
2. Monitor the FHR for reaction to this stress.

Clinical Implications

1. The presence of consistent and persistent late decelerations with most uterine contractions, regardless of their frequency, constitutes a positive (abnormal) OCT result. This is often associated with decreased baseline FHR variability, a lack of FHR acceleration with fetal movement, and a fetus at risk for intrauterine asphyxia.
2. The results of OCT can be categorized as follows:
   a. Negative: no late decelerations
   b. Positive: late decelerations following 50% or more of contractions, even if the frequency of the contractions is less than three in 10 minutes
   c. Equivocal: intermittent, late, or variable decelerations
   d. Unsatisfactory: less than three contractions within 10 minutes or a poor-quality tracing

Interventions

Pretest Patient Preparation

1. Explain test purpose and procedure.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel accordingly about meaning of fetal heart activity and movement.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

1. With all methods of OCT/CST, there is a risk for hyperstimulation, which could result in extended FHR decelerations that could be hypoxic for the fetus.
2. Contraindications for OCT/CST include:
   a. Third-trimester bleeding (unexplained vaginal bleeding)
   b. Preterm labor (premature)
   c. Presence of classic uterine incision
   d. Placenta previa

Nonstress Test (NST)

The NST can be performed in a hospital, clinic, or possibly home care setting. Test results reflect the functions of the fetal brain stem, autonomic nervous system, and heart.
Reference Values

**Normal** Negative result: reactive NST American College of Obstetricians and Gynecologists (ACOG) criteria for a reactive NST (with or without stimulation): two or more accelerations of FHR, peaking at least 15 beats/minute above the baseline FHR and lasting at least 15 seconds from baseline to baseline, within a 20-minute period

**Procedure**

1. Assess maternal vital signs, last oral intake (including medicines or street drugs), smoking history, and fetal movement history.
2. Apply external fetal monitor with woman positioned off her back in lateral tilt position.
3. After 26 weeks of gestation, this assessment of the FHR pattern without contractions evaluates fetal oxygenation. Fetal movement may or may not be identified by the woman during the test. If gestation is < 26 to 30 weeks, the fetus may not meet the criteria for a reactive NST yet still be a healthy fetus.
4. It is no longer recommended to feed the woman before this test because of the possibility of emergency delivery. Glucose does not alter the FHR pattern.
5. If unable to elicit fetal heart rate accelerations during NST, acoustic stimulation of a fetus that is not acidotic may evoke fetal heart rate accelerations that seem to predict fetal well-being. An artificial larynx (vibroacoustic stimulator) that is designed for fetal monitoring is placed on the mother’s abdomen, and the stimulus is activated for 1 to 2 seconds. The stimulus may be repeated up to three times for gradually increased durations up to 3 seconds to bring about fetal heart rate accelerations. Use of acoustic stimulation can shorten time needed for reactive NST and reduce false-positive test results.

**Clinical Implications** A nonreactive NST (positive test) consists of fewer than two accelerations of FHR (ACOG criteria). If the fetus does not react within the first 20 minutes, stimulation should be applied. The test is considered nonreactive if, after extension to 40 minutes, the ACOG criteria are not met. This extended testing minimizes the possibility of lack of activity owing to fetal sleep. If the FHR pattern is unclear, the test is considered inconclusive or unsatisfactory.

**Clinical Alert**

1. The NST is a screening test and can easily and safely be done once a week.
2. A nonreactive NST (positive test) should be followed by a CST.
3. Ultrasound studies and a fetal biophysical profile (FBP) may be needed after a nonreactive NST.
4. Nonrepetitive, brief (less than 30 seconds) variable decelerations may be noted in up to 50% of NSTs and do not indicate a compromised fetus or a need for interventions.
5. Repetitive variable decelerations (three in 20 minutes) or decelerations that last 60 seconds or longer indicate nonreassuring fetal heart rate pattern and increased risk for cesarean delivery.

**Interfering Factors** A false-positive result may be caused by fetal sleep, preterm gestation, smoking before the NST, congenital anomalies, or maternal use of drugs such as central nervous system depressants or beta blockers.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcome; counsel appropriately.
2. Advise the patient regarding need for weekly or twice-weekly testing according to physician's orders, if pregnancy history indicates risk factors for antepartum fetal demise. If NST is performed for a single occurrence of decreased fetal movement in uncomplicated pregnancy and reactive NST results, reassure patient that test need not be repeated.
3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Fetal Activity-Acceleration Determination (FAD)**

The fetal activity-acceleration determination (FAD) test often is not distinguished from NST, but it is different. In the FAD, both acceleration of FHR and fetal movement are evaluated.

**Reference Values**

**Normal** Negative result: reactive test Criteria are similar to those for NST, but fetal movement is also required: more than three discrete body or limb movements within 30 minutes. In a reactive test (well-oxygenated fetus), spontaneous accelerations of FHR begin at about the time of onset of fetal movement. This effect expresses the condition of the neurologic system and its effect on fetal movement and FHR.

**Procedure**

1. Remember that the procedure is the same as for the NST.
2. Give the woman a button to push when fetal movement occurs; pushing the button causes a mark to appear on the monitor strip.

**Clinical Implications**

1. A nonreactive FAD is ascertained in the same manner as is the NST. Results are of questionable validity before 30 weeks’ gestation. Follow-up for a nonreactive test should include an ultrasound study to assess fetal movement and tone.
2. A nonreactive FAD (positive result) is associated with greater risk for hypoxia.
Clinical Alert

Fetal movement tends to decrease as gestation progresses.

Interventions

Pretest Patient Preparation

1. Explain reason for testing and fetal heart rate monitoring.
2. This test may be performed in a hospital or clinic setting.
3. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome and counsel appropriately. Explain need for possible follow-up ultrasound.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Continuous Fetal Heart Rate (FHR) Monitoring

Continuous FHR monitoring is done (both before and during labor) to evaluate postterm pregnancy (>42 weeks); after a nonreactive stress or nonstress test; and in the presence of diabetes, preeclampsia, chronic hypertension, or intrauterine growth retardation. Normally, the rate is 100 to 150 beats/minute; accelerations occur with fetal movement, and a return of variable decelerations to baseline occurs with no evidence of decreasing baseline variability or increasing baseline rate.

Fetal Biophysical Profile (FBP) (Biophysical Profile [BPP])

The biophysical profile measurements, used in the later stages of pregnancy, assess fetal well-being. The BPP or FBP is more accurate and provides more information than does the NST alone. It can identify the fetus affected by hypoxia that is at risk for intrauterine distress or death. In high-risk pregnancies, testing usually begins by 32 to 34 weeks of gestation; those with severe complications may require earlier testing, at 26 to 28 weeks.

The FBP uses ultrasound imaging to evaluate five distinct parameters: (1) evidence of FHR (cardiac rate) accelerations (NST); (2) muscle tone; (3) fetal movement; (4) fetal breathing; and (5) volume of amniotic fluid. Based on sonographic evidence during a typical 20- to 30-minute survey, each parameter is assigned a value of 0 to 2 points (2 is optimal). The maximum number of points obtainable is 10; a score of 10 indicates a normal test without evidence of fetal distress. Generally, a score > 8 indicates fetal well-being. The FBP also provides the clinician with valuable information regarding fetal size and position, number of fetuses, placental location and grade, and evidence of specific fetal activities such as micturition and eye movements.

Another version of the FBP, termed the modified biophysical profile, has become a primary mode of antepartum fetal testing. The modified version includes the NST as a measure of fetal acid-base status and the amniotic fluid index (AFI) as a long-term placental function assessment. The modified FBP is normal if the NST is reactive and if the AFI is > 5. Abnormal results include nonreactive NST and AFI = 5.

In some laboratories, Doppler examinations of the umbilical vessels assess utero-fetal blood flow. Abnormal Doppler blood flow studies (umbilical artery velocimetry) may be detected before changes in NST, CST, or FBP are detectable. Abnormal Doppler umbilical artery waveforms become indicative of acidosis, hypoxia, and intrauterine growth retardation, which result in a poor outcome. Doppler velocimetry has demonstrated benefits for fetuses with suspected intrauterine growth retardation.

Reference Values

Normal Fetal well-being score: >8 points, based on normal NST, normal fetal muscle tone, movement, and breathing; and normal volume of amniotic fluid

Procedure

1. Explain test purpose and procedure.
2. Position the patient on her back (as for an obstetric sonogram). Apply a gel (coupling agent) to the skin of the lower abdomen. Then, move the ultrasound transducer across the lower abdominal area to visualize the fetus and surrounding structures.
3. Examining time is usually 30 minutes but may vary because of fetal age or fetal state.
4. A CST or NST is also done at this time (see pages 980 and 982).

Clinical Implications

1. Variables that influence FBP include fetal age, fetal behavioral states, maternal or fetal infection, hypoglycemia, hyperglycemia, and postmaturity.
2. If a fetus < 36 weeks of gestation does not have stable behaviors, a longer test may be needed. Infection may cause absence of FHR reactivity and fetal breathing movements. Frequency of fetal breathing increases during maternal hyperglycemia and decreases with maternal hypoglycemia. Other variables that influence FBP include use of therapeutic or nontherapeutic chemicals. Magnesium sulfate may decrease or eliminate fetal breathing movements and decrease FHR variability. Nicotine can decrease the profile parameters, and cocaine may also decrease the FBP score.
3. When the five major biophysical profile parameters can be observed, the fetus is considered to be free of distress. Generally, a score of 8 points indicates fetal well-being.
4. A score of 6 points is equivocal, and retesting should be done in 12 to 24 hours.
5. A score < 4 indicates the potential for or the existence of fetal distress. This warrants further testing or the consideration of delivery.

**Clinical Alert**

To assess the fetal state properly, a sonographic determination of eye movement and respiration must be done. If no eye movements and no respirations are evident, the fetus is most likely asleep. On the other hand, if rapid eye movement is evident but breathing is absent, the fetus is probably in distress.

**Interventions**

**Pretest Patient Preparation**

1. Explain the test purpose and procedure and include information regarding each part of the test and how it relates to fetal well-being.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes, counsel appropriately, and inform about further testing.
2. Instruct the patient regarding need for weekly or twice-weekly testing, if pregnancy history indicates risk for antepartum fetal demise.
3. Follow Chapter 1 guidelines regarding safe, effective, informed posttest care.

**Fetoscopy**

Fetoscopy allows direct observation of the fetus and facilitates fetal blood or skin sampling. It provides direct visualization of the fetus in 2- to 4-cm segments so that developmental defects can be more accurately identified. The fetal blood sample allows early diagnosis of disorders such as hemophilia A and B that are not amenable to detection through other means. Fetoscopy can also be used for therapeutic interventions such as shunt placement.

**Reference Values**

**Normal**
- Normal fetal development; no evidence of fetal developmental defects
- Negative for hemophilia types A and B and sickle cell anemia

**Procedure**

1. Obtain a properly signed and witnessed consent form.
2. Apply a local anesthetic to the mother's abdominal wall. Meperidine (Demerol), which crosses the placenta, may be given to the mother to quiet the fetus.
3. Use real-time ultrasound to locate the proper maternal abdominal area through which to make a small incision, and then insert the cannula and the trocar into the uterus.
4. After cannulation into the uterus, insert an endoscope (fetoscope), consisting of a fiberoptic light source and a self-focusing lens, and then manipulate for optimal views and fetal tissue sampling (eg, skin, blood, amniotic fluid).

**Clinical Implications**

Abnormal results reveal:

1. Fetal malformation
2. Neural tube defects
3. Sickle cell anemia
4. Hemophilia

**Clinical Alert**

1. Fetoscopy poses an increased risk for spontaneous abortions (5%–10%), preterm delivery (10%), amniotic fluid leakage (1%), and intrauterine fetal death.
2. Fetoscopy is offered only when the woman has a significant risk for producing a child with a major birth defect that can be diagnosed only by this method.

**Interventions**

**Pretest Patient Preparation**

1. The woman (or couple) should receive genetic counseling and a thorough explanation of the procedure and its benefits, risks, and limitations.
2. Antibiotics may be ordered before the procedure to prevent amnionitis. Assess for possible allergies to the drug.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Monitor mother and fetus for several hours after the procedure. Institute proper protocols for dealing with maternal blood pressure and pulse changes, FHR abnormalities, uterine activity, vaginal bleeding, or amniotic fluid leakage. Rh-negative mothers should receive human Rho(D) immune globulin (RhogAM) unless the fetus is also known to be Rh negative. Repeat ultrasound studies should be done to check amniotic fluid volume and fetal viability.
2. Instruct the patient to report any pain, bleeding, infected cannulation site, amniotic fluid leakage, or fever (amnionitis).
3. Interpret test outcomes and counsel appropriately.
Percutaneous Umbilical Blood Sampling (PUBS) (Cordocentesis)

Percutaneous umbilical blood sampling (PUBS) has somewhat replaced fetoscopy because of the risk factors associated with the latter test. PUBS, for which research is ongoing, is probably a safer and easier way to sample blood from the umbilical cord of the fetus in utero. Fetal blood can be examined for hemophilia, hemoglobinopathies, fetal infections, chromosomal abnormalities, fetal distress, fetal drug levels, and other blood studies.

Reference Values
Normal No abnormalities noted (see explanation of test)

Procedure

1. Scan with a real-time ultrasound transducer (placed into a sterile glove) to provide landmarks as a 20- to 25-gauge spinal needle is first inserted into the maternal abdomen and then guided into the fetal umbilical vein, 1 to 2 cm from the cord insertion site on the placenta.
2. Aspirate the fetal blood sample into a syringe containing anticoagulant to prevent clotting of the sample.

Clinical Alert
Risks include transient fetal bradycardia, maternal infection, premature labor, and a 1% to 2% incidence of fetal loss.

Clinical Implications Abnormal blood results reveal:

1. Hemoglobinopathies
2. Hemophilia A or B, other coagulation disorders
3. Fetal infection
4. Chromosome abnormalities, genetic diseases
5. Isoimmunization
6. Metabolic disorders
7. Fetal hypoxia

Interventions

Pretest Patient Preparation

1. Explain the procedure and its purpose, benefits, and risks. Obtain a properly signed and witnessed consent form.
2. Assist with relaxation exercises during the procedure. Antibiotics may be given before the test to prevent infection.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Monitor maternal vital signs and perform external fetal monitoring or an NST. Observe for signs of fetal distress.
2. Perform an ultrasound 1 hour after the procedure to ensure that there is no bleeding at the puncture site.
3. Interpret test outcomes and counsel appropriately about fetal therapy (eg, red blood cell and platelet transfusion and drug treatment).
4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Chorionic Villus Sampling (CVS)

Chorionic villus sampling (CVS) can provide very early diagnosis of fetal genetic or biochemical disorders. CVS involves extraction of a small amount of tissue from the villi of the chorion fondosum. This tissue is composed of rapidly proliferating trophoblastic cells that ultimately form the placenta. Although not a part of the fetus, these villi cells are genetically identical to the fetus and are considered fetal rather than maternal in origin.

CVS differs from amniocentesis in several respects. In amniocentesis, the cells examined are desquamated fetal cells; the cells sampled in CVS divide rapidly and are easier to culture. For this reason, karyotyping (see Chap. 11) can be performed much more rapidly, and diagnostic information can be provided within 24 hours, much faster than with amniotic fluid cells. Also, CVS can be performed much earlier in pregnancy, typically at 7 to 11 gestational weeks, whereas amniocentesis usually is performed after 16 weeks’ gestation, with results available several weeks later. CVS, therefore, has the advantage of providing first-trimester diagnosis, which is of particular value when the choice is made to abort an affected fetus because first-trimester terminations of pregnancy are medically safer.

CVS reveals chromosome abnormalities and fetal metabolic or blood disorders. However, because CVS cannot be used to measure AFP, it cannot detect neural tube defects or other disorders associated with increased AFP levels.

Indications for CVS include the following:

1. Abnormal ultrasound test
2. Fetus at risk for detectable mendelian disorders:
   a. Tay-Sachs disease
   b. Hemoglobinopathies
   c. Cystic fibrosis
   d. Muscular dystrophy
3. Birth of previous child with evidence of chromosome abnormality
4. Parent with known structural chromosomal rearrangement
5. Diagnosis of fetal infection

Reference Values
Normal: Negative for chromosomal and DNA abnormalities
No fetal metabolic enzyme or blood disorders

Procedure (Transcervical Method)

1. Position the mother on her back to permit ultrasound documentation of the number of fetuses in utero and their viability and localization of trophoblastic tissue. Ask the patient either to maintain a full bladder or to empty the bladder so as to optimize the sampling path. A bimanual pelvic examination is often performed concurrently with this preliminary ultrasound examination.
2. Have the patient assume a lithotomy position. Insert a sterile speculum after the vagina has been cleansed with an iodine-based antiseptic.
3. Introduce a sterile flexible catheter with a stainless-steel obturator into the vaginal canal and advance through the cervical canal into the trophoblastic tissue. The catheter is visually tracked by the ultrasound device.
4. Once the catheter is in place, attach syringe to the end of the catheter to extract approximately 5 mL of tissue. Immediately examine the tissue sample under a low-power microscope to determine that both quantity and tissue quality are acceptable.
5. Make up to three passes of the catheter. Use a new, sterile catheter each time. After sufficient tissue has been gathered, use ultrasound again to monitor fetal viability. Use the tissue sample for chromosome and enzyme analysis and for other tests.
6. Be aware that a transabdominal method may also be used. This method is similar to amniocentesis, except that the thin-walled needle is inserted into the chorionic bed.

Clinical Implications
Abnormal CVS results indicate:
1. Abnormal fetal tissue
2. Chromosome abnormalities
3. Fetal metabolic and blood disorders
4. Fetal infection

Interventions
Pretest Patient Preparation
1. Be aware that genetic counseling typically proceeds any CVS procedure.
2. Explain the purpose, procedure, and risks of the test.
3. Ensure that a legal consent form is signed by the mother and the father of the baby and is properly witnessed.
4. Have the patient drink four 8-ounce glasses of water about 1 hour before the examination. The patient should not void until instructed to do so.
5. Obtain baseline measurements of maternal vital signs and FHR.
6. Advise the patient that she may experience cramping as the catheter passes through the cervical canal.
7. Help the patient to relax.
8. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Monitor maternal vital signs and FHR every 15 minutes for the first hour after test completion.
2. Instruct the patient to notify her physician if she experiences abdominal pain, vaginal bleeding or abnormal discharge, elevated temperature, chills, or amniotic fluid leakage.
3. Interpret test outcomes and counsel appropriately. Rh-negative women usually receive RhoGAM.
4. Support the mother and significant others during decision making. Provide opportunity for questions and discussion.
5. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert
1. CVS is not considered a routine alternative to amniocentesis. The safety of the CVS procedure is related to the experience and skill of the examiner. In experienced hands, the rates of complications and fetal loss are only slightly greater than for amniocentesis. Risks include leakage of amniotic fluid, bleeding, intrauterine infection, spontaneous abortion, maternal tissue contamination of specimen, Rh isoimmunization, and fetal death (5%).
2. Transcervical CVS is difficult in patients who have a fundal placental implantation site or an extremely retroflexed or anteflexed uterus. In such patients, a transabdominal approach similar to that used for amniocentesis is employed.
3. CVS cannot detect neural tube defects or other disorders associated with abnormal maternal serum.
4. Some specialists advise that this procedure be reserved for evaluation of conditions that present relatively high genetic risks (eg, hemoglobinopathies).
5. An increased risk for severe limb deformities is associated with this procedure.

AMNIOTIC FLUID STUDIES

The fluid filling the amniotic sac serves several important functions. It provides a medium in which the fetus can readily move, cushions the fetus against possible injury, helps maintain an even temperature, and provides useful information concerning the health and maturity of the fetus. The origin of amniotic fluid is not completely understood. In early pregnancy, it is produced by the amniotic membrane covering the placenta and the cord. As the pregnancy progresses, it
is believed to be primarily a byproduct of fetal pulmonary secretions, urine, and metabolic products from the intestinal tract.

Initially, amniotic fluid is produced from the amniotic membrane cells. Later, most of it is derived from the maternal blood. The volume increases from about 30 mL at 2 weeks' gestation to 350 mL at 20 weeks. After 20 weeks, the volume ranges from 500 to 1000 mL. The volume of amniotic fluid changes continuously because of fluid movement in both directions through the placental membrane. Later in pregnancy, the fetus contributes to amniotic fluid volumes through excretion of urine and swallowing of amniotic fluid. The fetus also absorbs up to 400 mL of amniotic fluid every 24 hours through its gastrointestinal tract, bloodstream, and umbilical artery exchanges across the placenta. Probably, some fluid is also absorbed by direct contact with the fetal surface of the placenta. Amniotic fluid contains castoff cells from the fetus and resembles extracellular fluid with suspended, undissolved material. It is slightly alkaline and contains albumin, urea, uric acid, creatinine, lecithin, sphingomyelin, bilirubin, fat, fructose, epithelial cells, leukocytic enzymes, and lanugo hair.

When amniocentesis is advised early in pregnancy (15 to 18 weeks), the purpose is to study the fetal genetic makeup and to determine developmental abnormalities. Fetal cells are separated from the amniotic fluid by centrifugation and are placed in a tissue culture medium so that they can be grown and harvested for subsequent karyotyping to identify chromosome disorders. Testing in the third trimester is done to determine fetal age and well-being, to study blood groups, or to detect amnionitis.

**Amniocentesis**

Amniotic fluid is aspirated by means of a needle guided through the mother's abdominal and uterine walls into the amniotic sac. Amniocentesis is preferably performed after the 15th week of pregnancy. By this time, amniotic fluid levels have expanded to 150 mL, so that a 10-mL specimen can be aspirated. If the purpose of amniocentesis is to ascertain fetal maturity, it should be done after the 35th week of gestation.

Amniocentesis provides a method to detect fetal abnormalities in situations in which the risk for an abnormality may be high. The test can evaluate fetal hematologic disorders, fetal infections, inborn errors of metabolism, and sex-linked disorders. It is not done to determine the sex of the fetus simply out of curiosity.

Chromosomal abnormalities and neural tube defects such as anencephaly, encephalocele, spina bifida, and myelomeningocele can be determined, as can estimates of fetal age, fetal well-being, and pulmonary maturity.

The development of significant maternal Rh antibody titers or a history of previous erythroblastosis can be an indication for amniocentesis.

**High-Risk Parents Who Should Be Offered Prenatal Diagnosis**

1. Women of advanced maternal age (>35 years) who are at risk for having a child with a chromosome abnormality, especially trisomy 21. At maternal age 35 to 40 years, the risk for Down's syndrome is 1% to 3%; at age 40 to 45, it is 4% to 12%; and at > 45 years, the risk is 12% or greater.
2. Women who have previously borne a trisomic child or a child with another kind of chromosome abnormality.
3. Parents of a child with spina bifida or anencephaly or a family history of neural tube disorders.
4. Couples in which either parent is a known carrier of a balanced translocation chromosome for Down's syndrome.
5. Couples in which both partners are carriers for a diagnosable metabolic or structural autosomal recessive disorder.
6. Couples in which either partner or a previous child is affected with a diagnosable metabolic or structural dominant disorder.
7. Women who are presumed carriers of a serious X-linked genetic disorder.
8. Couples from families whose medical history reveals mental retardation, ambiguous genitalia, or parental exposure to toxic environmental agents (eg, drugs, irradiation, infections).
9. Couples whose personal and family medical history reveals multiple miscarriages, stillbirths, or infertility.
10. Parents with anxiety about the health status of potential offspring.
11. Women with abnormal ultrasound results.

**Clinical Alert**

The in utero diagnosis of many genetic disorders may lead the parents to consider abortion as an option for dealing with an unfavorable situation. Because this can be a very difficult and controversial choice, communication between the parents and the health care team must take place in a nonjudgmental, nonthreatening manner.

**Reference Values**

*Normal* Normal amniotic fluid constituents and properties vary according to the age of fetus and the laboratory methods used; pH is slightly alkaline. See descriptions of individual tests.

**Procedure (in Combination with Ultrasound)**

1. Position the patient on her back with her arms behind her head to prevent touching of the abdomen and the sterile field during the procedure (see Obstetric Sonogram in Chap. 13).
2. Perform ultrasound scanning before the procedure to assess fetal number, viability, and position. An appropriate pocket of amniotic fluid is localized in the scan. The tap site should be located away from the fetus, from the site of umbilical cord insertion, and from any thick placental segments.
3. Cleanse the skin thoroughly with an appropriate antiseptic solution and properly drape with sterile drapes. Inject a local anesthetic slowly at the puncture site.
4. Advance a 3.5-inch spinal needle (20- to 22-gauge) with stylet through the abdominal and uterine walls into the amniotic sac but away from the fetus and, when possible, from the placenta. Use continuous ultrasound...
surveillance to track the position of the fetus. Should the fetus move close to the needle, withdraw the needle.
5. Once the needle is properly positioned, remove the stylet and attach a syringe to the needle to permit aspiration of
a 20- to 30-mL specimen. Discard the first 0.5 mL of aspirated fluid to prevent contamination by maternal cells or
blood.
6. Withdraw the needle, and place an adhesive bandage over the puncture site. Postprocedure ultrasound scanning
confirms fetal viability.
7. Place the amniotic fluid specimen in a sterile brown or foil-covered silicone container to protect it from light and
thereby prevent breakdown of bilirubin. Label the container properly. Include the estimated weeks of gestation and
the expected delivery date. Deliver the sample to the laboratory immediately.
8. Be aware that the laboratory workup for genetic diagnoses usually takes 2 to 4 weeks to complete. However,
specimens obtained for determination of fetal age (eg, creatinine) take 1 to 2 hours; determinations of the
lecithin-to-sphingomyelin (L/S) ratio and phosphatidyl glycerol take 3 to 4 hours; Gram stain to rule out infection
takes one-half hour, and cultures take 48 to 72 hours.
9. The procedure may have to be repeated if no amniotic fluid is obtained or if there is failure of cell growth or culture
results are negative.
10. Record the type of procedure done, date, time, name of physician performing the test, maternal-fetal response, and
disposition of specimen.

Clinical Implications

1. Elevated amniotic fluid alpha-fetoprotein (AFP) can indicate possible neural tube defects as well as multiple
gestations, fetal death, abdominal wall defects, teratomas, Rh sensitization, and fetal distress.
2. Decreased AFP is associated with fetal trisomy 21 (Down’s syndrome).
3. Creatinine levels are reduced in fetal prematurity. At 37 weeks of gestation, creatinine in amniotic fluid should be >2
mg/dL (>15 µmol/L). 
4. Increased or decreased total amniotic fluid volumes are associated with certain types of arrested fetal development.
5. Increased bilirubin levels are associated with impending fetal death. (See page 1005 for normal values.)
6. Amniotic fluid color changes are associated with fetal distress and other disorders such as chromosome
abnormalities.
7. Sickle cell anemia and thalassemia can be detected through analysis of amniotic fibroblast DNA.
8. X-linked disorders are not routinely diagnosed in utero. However, because these disorders affect only men, the fetal
sex may need to be determined when the mother is a known carrier of the X-linked gene in question (eg, hemophilia, Duchenne's muscular dystrophy).
9. Screening for carrier state or affected fetus is done through chromosomal testing.
10. The presence of some of the more than 100 detectable metabolic disorders can be detected in the amniotic fluid
sample. Examples include Tay-Sachs disease, Lesch-Nyhan syndrome, Hunter's syndrome, Hurler's syndrome, and
various hemoglobinopathies. Hereditary metabolic disorders are caused by absence of an enzyme due to deletion
or by alteration of the structure or synthesis of an enzyme due to gene mutation. If the enzyme in question is
expressed in amniotic fluid cells, it can potentially be used for prenatal diagnosis. An unaffected fetus would have a
normal enzyme concentration, a clinically normal carrier of the gene defect would have perhaps half of the normal
enzyme level, and an affected fetus would have a very small amount or none of the enzyme.
11. For disorders in which an abnormal protein is not expressed in amniotic fluid cells, other test procedures are
necessary, such as DNA restriction endonuclease analysis.

Interfering Factors

1. Fetal blood contamination can cause false-positive results for AFP.
2. False-negative and false-positive errors in karyotyping can occur.
3. Polyhydramnios may falsely lower bilirubin values as a result of dilution.
4. Hemolysis of the specimen can alter test results.
5. Oligohydramnios may falsely increase some amniotic fluid analysis values, especially bilirubin; this can lead to
errors in predicting the clinical status of the fetus.

Interventions

Pretest Patient Preparation

1. Ensure that elective genetic counseling includes a discussion of the risk for having a child with a genetic defect and
problems (eg, depression, guilt) associated with selective abortion. The father should be present and should be a
partner in the decision-making process. In genetic counseling, do not coerce the parents into undergoing abortion
or sterilization; this should be an individual choice.
2. Explain test purpose, procedure, and risks; assess for contraindications.
3. Ensure that a properly signed and witnessed legal consent form is obtained.
4. Instruct the patient to empty her bladder just before the test.
5. Obtain baseline measurements of fetal and maternal vital signs. Monitor fetal signs for 15 minutes.
6. Alert the patient to the possibility that transient feelings of nausea, vertigo, and mild cramping may occur during the
procedure. Help the patient to relax.
7. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Check maternal blood pressure, pulse, respiration, and fetal heart tone every 15 minutes for the first half hour after
test completion. Palpate the uterine fundus to assess fetal and uterine activity; monitor for 20 to 30 minutes with an
external fetal monitor, if one is available.
2. Position the mother on her left side to counteract supine hypotension and to increase venous return and cardiac
output.
3. Instruct the patient to notify her physician if she experiences amniotic fluid loss, signs of onset of labor, redness and inflammation at the insertion site, abdominal pain, bleeding, elevated temperature, chills, unusual fetal activity, or lack of fetal movement.

4. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

1. Fetal loss attributable to the procedure is <0.5%. Repeat amniocentesis is necessary in 0.1% of cases.

2. Fetal complications include:
   a. Spontaneous abortion
   b. Injury to the fetus (fetal puncture)
   c. Hemorrhage
   d. Infection
   e. Rh sensitization if fetal blood enters the mother's circulation

3. Maternal complications include:
   a. Hemorrhage
   b. Hematomas

4. This test is contraindicated in women with a history of premature labor or incompetent cervix and in the presence of placenta previa or abruptio placentae. If the amniotic fluid is bloody (blood is usually of maternal origin) and if a significant number of fetal cells (Kleihauer-Betke–positive smear) are present in the amniotic fluid of an Rh-negative mother, administration of RhoGAM should be considered. Some doctors prefer to administer RhoGAM to all Rh-negative mothers for amniocentesis, unless they are already sensitized at that time.

5. Families need to know that prenatal diagnoses based on amniotic fluid assay are not infallible; sometimes, results do not reflect the true fetal status. Findings from amniocentesis cannot guarantee a normal or abnormal child; they can only determine the relative likelihood of specific disorders within the limits of laboratory measurements. Some conditions cannot be predicted by this method, including nonspecific mental retardation, cleft lip and palate, and phenylketonuria (PKU).

6. Accurate and optimally safe results from amniocentesis are possible only if the following protocols are observed:
   a. Gestation =15 weeks
   b. Ultrasound monitoring to locate suitable pools of amniotic fluid, outline the placenta, exclude the presence of a multiple pregnancy, and accurately estimate fetal maturity. These considerations are necessary to correctly interpret AFP values in amniotic fluid and maternal blood.
   c. Precise and meticulous amniocentesis technique, including use of 20- or 22-gauge needle
   d. Maximum of two needle insertion attempts for a single tap
   e. Administration of RhoGAM for the Rh-negative woman

7. Cytogenetic analysis can produce results that are 99.8% accurate.

8. Techniques have been developed for performing amniocentesis in the presence of twin fetuses. Amniotic fluid is aspirated from one of the amniotic sacs, and a small amount of contrast material is injected into the sac. When the adjacent sac is tapped and produces clear amniotic fluid, the clinician is assured that each sac has been tapped and each fetus will be accurately assessed.

9. An anteriorly located placenta does not preclude amniocentesis. A thin portion of placenta can be traversed during amniocentesis with no apparent increase in postamniocentesis complications.

Amniotic Fluid Alpha Fetoprotein (AFP)

AFP is synthesized by the embryonic liver and is the major protein (glycoprotein) found in fetal serum. It resembles albumin in molecular weight, amino acid sequence, and immunologic characteristics. However, it is not normally detectable after birth. Ordinarily, high levels of fetoproteins are found in the developing fetus, and low levels exist in maternal serum and amniotic fluid.

The amniotic fluid AFP test is used to diagnose fetal neural tube defects (malformations of the central nervous system); fetoprotein leaks into the amniotic fluid during such pregnancies. The causes of neural tube defects are not known; however, a genetic component is assumed because an increased risk for recurrence exists. Neural tube defects usually exhibit polygenic (multifactional) traits. In cases of anencephaly and open spina bifida, both maternal serum alpha-fetoprotein (MS-AFP) and amniotic fluid AFP concentrations are abnormal by the 18th week of gestation.

Additionally, AFP measurements have been used as indicators of fetal distress; in such cases, both amniotic fluid AFP and MS-AFP may be increased. However, final confirmation must come from further studies.

Reference Values

**Normal** 12–16 weeks Peak at 12–16 weeks is 14.5 µg/L or 196 pmol/L. Values vary considerably according to age of fetus and laboratory methods used. Values peak at 12–16 gestational weeks and then gradually decline to term.

**Procedure** In the laboratory, amniotic fluid is analyzed for concentration of AFP.

**Clinical Implications** Increased amniotic AFP levels are associated with:

1. Neural tube defects such as anencephaly (100% reliable), encephalocele, spina bifida, and myelomeningocele (90% reliable)
2. Congenital nephrosis
3. Omphalocele
4. Turner's syndrome with cystic hydromas
5. Gastrointestinal tract obstruction
6. Missed abortion
7. Fetal distress
8. Imminent or actual fetal death
9. Severe Rh immunization
10. Esophageal or duodenal atresia
11. Fetal liver necrosis secondary to herpes virus infection
12. Sacrococcygeal teratoma
13. Spontaneous abortion
14. Trisomy 13
15. Urinary obstruction (eg, fetal bladder neck obstruction with hydronephrosis)
16. Cystic fibrosis

Interfering Factors

1. Fetal blood contamination causes increased AFP.
2. Increased AFP is associated with multiple pregnancies.
3. False-positive (0.1%–0.2%) results may be associated with fetal death, twins, or genetic anomalies, but sometimes no explanation can be given for the results.

Clinical Alert

1. Any parents who have already produced a child with a neural tube defect should be offered antenatal studies in anticipation of future pregnancies. If one parent has spina bifida, the pregnancy should be closely monitored.
2. High-resolution ultrasound studies must be used to confirm increased AFP levels.

Interventions

Pretest Patient Preparation

1. Explain the test purpose and the meaning of positive and negative test results.
2. Provide for genetic counseling.
3. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes, counsel, and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Amniotic Fluid Total Volume

Measurement of amniotic fluid total volume is helpful for estimating the changes in total amounts of certain substances that circulate in the amniotic fluid, including bilirubin, creatinine, and surface-active agents. Knowledge of total amniotic fluid volume is important because marked changes in the amount of amniotic fluid can decrease the predictive value of serial concentration measurements of specific substances. This measurement is most important when test results do not agree with the clinical picture.

Reference Values

<table>
<thead>
<tr>
<th>Weeks of Gestation</th>
<th>Average Volume (mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12</td>
<td>Approximately 50</td>
</tr>
<tr>
<td>15</td>
<td>350</td>
</tr>
<tr>
<td>20</td>
<td>450</td>
</tr>
<tr>
<td>25</td>
<td>750</td>
</tr>
<tr>
<td>30 to 35</td>
<td>1500</td>
</tr>
</tbody>
</table>

After 35 weeks, values decrease to 1250 mL at term.

Procedure

1. Study a sample of amniotic fluid with the use of a solution of paraaminohippuric acid (PAH) for absorbency and dilution to calculate the probable amniotic fluid volume in milliliters.
2. Correct amniotic fluid total volume by multiplying the measured levels of specific substance by the actual fluid volume divided by average volume (for gestation age).

Clinical Implications

1. Polyhydramnios (increased amniotic fluid, >2000 mL) is suggested by a total intrauterine volume > 2 standard deviations above the mean for a given gestational age. It is estimated that 18% to 20% of fetuses in such pregnancies have congenital anomalies, the two most common being anencephaly and esophageal atresia (fetal swallowing is greatly impaired). The remainder have involvement secondary to Rh disease, diabetes, or other, unknown causes. Polyhydramnios is also associated with multiple births (eg, twins).
2. Oligohydramnios (reduced volume of amniotic fluid, <300 mL) is suggested by a total intrauterine volume > 2 standard deviations below the mean occurring before the 25th week of gestation. A disturbance of kidney function caused by renal agenesis or kidney atresia can result in oligohydramnios (fetal urination is impaired). After 25 weeks, the suspected causes of decreased amniotic fluid volume are premature rupture of membranes, intrauterine growth retardation, and postterm pregnancy.
If either polyhydramnios or oligohydramnios is suspected, the fetus should be screened with ultrasound to detect physical anomalies.

**Interventions**

**Pretest Patient Preparation**

1. Explain the reason for amniotic fluid testing and the meaning of results.
2. See Chapter 1 guidelines for [safe, effective, informed pretest care](#).

**Posttest Patient Aftercare**

1. Interpret amniotic fluid test results and monitor appropriately.
2. Follow Chapter 1 guidelines for [safe, effective, informed posttest care](#).

**Amniotic Fluid Index (AFI)**

**Reference Values**

*Normal* At term, the AFI is usually between 8 and 18 cm. Values < 5 cm indicate oligohydramnios, and those > 24 cm indicate polyhydramnios.

**Procedure**

1. The pregnant woman lies supine with displacement of the uterus to the left. The abdomen is divided into four quadrants.
2. Ultrasound is used to locate the largest pocket of amniotic fluid in each of the four quadrants, and each pocket is measured vertically. The four values are added together to obtain the AFI. The advantage of this test is that serial follow-up measurements can be done.

**Clinical Implications**

1. Oligohydramnios and polyhydramnios are indicators of poor outcome in pregnancy.
2. An AFI lower than the 2.4 percentile for a certain gestational age is considered to represent oligohydramnios.
3. Oligohydramnios can indicate chronic uteroplacental insufficiency or renal anomaly.
4. An AFI higher than the 97.5 percentile for a certain gestational age is considered to indicate polyhydramnios.

Polyhydramnios is associated with upper gastrointestinal tract obstruction or malformation (eg, tracheoesophageal fistula, hydrops fetalis).

**Interfering Factors** False-positive results can occur in a severely dehydrated woman.

**Interventions**

**Pretest Patient Preparation**

1. Explain the reason for the AFI procedure.
2. See Chapter 1 guidelines for [safe, effective, informed pretest care](#).

**Posttest Patient Aftercare**

1. Explain the test results to the patient. Prepare the patient for follow-up procedures or need for delivery of the infant.
2. Follow Chapter 1 guidelines for [safe, effective, informed posttest care](#).

**Amniotic Fluid Creatinine**

Creatinine, a byproduct of muscle metabolism found in amniotic fluid, reflects increased fetal muscle mass and the ability of the maturing kidney (ie, glomerular filtrating system) to excrete creatinine into the amniotic fluid. The amniotic fluid creatinine concentration progressively increases as pregnancy advances. The mother's blood creatinine level should be known before the amniotic fluid creatinine value is interpreted.

Creatinine indicates fetal physical maturity and correlates reasonably well with the level of lung maturity. Normal lung development is dependent on normal kidney development. As pregnancy progresses, the amniotic fluid creatinine level increases. A value of 2 mg/dL (177 µmol/L) is accepted as an indicator that gestation is at 37 weeks or more. However, the use of this value alone to assess maturity is not advised for several reasons. A high creatinine concentration may reflect fetal muscle mass but not necessarily kidney maturity. For example, a large fetus of a diabetic mother may have high creatinine levels because of increased muscle mass. Conversely, a small, growth-retarded infant of a hypertensive mother may have low creatinine levels because of decreased muscle mass. Creatinine levels can be misleading if they are used without other supporting data. So long as maternal blood creatinine levels are not increased, amniotic fluid creatinine measurements have a certain degree of reliability if they are interpreted in conjunction with other maturity studies.

**Reference Values**

*Normal* A value > 2 mg/dL or > 177 µmol/L indicates fetal maturity (at 37 weeks) if maternal creatine is normal.

**Procedure**

1. Obtain an amniotic fluid sample of at least 0.5 mL.
2. Protect the specimen from direct light.
3. Obtain maternal venous blood sample.

**Clinical Implications** Creatinine levels lower than expected may occur in the following situations:

1. Early in the gestational cycle (not yet at 37 weeks)
2. Fetus smaller than normal (growth retarded)
3. Fetal kidney abnormalities
4. Prematurity

**Interfering Factors** Causes of elevated amniotic fluid creatinine concentrations that are not consistent with gestational age include abnormal maternal creatinine, diabetes, and preeclampsia.

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose of the test.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Amniotic Fluid Lecithin-to-Sphingomyelin (L/S) Ratio (Surfactant Components)**

Lecithin and sphingomyelin have detergent (surfactant) ability. These substances, produced by lung tissue, stabilize the neonatal alveoli to prevent their collapse on expiration and consequent atelectasis. The amount of lecithin in amniotic fluid is less than the amount of sphingomyelin until 26 weeks of gestation; at 30 to 32 weeks of gestation, the two lipid values are about equal. At 35 weeks, lecithin level rises abruptly, but sphingomyelin stays constant or decreases slightly. Saturated phosphatidylcholine, a subfraction of total lecithins, is a major surface-active component of lung surfactant.

The relationship between the phospholipids and the surface-active agents, lecithin and sphingomyelin, is used as an index of fetal lung maturity. If early delivery is anticipated because of conditions such as diabetes, premature rupture of membranes, maternal hypertension, placental insufficiency, or erythroblastosis (Rh disease), the L/S ratio can be used to predict whether the fetal lung will function properly at birth. When early delivery is necessary for fetal viability, the result may be prematurity, pulmonary immaturity, or perinatal mortality. The L/S ratio should be determined on all repeat cesarean sections before delivery to ascertain when fetal lungs are functionally mature. Sphingomyelin exhibits surface-active properties in the lung but plays no role in the surfactant system except to be used as a convenient marker.

**Reference Values**

**Normal** A ratio of 2:1 or greater indicates pulmonary maturity

**Procedure**

1. Withdraw at least 3 mL of amniotic fluid, or collect from a free flow of fluid from the vagina in cases of ruptured membranes.
2. Centrifuge the fluid and prepare for analysis, and read the results in a reflectance densitometer. Calculate the L/S ratio.

**Clinical Implications**

1. A decreased L/S ratio (<1.5:1) is often associated with pulmonary immaturity and respiratory distress syndrome (RDS).
2. An L/S ratio > 2:1 signifies fetal lung maturity. The occurrence of RDS is extremely unlikely.
3. An L/S ratio between 1.5:1 and 1.9:1 indicates possible mild-to-moderate RDS (50% risk).
4. Fetuses of women with insulin-dependent diabetes develop RDS at higher ratios. The L/S ratio should be >3.5:1 for these infants.
Clinical Alert

1. If the L/S ratio is < 1.5:1, it is preferable to delay induced delivery until the fetal lung becomes more mature.
2. Fetal lung maturity appears to be regulated by hormonal factors, some stimulatory and others possibly inhibitory. For this reason, hormones such as betamethasone (Celestone) are given in two doses, administered 12 to 18 hours apart, if premature labor occurs.
3. Under certain stressful conditions, premature fetal lung maturation may be seen. This accelerated fetal lung maturation is thought to be a protective mechanism for the preterm fetus should delivery actually occur.
   a. Premature rupture of the membranes. Prolonged rupture of the membranes (after 72 hours) has an acute negative effect on lung maturation.
   b. Acute placental function
   c. Placental insufficiency
   d. Chronic abruptio placentae
   e. Renal hypertensive disease caused by degenerative forms of diabetes
   f. Cardiovascular hypertensive disease associated with drug abuse
   g. Severe pregnancy-induced hypertension
4. Delayed fetal lung maturation may be seen in the following conditions; in these instances, a higher L/S ratio (>3:5) may be necessary to ensure adequate fetal lung maturity:
   a. Infants born to mothers with insulin-dependent diabetes
   b. Infants born to mothers with nonhypertensive glomerulonephritis
   c. Hydrops fetalis
5. A lung profile of amniotic fluid to evaluate lung maturity looks not only for lecithin but also for two other phospholipids—phosphatidyl glycerol (PG) and phosphatidylinositol (PI). PI increases in the amniotic fluid after 26 to 30 weeks of gestation, peaks at 35 to 36 weeks, and then decreases gradually. PG appears after 35 weeks and continues to increase until term; measurements are classified as positive PG or negative PG. The lung profile is a useful adjunct in evaluating the L/S ratio. It appears that lung maturity can be confirmed in most pregnancies if PG is present (positive) in conjunction with an L/S ratio of 2:1. PG may provide stability that makes the infant less susceptible to RDS when experiencing hypoglycemia, hypoxia, or hypothermia. The PG measurement is especially useful in borderline cases and in class A, B, and C diabetes when pulmonary maturation is delayed.

Interfering Factors

1. High false-negative result rates
2. Unpredictability or borderline values
3. Unpredictability of contaminated blood specimens
4. Occasional false-positive values associated with conditions such as Rh disease, diabetes, or severe birth asphyxia.

Interventions

Pretest Patient Preparation

1. Explain the reason for testing and the meaning of results.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Amniotic Fluid Shake Test (Foam Stability Test)

The shake test is a qualitative measurement of the amount of pulmonary surfactant contained in the amniotic fluid. It is quick and inexpensive. It is a “bedside test” of lung maturity. In an obstetric emergency, an immediate decision about delivery can be made. The advantage of this test over the L/S ratio is that a physician, technician, or nurse can perform it and the results are highly reliable. The L/S ratio usually is not determined when the shake test is positive because the shake test also indicates fetal maturity. A table of dilutions is used to determine the stage of lung maturity.

Reference Values

Normal Positive: persistence of a foam ring for 15 minutes after shaking (at an amniotic fluid-alcohol dilution of 1:2) indicates lung maturity

Procedure

1. Remember that the test is based on the ability of amniotic fluid surfactant to form a complete ring of bubbles on the surface of the amniotic fluid in the presence of 95% ethanol.
2. Place a mixture of 95% ethanol and amniotic fluid in an appropriate container and shake for 15 seconds. A commercial kit may be used.

Clinical Implications

1. If a complete ring of foam forms and persists for 15 minutes, the test is positive.
2. If no ring of bubbles forms, the test is negative.
3. The test has a high false-negative rate but a low false-positive rate. The L/S ratio must be > 4:1 for this test to be positive.

Interfering Factors
1. Blood or meconium contamination can alter results.
2. Contamination of glassware or reagents can alter test results.

**Interventions**

**Pretest Patient Preparation**

1. Explain the reason for testing and the meaning of results.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test outcomes and counsel appropriately.
2. See Chapter 1 guidelines for safe, effective, informed posttest care.

**Amniotic Fluid Foam Stability Index (FSI)**

The foam stability index (FSI) is a modification of the shake test. It provides a functional measurement of fetal lung maturity based on the surface tension properties of surfactant phospholipids.

**Reference Values**

**Normal** FSI: >0.47

**Procedure**

1. Mix a fixed amount of undiluted amniotic fluid with increasing volumes of ethanol.
2. Shake the sample and observe for foam.
3. Document the largest column of ethanol in which the amniotic fluid can form and support foam. This test is almost as reliable as the L/S ratio in normal pregnancies, and it seems to have a lower false-positive rate than the shake test.

**Clinical Implications**

FSI of >0.48 is termed mature; a value of <0.46 is termed immature.

**Interfering Factors**

1. Blood or meconium contamination can produce a false mature result.
2. The test is not reliable for amniotic fluid collected from the vagina.

**Amniotic Fluid Fern Test**

Fern production is a result of the concentration of electrolytes, especially sodium chloride, in the cervical glands; it is under the control of estrogen. Close to term, amniotic fluid shows a typical fern pattern similar to that seen in cervical mucus; this indicates a predominantly estrogen effect rather than progesterone.

This study differentiates urine from amniotic fluid. It is done to determine whether the fluid passed is urine or prematurely leaked amniotic fluid. This is a relatively fast and inexpensive test that can be easily done.

**Reference Values**

**Normal** Positive test for presence of amniotic fluid

**Procedure**

1. Perform a vaginal examination with the use of a sterile speculum.
2. Place a few drops of fluid on a slide and allow to dry.
3. Look for a fern or “palm leaf” pattern (arborization) under the microscope.

**Clinical Implications**

1. A positive test shows the fern pattern indicative of amniotic fluid.
2. A negative test shows no ferning or crystallization; this indicates little or no estrogen effect.
3. No fern pattern is seen if the specimen is urine.

**Interfering Factors**

Blood contaminating the specimen inhibits fern formation.

**Clinical Alert**

Urine can also be differentiated from amniotic fluid if the fluid is tested for the presence of urea, nitrogen, potassium, and creatinine and the absence of AFP.

**Interventions**

**Pretest Patient Preparation**

1. Explain test purpose and procedure.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
1. Interpret test results and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Amniotic Fluid Color**

Amniotic fluid specimens may vary from no color to a pale straw-yellow color. White particles of vernix caseosa from fetal skin and lanugo hair may be present. In certain disorders such as missed abortion, chromosomally abnormal fetus, and fetal anencephaly, the amniotic fluid color is altered.

<table>
<thead>
<tr>
<th>Clinical Alert</th>
</tr>
</thead>
<tbody>
<tr>
<td>Every amniotic fluid specimen should be visually inspected for color.</td>
</tr>
</tbody>
</table>

**Clinical Implications**

1. **Yellow** amniotic fluid indicates blood incompatibility, erythroblastosis fetalis, or presence of bile pigment released by red blood cell hemolysis (fetal bilirubin).
2. **Dark yellow** aspirate indicates probable fetal involvement.
3. **Red** color indicates blood, in which case it must be determined whether the blood is from the mother or the fetus. Fetal blood in the amniotic fluid is of grave concern.
4. **Green, opaque** fluid indicates meconium contamination. The fetus passes meconium because of hyperperistalsis in response to a stressor that may be very transient or may be more serious and protracted (eg, hypoxia). A very good correlation states that the more meconium present, the more severe and immediate the stressor. Additional assessments, such as amnioscopy and amniography, must be made to determine whether the fetus is experiencing ongoing episodes of hypoxia or other stressors. Green color can also indicate erythroblastosis but is not necessarily indicative of it.
5. **Yellow-brown, opaque** fluid may indicate intrauterine death and fetal maceration (although not necessarily from erythroblastosis), oxidized hemoglobin, or maternal trauma.

**Reference Values**

Normal Sound: colorless or pale straw-yellow color

**Procedure**

1. Observe color changes and staining through amnioscopy before the amniotic membranes have ruptured.
2. Place an amnioscope into the vagina and against the fetal presenting part. Visualize the amniotic fluid through the amniotic membranes. Problems with amnioscopy include inadvertent rupture of membranes, insufficient dilatation of the cervix and consequent difficulty inserting the amnioscope, intrauterine infection, and occasional difficulty in interpreting amniotic fluid color.
3. Be aware that the test may also be difficult to perform if the patient is in active labor.

<table>
<thead>
<tr>
<th>Clinical Alert</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meconium staining may also be observed when an amniocentesis is done. After the membranes have ruptured, meconium staining may be observed in the vaginal discharge. Once meconium staining is identified, more assessments (eg, FHR patterns) must be made before delivery is contemplated to determine whether the fetus is experiencing ongoing episodes of hypoxia.</td>
</tr>
<tr>
<td>The presence of meconium in the amniotic fluid is normal in breech presentations.</td>
</tr>
</tbody>
</table>

**Interventions**

**Pretest Patient Preparation**

1. Explain the purpose of the test and the procedure if amnioscopy is done.
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret the test results, counsel, and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Amniotic Fluid Bilirubin Optical Density (OD)**

Bilirubin is a pigment acquired by the amniotic fluid during its circulation through the gastrointestinal tract. Bilirubin may be found in amniotic fluid as early as the 12th week of gestation. It reaches its highest concentration between 16 and 30 weeks. As the pregnancy continues, the amount of bilirubin progressively decreases until it finally disappears near term. Bilirubin levels increase in the presence of erythroblastic fetuses and fetuses with anencephaly or intestinal obstruction.

This measurement is used to monitor the fetal state in an Rho-negative pregnant women who has a rising anti-Rho antibody titer. The rising titer is synonymous with Rh erythroblastosis fetalis or hemolytic disease of the newborn (HDN). This determination usually is not made before 20 to 24 weeks’ gestation because no therapy is available to the fetus before that time. Close to term, the amniotic fluid bilirubin pigment concentration normally decreases in the absence of Rh sensitization.
Determination of optical density (OD) is only one of several laboratory methods used to measure bilirubin. The degree of hemolytic disease falls into three zones, using OD measurement and a wavelength (absorbance) of 450 nm (the Liley or Diazo method).

1. If OD = 0.28 to 0.46 (zone 1, low zone 2+ OD) at 28 to 31 weeks, the fetus will not be affected or will have very mild hemolytic disease.
2. If OD = 0.47 to 0.90 (zone 2, middle zone, 3+ OD), there is a moderate effect on the fetus. The fetal age and the trend in OD indicate the need for intrauterine transfusion and early delivery.
3. If OD = 0.91 to 1.0 (zone 3, high zone, 4+ OD), the fetus is severely affected, and fetal death is a possibility. In this case, a decision concerning delivery or intrauterine transfusion, depending on the age of the fetus, should be made. After 32 to 33 weeks of gestation, early delivery and extratherine treatment are preferred.
4. An OD < 0.04 indicates fetal maturity and well-being.

Reference Values

- Normal OD < 0.02 mg/dL or < 0.34 µmol/L at 450 nm absorbence wavelength by the Liley method, or 0.025 mg/dL (0.43 µmol/L) by the Diazo method, indicates maturity

Procedure

1. Collect 5 to 10 mL of amniotic fluid in a light-proof container.
2. Send the fluid to the laboratory immediately.
3. Be aware that the specimen may be refrigerated for up to 24 hours. It can be frozen if a longer time will elapse before analysis.
4. Avoid blood in the specimen. If initial aspiration produces a bloody fluid, the needle should be repositioned to obtain a specimen free of red cells. If a blood-free specimen cannot be obtained, the specimen must be examined at once, before hemolysis occurs.
5. Indicate weeks of gestation on laboratory request form.

Clinical Implications

- Increased OD is found in:
  1. Erythroblastosis fetalis
  2. Other fetal hemolytic diseases
  3. Maternal infectious hepatitis
  4. Maternal sickle cell crisis

Interfering Factors

- Blood, hemoglobin, or meconium in the specimen can produce inaccurate results.
- Maternal use of steroids interferes with the test.
- Exposure of the amniotic fluid to light compromises the test.
- Fetal acidosis interferes with the test.

Interventions

Pretest Patient Preparation

1. Explain the test purpose and the meaning of test results.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcome; monitor and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Amniotic Fluid and Desaturated Phosphatidylcholine (DSPC)

Desaturated phosphatidylcholine (DSPC) is the major component (50%) of fetal pulmonary surfactant. The concentration in amniotic fluid can be measured by separating DSPC from unsaturated lecithin. Phosphatidylcholine is the second most surface-active component of surfactant.

This test is a direct measure of primary phospholipid in surfactant and is used in the assessment of fetal lung maturity.

Reference Values

- Normal Presence of DSPC is evidence that fetus is at least 36–37 weeks in development

Procedure Obtain and examine amniotic fluid for the primary phospholipid (DSPC).

Clinical Implications
1. Normal levels are consistent with fetal lung maturity and indicate a negligible risk for RDS.
2. Low levels are associated with immaturity and a high risk for RDS.

Interfering Factors
Results may be altered by changes in amniotic fluid volume (oligohydramnios or polyhydramnios).

Interventions

Pretest Patient Preparation

1. Explain test purpose and the amniotic fluid sampling procedure.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes; monitor and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

ADDITIONAL MEASUREMENTS

Group B Streptococcal (GBS) Screening

GBS is a frequent cause of newborn pneumonia. Sepsis and meningitis are also common problems of GBS disease. Preterm infants are more susceptible to GBS disease, but it occurs most often in full-term infants.

The Centers for Disease Control and Prevention guidelines recommend that all pregnant women should be screened for anogenital group B streptococcal colonization at 35–37 weeks’ gestation.

Reference Values
Normal
A negative vaginal and anorectal culture indicates GBS has not colonized in the cultured sites. A positive culture indicates a GBS carrier and results should be recorded on the prenatal record so that it is available to the health care providers at the time and place of delivery.

Procedure

1. Use a single standard culture swab to swab the distal vagina and anorectum, or two separate swabs can be used. See Chapter 7 for more information. Place swabs in transport medium if laboratory is off site.
2. Report results on prenatal record and ensure that a copy is available at the hospital where delivery of the infant is anticipated.

Clinical Implications

1. Intrapartum antibiotic prophylaxis should be considered with positive culture results by weighing the risks and benefits of treatment with each GBS carrier who is pregnant.
2. Intravenous penicillin G is preferred treatment, but ampicillin is an alternative, and clindamycin and erythromycin can be used for those women with penicillin allergy.
3. Prophylaxis should be continued throughout active labor until delivery.
4. Women who previously have given birth to an infant with GBS disease do not need to be screened prenatailly but should receive intrapartum treatment.

Interventions

Pretest Patient Preparation

1. Explain the screening test to the patient, including the risks of GBS disease to newborn.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert

1. Oral antibiotics should not be used to treat pregnant women with a positive anogenital GBS culture because they are not effective in eliminating the carrier status or preventing neonatal disease. Treatment should take place intrapartum.
2. Also, if symptomatic or asymptomatic GBS bacteria are detected in pregnancy, treatment should be considered at the time of diagnosis because this usually indicates a heavily colonized individual. (See Chap. 7 for more information.) Intrapartum treatment is also indicated for this individual.

Posttest Patient Aftercare

1. Interpret test outcomes; monitor and counsel appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Fetal Oxygen Saturation Monitoring (FSpO₂)

FSpO₂ monitoring is used with external fetal monitoring (EFM) as an additional means of assessment when the FHR is nonreassuring or not interpretable. The FSpO₂ monitoring system involves a single-use sterile disposable sensor that is inserted through the cervix into the uterus and rests against the fetal temple, cheek, or forehead. The sensor is a reflectance sensor in which a light source and a photodetector are placed next to each other. Back scatter of light (light
Absorption by pulsing arterial blood) is measured from a vascular bed under the sensor, and when it is reflected back to photodetector, reflected light is analyzed and displayed on the monitor and on the FHR paper tracing. FSpO$_2$ monitoring should only be used after maternal membranes have ruptured and on a singleton fetus in vertex presentation with a gestational age of 36 weeks or greater.

**Reference Values**

**Normal** With FSpO$_2$ of 30% or above between uterine contractions with a nonreassuring fetal heart rate, the fetus can be assumed to be adequately oxygenated. Normal range for FSpO$_2$ is 30%–70%. Continued FSpO$_2$ readings of below 30% for more than 10 minutes are likely to lead to progressive fetal hypoxemia, acidemia, and deterioration in fetal well-being.

**Procedure**

1. Be aware that FSpO$_2$ is indicated if nonreassuring FHR pattern.
2. Perform Leopold's maneuvers to determine fetal position and sterile vaginal exam to assess dilatation, station, and presentation.
3. Apply sensors when membranes are ruptured and cervical dilatation of 2 cm or greater has been achieved with the fetus at station of -2 or below and vertex presentation.
4. Insert a single-use sterile sensor (proficiency in fetal scalp electrode insertion or intrauterine pressure catheter insertion is necessary). Insert the sensor perpendicular to sagittal suture. Insertion should be done between uterine contractions.
5. Attach the sensor to the fetal oxygen saturation monitor. This monitor may be able to interface with a fetal heart rate monitor and record as a continuous line with uterine activity.
6. Document FSpO$_2$ on labor flow sheet as a range (eg, 40%–45%), and follow standard documentation intervals of other fetal assessments such as FHR.

**Clinical Implications**

1. Single measurements of FSpO$_2$ are not useful; need to track trends.
2. FSpO$_2$ monitoring along with the use of fetal monitoring provides the ability to detect a compromised fetus and a healthy fetus with nonreassuring FHR.
3. Provides data that a fetus with a nonreassuring FHR pattern can safely continue in labor and reduce unnecessary interventions during labor and birth, therefore improving maternal-fetal outcomes and decreasing costs.

**Interfering Factors**

1. Vernix can cause interruption in FSpO$_2$ monitoring if present in significant quantity. Meconium does not interfere.
2. Strong uterine contractions may cause temporary loss of the signal from sensor at peak of uterine contractions.
3. Fetal and maternal movement can displace sensor.

**Clinical Alert**

Contraindications include the following:

1. Documented or suspected placenta previa
2. Ominous FHR pattern requiring immediate intervention
3. Need for immediate delivery not related to FHR pattern
4. Active genital herpes, hepatitis B and E, or other infections that preclude internal monitoring
5. Seropositivity for HIV

**Interventions**

**Pretest Patient Preparation**

1. Explain the reason for monitoring and the procedure involved.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Evaluate test outcomes; counsel and monitor appropriately during labor. Treat accordingly.
2. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

**Fetal Nuchal Translucency (FNT)**

FNT is a noninvasive prenatal screening tool used to alert health care providers of potential fetal abnormalities. Ultrasonography is used to assess for a fluid collection at the nape of the fetal neck. An abnormal fluid collection may be due to genetic disorders or fetal physical abnormalities. The test is most accurate at 10–14 weeks' gestation. This screening test cannot identify fetal abnormalities; it is used as a screening tool for pregnancies that need diagnostic testing. If the FNT exceeds the normal range, then diagnostic genetic testing is recommended. There is a relationship between increased fluid in nuchal area and cardiac abnormalities. The nuchal edema is thought to be a compensation factor in the fetus. It may indicate trisomy 18, trisomy 21 (Down's syndrome), or fetal cardiac anomalies.

**Reference Values** Abnormal FNT during 10–14 weeks’ gestation that is greater than 2.5 mm of fluid in the fetal neck is considered abnormal. A nuchal translucency measurement of 3 mm or more is highly suggestive of fetal abnormalities.

**Procedure** FNT screening is performed by specially trained practitioners.
1. Explain the test purpose and the procedure.
2. Position the patient on her back as you would for obstetric ultrasound. A coupling gel is applied to the skin of the lower abdomen, and the ultrasound transducer is moved across the abdomen to visualize the fetal neck for fluid accumulation.
3. FNT is determined by ultrasound measurement of fluid in the nape of the neck between 10 and 14 weeks’ gestation.

Clinical Implications

1. FNT screening can alert clinicians to potential fetal abnormalities.
2. Analyzing maternal serum level of beta-hCG and PAPPA levels along with FNT increases accuracy of testing for pregnancy that is at risk for fetal abnormalities.

Interfering Factors

1. There is a small but significant difference in FNT of some ethnic groups (Caucasian and Asian fetuses have larger measurements than African and Caribbean fetuses).
2. Nuchal cord will decrease accuracy. Color Doppler ultrasound needs to be used in these cases to differentiate cord from fluid.
3. Improper caliper placement during ultrasound

Interventions

Pretest Patient Preparation

1. Explain the test purpose and procedure to the patient.
2. Refer to Chapter 1 guidelines for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test outcomes; monitor and counsel appropriately. Educate patient about further testing (genetic) and genetic counseling.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

BIBLIOGRAPHY

American College of Obstetricians and Gynecologists, Washington, DC, Antepartum Fetal Surveillance, ACOG Practice Bulletin, No. 9, October 1999


American College of Obstetricians and Gynecologists, Washington, DC, First-Trimester Screening for Fetal Anomalies with Nuchal Translucency. ACOG Committee Opinion, October 1999


Centers for Disease Control and Prevention: Recommendations for GBS disease prevention. Online: www.cdc.gov/mmwr/preview/mmwrhtml/00043277.htm

Children's Hospital of Philadelphia—Center for Fetal Diagnosis and Treatment Online: http://fetalsurgery.chop.edu/fetalnews.shtml


Wisconsin Association for Perinatal Care: Laboratory Testing During Pregnancy—Recommendations of Perinatal Testing Committee, Madison, WI, September 2000, pp. 6–25.
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Chart 16.2 Grading of Coronary Occlusions

OTHER ORGANS AND BODY FUNCTIONS

Magnetic Resonance Imaging (MRI); Magnetic Resonance Angiography (MRA); Magnetic Resonance Spectroscopy (MRS)

Clinical Alert

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OVERVIEW OF SPECIAL STUDIES

These special studies have been selected for discussion because of their great diagnostic value in identifying diseases and disorders of certain organs and systems. Tests after death serve to identify previously undiagnosed disease; evaluate accuracy of predeath diagnosis; provide information about sudden, suspicious, or unexplained deaths; assist in organ donation and postmortem legal investigations; and promote quality control in health care settings.

THE EYE

Visual Field Testing

This procedure is used in conjunction with basic eye tests to evaluate and rule out glaucoma. The visual field exam may detect diseases that affect the eye, optic nerve, or brain. Small blind spots in the visual field begin to appear early in glaucoma.

Reference Values

Normal Negative for blind spots

Procedure

1. Remember that the visual field test presents dimmer and dimmer targets of what the eye can see until they reach the limit.
2. Check one eye at a time.
3. Inform patient that procedure time is about 45 minutes for each eye.

Clinical Implications
1. Abnormal findings show the blind spots that appear in glaucoma.
2. Repeat testing for positive findings will show larger spots and progression of disease (see Fig. 16.1).

**FIGURE 16.1** Humphrey Field Analyzer II and visual field grid. The darker the symbol (from a single dot to a black square), the less likely the field is normal in that location. (Source: Zeiss Humphrey Systems, Dublin, CA, USA)

**Interventions**

**Pretest Patient Care**
1. Explain purpose and procedure for visual field testing.
2. Alert patient that no pain should be involved. The only discomfort is related to feeling sleepy, feeling like the patient is being hypnotized.
3. Be aware that elevated intraocular pressure, family history, age, and ethnicity are among the risk factors for developing glaucoma.
4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
1. Interpret test outcomes (see Fig. 16.2) and counsel appropriately, especially about need for further testing and possible treatment.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**NOTE**

For the previous section, information was produced by the Eye Clinic of Wisconsin. Laser Diagnostic Technologies of San Diego, California, supplied the technical information.

**Retinal Imaging**

Retinal imaging is a new technology (scanning laser ophthalmoscope [SLO]) performed by an optometrist and used to evaluate the back of the eye. The retina, or the inside layer at the back of the eye, is responsible for the majority of vision (see Fig. 16.3).
Cause of vision changes and general health can be diagnosed by viewing the retina. SLO technology uses red and green lasers to detect eye disease and monitor treatment. Green laser (532 mm) scans the sensory retina through the pigment epithelium layers of the retina. Red laser (633 mm) scans the deeper structures of the retina, from the pigment epithelium deep into the choroid. Unlike conventional imaging, Optomap retinal images are made at varying depths, providing additional diagnostic information. This provides up to a 200-degree internal view of the retina and is captured within 0.25 seconds. Even though the patient may not be aware vision is affected, signs of systemic disease such as diabetes, hypertension, and retinal disease like macular degeneration may be seen.

Reference Values
Normal Retinal scan: healthy eye with no diseases noted

Procedure
1. Be aware that no dilation is necessary. Through digitally imaging the back of the eye, SLO technology uses different colored lasers to scan a picture of the retina onto a computer screen (Optomap). This exam allows for a more thorough exam of the retina than the routine ophthalmoscope exam.
2. Recommend this exam for all patients during routine eye exams to follow the health of the eyes more accurately.
3. Tell patient that procedure time is approximately 25 minutes.

Clinical Implications Abnormal results show evidence of bleeding in eye associated with (most often) diabetes, hypertension, or macular degeneration.

Interventions
Pretest Patient Care
1. Explain the purpose of the eye test and procedure.
2. Do not administer eye drops for test.
3. Remember that if patient has contact lenses, these may be worn during the procedure.
4. Be aware that generally, there is minimal or no discomfort during the procedure. However, some individual's eyes may be sensitive to the flashing of light of the lasers.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Interpret test results. If bleeding or related problems that may require surgery are identified, the doctor may refer patient to another eye specialist.
2. Refer the patient to the appropriate medical specialist if problems related to medical diagnosis, such as bleeding in eye related to diabetes or hypertension, are identified.
3. Instruct the patient to return for follow-up a year later for repeat tests to evaluate disease progression.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Retinal Nerve Fiber Analysis
This procedure evaluates glaucoma by use of microscopic laser technology to precisely measure the thickness of the retinal nerve fiber of the eye and is recorded in computerized data for analysis. It is the nerve layer that receives and transmits images that gives us vision. The instrument is portable, and an ophthalmologist is the clinician who uses this technology.

Reference Values
Normal No abnormalities of retinal nerve fiber Normal thickness of retinal nerve layer

Procedure
1. Remember that no eye drops (dilation) are necessary.
2. Have the patient sit upright in the examining chair.
3. Place the patient's forehead and chin in cuplike holders and check one eye at a time. Twenty sectional images are obtained in less than 1 second and then analyzed to determine thickness of nerve layer.

Clinical Implications Abnormal appearance of the optic nerve is associated with changes in the eye that occur in glaucoma. Changes may be associated with vision loss.

Interventions
Pretest Patient Care
1. Explain test purpose and procedure. No pain or discomfort is associated with this test. There are no bright flashes of light.
2. Be aware that contact lenses may be left in place.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Evaluate outcomes, counsel appropriately, and explain if there is need for further testing and possible treatment of abnormal outcomes.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Fluorescein Angiography (FA)
The purpose of this test is to detect vascular disorders of the retina that may be the cause of poor vision. Fluorescein, a yellow-red contrast substance, is injected intravenously over a 10–15 second period. Under ideal conditions, retinal capillaries 5–10 µm in diameter can be visualized using FA. Images of the eye, taken by a special camera, are studied to detect the presence of retinal disorders.

Reference Values
Normal Normal retinal vessels, retina, and circulation

Procedure
1. Give a series of three drops to dilate the pupil of the eye.
   a. Complete dilatation occurs within 30 minutes of giving the last drop.
   b. When dilatation is complete, take a series of color photographs of both eyes.
2. Have the patient sit with the head immobilized in a special headrest in front of a fundus camera.
3. Inject fluorescein dye intravenously.
4. Take a series of photographs as the dye flows through the retinal blood vessels over a period of 3–4 minutes.
5. Take a final series of photographs 8–10 minutes after the injection.
6. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Alert
1. Choroidal circulation is not seen with color photographs.
2. Some patients may experience nausea for a short period of time following the injection.
3. The eye drops may sting or cause a "burning" sensation.

Clinical Implications Abnormal results reveal:
1. Diabetic retinopathy
2. Aneurysm
3. Macular degeneration
4. Diabetic neovascularization
5. Blocked blood vessels
6. Leakage of fluid from vessels

Interventions
Pretest Patient Care
1. Determine whether the patient has any known allergies to medications or contrast agent.
2. Instruct patient about the purpose, procedure, and side effects of the test.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Inform the patient that the patient may experience color changes in the skin (yellowish) and urine (bright yellow or green) for 36–48 hours after the test.
2. Advise the patient to wear dark glasses and not to drive while the pupils remain dilated (4–8 hours). During this time, patients are unable to focus on nearby objects and react abnormally to changes in light intensity.
3. Interpret test results and monitor appropriately.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Electro-oculography (EOG)
This test of retinal function is used in the study of suspected hereditary and acquired degeneration of the retina. As a measurement of retinal function, electro-oculography (EOG) serves primarily to complement electrotretinography (ERG) by determining the functional state of retinal pigment epithelium, as in retinitis pigmentosa. EOG determines the electrical potential of the eye at rest in both darkness and light. Normally, the potential difference between the front and back of the eye should increase as light intensity increases.

Reference Values
Normal >1.85 ratio (Arden ratio: maximum height of the retinal potential in light divided by the minimum height of the potential in the dark)

Procedure
1. Have the patient sit in the examining chair.
2. Place skin surface electrodes in the inner and outer canthi of the eye. The electrical potentials are recorded on a polygraph unit.
3. Make two recordings:
   a. Make recordings after 15 minutes with the patient in total darkness with measurement of eye movement through a known angle.
   b. Ask the patient to again move the eyes through the same angle, this time with the integrating sphere lighted.
4. Tell patient that total examination time is 40–45 minutes.

Clinical Alert

If FA and EOG are both ordered, the EOG must be done first because the eye is dilated for the FA test but not for the EOG test. However, when an ERG and an FA are performed on the same day, the FA should be done first to avoid corneal edema caused by the corneal electrode used in the ERG procedure. The waiting time between FA and ERG should be at least 2 hours.

Clinical Implications

1. An Arden ratio of 1.60–1.84 is probably abnormal; a ratio of 1.20–1.59 is definitely abnormal; whereas a ratio < 1.20 is flat. The outcome is usually reported as normal or abnormal.
2. The EOG ratio decreases in most retinal degeneration, eg, retinal pigmentosa; this sometimes parallels the decrease on the ERG examination.
3. In Best's disease (congenital macular degeneration), the EOG is abnormal; however, the ERG is normal.
4. In retinopathy, due to toxins such as antimalarial drugs, the EOG may show abnormalities earlier than the ERG.
5. Supernormal EOGs have been noted in albinism and aniridia (loss of all or part of iris) in which the common factor seems to be chronic excessive light exposure resulting in retinal damage.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Clinical Alert

For the most part, the patient will experience little to no discomfort.

Posttest Patient Aftercare

1. Interpret test results and monitor appropriately.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Electroretinography (ERG)

The electroretinography (ERG) is used to study hereditary and acquired disorders of the retina, including partial and total color blindness (achromatopia), night blindness, retinal degeneration, and detachment of the retina in cases in which the ophthalmoscopic view of the retina is prohibited by some opacity, eg, vitreous hemorrhage, cataracts, or corneal opacity. When these disorders exclusively involve either the rod system or the cone system to a significant degree, the ERG shows corresponding abnormalities.

In this test, an electrode is placed on the eye to obtain the electrical response to light. When the eye is stimulated with a flash of light, the electrode will record potential (electric) change that can be displayed and recorded on an oscilloscope. The ERG is indicated when surgery is considered in cases of questionable retinal viability.

Reference Values

Normal Normal A and B waves

NOTE

“A” waves are produced by photoreceptor cells and “B” waves by Müller cells.

Procedure

1. Have patient hold eyes open during the procedure.
2. Remember that the patient may be sitting up or lying down.
3. Instill topical anesthetic eye drops.
4. Place bipolar cotton wick electrodes, saturated with normal saline, on the cornea.
5. Use two states of light adaptation to detect rod and cone disorders along with different wavelengths of light to separate rod and cone function. Normally, the more intense the light, the greater the electrical response.
   a. Room (ambient) light
   b. Room darkened for 20 minutes, then a white light is flashed
   c. Bright flash (In cases of trauma, when there is vitreous hemorrhage, a much more intense flash of light must be used.)
6. Use chloral hydrate or a general anesthesia for infants and small children who are being tested for a congenital abnormality.
7. Be aware that total examining time is about 1 hour.
Clinical Implications

1. Changes in the ERG are associated with:
   a. Diminished response in ischemic vascular diseases, eg, arteriosclerosis, giant cell arteritis
   b. Siderosis (poisoning of the retina when copper is imbedded intraocularly [this is not associated with stainless steel foreign bodies])
   c. Drugs that produce retinal damage, eg, chloroquine, quinine
   d. Retinal detachment
   e. Opacities of ocular media
   f. Decreased response, eg, in vitamin A deficiency or mucopolysaccharidosis

2. Diseases of the macula do not affect the standard ERG. Macular disorder can be detected using a focal ERG.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test.
2. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Clinical Alert

1. For the most part, the patient will experience little to no discomfort. The electrode may feel like an eyelash in the eye.
2. Caution the patient not to rub his or her eyes for at least 1 hour after testing to prevent accidental corneal abrasion.

Posttest Patient Aftercare

1. Interpret test results and monitor appropriately.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Eye and Orbit Sonograms

Ultrasound can be used to describe both normal and abnormal tissues of the eyes when no alternative visualization is possible because of opacities caused by inflammation, hemorrhage, or dense cataracts. This information is valuable in the management of eyes with large corneal leukomas or conjunctival flaps and in the evaluation of the eyes for keratoprosthesis. Orbital lesions can be detected and distinguished from inflammatory and congestive causes of exophthalmus with a high degree of reliability. An extensive preoperative evaluation before vitrectomy or surgery for vitreous hemorrhages is also done. In this case, the vitreous cavity is examined to rule out retinal and choroidal detachments and to detect and localize vitreoretinal adhesions, choroidal lesions, and intraocular foreign bodies. It can also be used to detect optic nerve drusen. Persons who are to have intraocular lens implants after removal of cataracts must be measured for the length of the eye (within 0.1 mm).

Reference Values

Normal Normal image pattern indicating normal soft tissue of eye, retrobulbar orbital areas, retina, choroid, and orbital fat

Procedure

1. Anesthetize the eye area by instilling eye drops.
2. Ask the patient to fix the gaze and hold very still. (If imaging a lesion, movement is required for a retinal detachment evaluation.)
3. Place a small, very-high-frequency transducer directly on the eye, or position over a water standoff pad placed onto the eye surface.
4. Take multiple images.
5. Be aware that if a lesion in the eye is detected, as much as 30 minutes may be required to differentiate the pathologic process accurately.
6. Remember that orbital examination can be done in 8–10 minutes.
7. See Chapter 1 guidelines for intratest care.

Clinical Implications

1. Abnormal patterns are seen in:
   a. Alkali burns with corneal flattening and loss of anterior chamber
   b. Detached retina
   c. Keratoprosthesis
   d. Extraocular thickening in thyroid eye disease
   e. Pupillary membranes
   f. Cycloidal membranes
   g. Vitreous opacities
   h. Orbital mass lesions
   i. Inflammatory conditions
   j. Vascular malformations
   k. Foreign bodies
   l. Hypotony
   m. Optic nerve drusen
   n. Congenital cataract
o. Posterior vitreous detachment  
p. Retinoschisis  
q. Choroidal hemorrhage/detachment  
r. Trauma  

2. Abnormal patterns are also seen in tumors of various types based on specific ultrasonic patterns:  
   1. Solid tumors (eg, meningioma, glioma, neurofibroma)  
   2. Cystic tumors (eg, mucocele, dermoid, cavernous hemangioma)  
   3. Angiomatosus tumors (eg, diffuse hemangioma)  
   4. Lymphangioma  
   5. Infiltrative tumors (eg, metastatic lymphoma, pseudotumor)

Clinical Alert
1. When a ruptured globe is suspected or surgery has been performed, ophthalmic ultrasound can be performed over a closed eyelid.  
2. Caution must be used to avoid excessive pressure applied to the globe causing expulsion of the contents and increased risk for introduction of bacteria.

Interfering Factors If at some time the vitreous humor in a particular patient had been replaced by gas or silicone oil, no result may be obtained.

Interventions

Pretest Patient Care
1. Explain the purpose and procedure of the test.  
2. See Chapter 1 guidelines for safe, effective, informed pretest care.

Clinical Alert
1. For the most part, the patient will experience little to no discomfort.  
2. Caution the patient not to rub his or her eyes until the effects of the anesthesia have disappeared to prevent accidental corneal abrasion.  
3. Minor blurred vision may be experienced for a short time.

Posttest Patient Aftercare
1. Interpret test results and monitor appropriately.  
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

BRAIN AND NERVOUS SYSTEM

Electroencephalography (EEG) and Epilepsy/Seizure Monitoring

The EEG measures and records electrical impulses from the brain cortex. It is used to investigate causes of seizures, to diagnose epilepsy, and to evaluate brain tumors, brain abscesses, subdural hematomas, cerebral infarcts, and intracranial hemorrhages, among other conditions. It can be a tool for diagnosing narcolepsy, Parkinson’s disease, Alzheimer’s disease, and certain psychoses. It is common practice to consider the EEG pattern, along with other clinical procedures, drug levels, body temperature, and thorough neurologic examinations, to establish electrocerebral silence, otherwise known as “brain death.” The American Electroneurodiagnostic Society sets guidelines for obtaining these recordings. When an electrocerebral silence pattern is recorded in the absence of any hope for neurologic recovery, the patient may be declared brain dead despite cardiovascular and respiratory support.

Epilepsy/seizure monitoring using simultaneous video and EEG recordings (online computer) is done to verify a diagnosis of epilepsy, when seizures begin, and how they appear. The results differentiate and define seizure type, localize region of seizure onset, quantify seizure frequency, and identify candidates for medical implantation of vagus nerve stimulator or surgical treatment of seizures. Hospital admission is required.

Reference Values

Normal

1. Normal, symmetric patterns of electrical brain activity
2. Range of alpha: 8–11 Hertz (cycles per second)
3. Seizure monitoring: expected outcome of at least three typical recorded seizures that may be different from what the patient usually experiences because medications have been reduced; also, onset area and type of seizures
4. No cross-circulation of internal carotid arteries
5. Evidence of hemispheres to support language and memory

Procedure

1. Be aware that an EEG can be done at any time. Scalp hair should be recently washed.
2. Fasten electrodes containing conduction gel to the scalp with a special skin glue or paste. Seventeen to 21 electrodes are used according to an internationally accepted measurement known as the 10–20 System. This system correlates electrode placement with anatomic brain structure.
3. Place the patient in a recumbent position, instruct to keep the eyes closed, and encourage to sleep during the test (resting EEG). (Seizure activating procedure [see number 4, number 5 and number 6]).
4. Before beginning the EEG, some patients may be instructed to breathe deeply through the mouth 20 times per
minute for 3 minutes. This hyperventilation may cause dizziness or numbness in the hands or feet but is nothing to be alarmed about. This activating breathing procedure induces alkalosis, which causes vasoconstriction, which in turn may activate a seizure pattern.

5. Place a light flashing at frequencies of 1 to 30 times per second close to the face. This technique, called photic stimulation, may cause an abnormal EEG pattern not normally recorded.

6. Be aware that certain persons may be intentionally sleep deprived before the test to promote sleep during the test. Administer an oral medication to promote sleep (eg, Valium chloral hydrate). The sleep state is valuable for revealing abnormalities, especially different forms of epilepsy. Make recordings while the patient is falling asleep, during sleep, and while the patient is waking.

7. Remove electrodes, glue, and paste after the EEG. The patient may then wash the hair.

8. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

**Procedure for Seizure Monitoring**

1. Apply electrodes, take EEG, and explain video and EEG monitoring (for up to 6 days). An electrode panel is applied and must be covered when patient eats. Patient remains in bed except to use the bathroom; a helmet is worn when out of bed.

2. Perform neuropsychologic testing to evaluate memory (remember objects), language (circles, squares), and problem solving (4–6 hours of testing).

3. A cerebral angiogram to assess cross-circulation in carotids is followed by a Wada test to determine the dominant hemisphere for language and if opposite hemisphere can support memory. An intravenous line is started and a catheter is threaded through the femoral artery to the internal carotid to inject sodium amobarbital to "put the brain to sleep" for 5 minutes in each half of the brain. The Wada test is also known as the intracarotid amytal test or the Brevital (when sodium methohexital is used) test.

4. Perform a functional brain MRI. Procedure time is about 90 minutes. Patient wears earphones and is asked to respond to questions, sounds, and pictures by pressing a special button.

5. A combined PET/CT scan is often done to provide further information about brain hemispheres.

**Clinical Implications**

1. Abnormal EEG pattern readings reveal seizure activity (eg, grand mal epilepsy, petit mal epilepsy) if recorded during a seizure. If a patient suspected of having epilepsy shows a normal EEG, the test may have to be repeated using sleep deprivation or special electrodes. The EEG may also be abnormal during other types of seizure activity (eg, focal [psychomotor], infantile myoclonic, or jacksonian seizures); between seizures, 20% of patients with petit mal epilepsy and 40% with grand mal epilepsy show a normal EEG pattern, and the diagnosis of epilepsy can be made only by correlating the clinical history with the EEG abnormality, if one exists.

2. An EEG may often be normal in the presence of cerebral pathology. However, most brain abscesses and glioblastomas produce EEG abnormalities.

3. Electroencephalographic changes due to cerebrovascular accidents depend on the size and location of the infarcts or hemorrhages.

4. Following a head injury, a series of EEGs may be helpful in predicting the likelihood of posttraumatic epilepsy, especially if a previous EEG is available for comparison.

5. In cases of dementia, the EEG may be normal or abnormal.

6. In early stages of metabolic disease, the EEG is normal; in the later stages, it is abnormal.

7. The EEG is abnormal in most diseases or injuries that alter the level of consciousness. The more profound the change in consciousness, the more abnormal the EEG pattern.

8. Abnormal procedure results, eg, identification of major connections between the anterior and posterior circulation, or abnormal connection between the internal carotid arteries, or isolation of seizure onset and number and types of seizures.

**Interfering Factors**

1. Sedative drugs, mild hypoglycemia, or stimulants can alter normal EEG tracings.

2. Oily hair, hair spray, and other hair care products interfere with the placement of EEG patches and the procurement of accurate EEG tracings.

3. Artifacts can appear in technically well-performed EEGs. Eye and body movements cause changes in brain wave patterns and must be noted so that they are not interpreted as abnormal brain waves.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure to allay patient fears and concerns. Emphasize that the EEG is not painful, that it is not a test of thinking or intelligence, that no electrical impulses pass through the body, and that it is not a form of shock therapy. The transmitted impulses are magnified at least 1 million times and transcribed to permanent hard copy for further study.

2. Explain seizure monitoring procedures, purposes, and risks. Risks of angiogram and Wada test include allergy to sodium amobarbital, cross-circulation leading to respiratory arrest, and stroke related to allergy to contrast agent used in angiogram.

3. Allow food if the patient is to be sleep deprived. However, no coffee, tea, or cola is permitted within 12 hours of the test. Emphasize that food should be eaten to prevent hypoglycemia.

4. Allow, but do not encourage, smoking before the test.

5. Have patient wash and thoroughly rinse hair with clear water the evening before the EEG so that the EEG patches remain firmly in place during the test. Tell patient to not apply conditioners or oils after shampooing.

6. Be aware that if a sleep study is ordered, the adult patient should sleep as little as possible the night before (ie, stay up past midnight) so that sleep can occur during the test.

7. Call the EEG department for special instructions if a sleep-deprivation study is ordered for a child.
8. Medications are generally reduced before the Wada test. A liquid breakfast is permitted.
9. EEG and video monitoring of seizures occur for up to 6 days, with medications gradually reduced by 1/3 for 3 days.
10. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Wash the hair after the test. Application of oil to the adhesive before shampooing can ease its removal.
2. Allow the patient to rest after the test if a sedative was given during the test. Put bedside rails in the raised position for safety. Resume medications (if reduced preprocedure).
3. Be aware that skin irritation from the electrodes usually disappears within a few hours.
4. Interpret test results and monitor appropriately. If a repeat testing is necessary, provide explanations and support to the patient. Explain possible treatment of uncontrolled seizures, eg, newer antiseizure medications, surgical implantation of vagus nerve stimulator. Explain role of female hormones in epilepsy: seizures worsened by hormones, adult epilepsy involves areas of the brain sensitive to reproductive hormone, and at menopause, seizures tend to increase, worsen, or lessen.
5. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Evoked Responses/Potentials: Brain Stem Auditory Evoked Response (BAER); Visual Evoked Response (VER); Somatosensory Evoked Response (SSER)

These tests use conventional EEG recording techniques with specific electrode site placement for each procedure and include computer data processing to evaluate electrophysiologic integrity of the auditory, visual, and sensory pathways. These are brain responses “time-locked” to some event. See Chart 16.1 for wave and standard deviation (SD) measurements.

<table>
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<tr>
<th>Chart 16.1 Wave and Standard Deviation Measurements for Evoked Response Potential</th>
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<tr>
<td>Normal Potentials, Brain-Stem Auditory Evoked Response (BAER), and Visual Evoked Response (VER)</td>
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<td>Absolute latency, measured in milliseconds (msec), of the first five waveforms at a sound stimulation rate of 11 clicks/second</td>
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</tbody>
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<table>
<thead>
<tr>
<th>Wave Mean ± Standard Deviation (SD)</th>
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<tbody>
<tr>
<td>I 1.7 ± 0.15</td>
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<tr>
<td>II 2.8 ± 0.17</td>
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<tr>
<td>III 3.9 ± 0.19</td>
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<tr>
<td>IV 5.1 ± 0.24</td>
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<td>V 5.7 ± 0.25</td>
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<th>Normal Visual Evoked Response (VER)</th>
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<tr>
<td>Absolute latency, measured in milliseconds of the first major positive peak (P&lt;sub&gt;100&lt;/sub&gt;)</td>
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<thead>
<tr>
<th>Wave Mean ± SD Range</th>
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<tr>
<td>P&lt;sub&gt;100&lt;/sub&gt; 102.3 ± 5.189–114</td>
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<tr>
<th>Normal Somatosensory Evoked Response (SSER)</th>
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<tr>
<td>Absolute latency of major waveforms, measured in milliseconds at a stimulation rate of 5 impulses/second</td>
</tr>
</tbody>
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<table>
<thead>
<tr>
<th>Wave Mean ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>EP 9.7 ± 0.7</td>
</tr>
<tr>
<td>A 11.8 ± 0.7</td>
</tr>
<tr>
<td>B 13.7 ± 0.8</td>
</tr>
<tr>
<td>II 11.3 ± 0.8</td>
</tr>
<tr>
<td>III 13.9 ± 0.9</td>
</tr>
<tr>
<td>N&lt;sub&gt;2&lt;/sub&gt; 19.1 ± 0.8</td>
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<tr>
<td>P&lt;sub&gt;2&lt;/sub&gt; 22.0 ± 1.2</td>
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</table>
Brain Stem Auditory Evoked Response. This study allows evaluation of suspected peripheral hearing loss, cerebellopontine angle lesions, brain stem tumors, infarcts, multiple sclerosis, and comatose states. Special stimulating techniques permit recording of signals generated by subcortical structures in the auditory pathway. Stimulation of either ear evokes potentials that can reveal lesions in the brain stem involving the auditory pathway without affecting hearing. Evoked potentials of this type are also used to evaluate hearing in newborns, infants, children, and adults through electrical response audiometry.

Visual Evoked Response. This test of visual pathway function is valuable for diagnosing lesions involving the optic nerves and optic tracts, multiple sclerosis, and other disorders. Visual stimulation excites retinal pathways and initiates impulses that are conducted through the central visual path to the primary visual cortex. Fibers from this area project to the secondary visual cortical areas on the brain's occipital convexity. Through this path, a visual stimulus to the eyes causes an electrical response in the occipital regions, which can be recorded with electrodes placed along the vertex and the occipital lobes. It is also used to assess development of blue-yellow pathway in infants.

Somatosensory Evoked Response. This test assesses spinal cord lesions, stroke, and numbness and weakness of the extremities. It studies impulse conduction through the somatosensory pathway. Electrical stimuli are applied to the median nerve in the wrist or peroneal nerve near the knee at a level near that which produces thumb or foot twitches. The potentials it takes for the current to travel along the nerve to the cortex of the brain is then measured. Somatosensory evoked responses can also be used to monitor sensory pathway conduction during surgery for scoliosis or spinal cord decompensation and/or ischemia. Loss of the sensory potential can signal impending cord damage.

Procedures

1. Obtain brain stem auditory evoked responses through electrodes placed on the vertex of the scalp and on each earlobe. Stimuli in the form of clicking noises or tone bursts are delivered to one ear through earphones. Because sound waves delivered to one ear can be heard by the opposite ear, a continuous masking noise is simultaneously delivered to the opposite ear.
2. Place electrodes used in visually evoked response on the scalp along the vertex and occipital lobes. Ask the patient to watch a checkerboard pattern flash for several minutes, first with one eye, then with the other, while brain waves are recorded.
3. Record somatosensory evoked responses through several pairs of electrodes. Apply electrical stimuli to the median nerve at the wrist or to the peroneal nerve near the knee. Scalp electrodes placed over the sensory cortex of the opposite hemisphere of the brain pick up the signals and measure, in milliseconds, the time it takes for the current to travel along the nerve to the cortex of the brain.
4. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications

1. Abnormal BAERs are associated with the following conditions:
   a. Acoustic neuroma
   b. Cerebrovascular accidents
   c. Multiple sclerosis
   d. Lesions affecting any part of the auditory nerve or brain stem area
2. Abnormal VERs are associated with the following conditions:
   a. Demyelinating disorders such as multiple sclerosis
   b. Lesions of the optic nerves and eye (prechiasma defects)
   c. Lesions of the optic tract and visual cortex (postchiasma defects)
   d. Abnormal visual evoked potentials may also be found in persons without a history of retrobulbar neuritis, optic atrophy, or visual field defects. However, many patients with proven damage to the postchiasma visual path and known visual field defects may have normal visual evoked potentials.
3. Abnormal SSERs are associated with the following conditions:
   a. Spinal cord lesions
   b. Cerebrovascular accidents
   c. Multiple sclerosis
   d. Cervical myelopathy accident

Interfering Factors

1. Some difficulty in interpreting brain stem evoked potentials may arise in persons with peripheral hearing defects that alter evoked potential results (ie, subthreshold stimulation of peripheral nerves and inadequate skin preparation).
2. Maximum depolarization stimulation is divided into two protocols:
   a. Brachial plexus (BP) protocol involves stimulation the median, ulnar, and superficial sensory radial nerves just proximal to the wrist.
   b. Lumbosacral (LS) protocol involves stimulating the posterior tibial and common peroneal nerves, which are the primary divisions of the lumbosacral plexus forming the sciatic nerve.

Interventions

Pretest Patient Care

1. Explain the test purpose and procedure.
2. Have patient wash and rinse hair before testing. Instruct patient not to apply any other hair preparations.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
Posttest Patient Aftercare

1. Allow patient to wash hair (assist if necessary). Remove gel from other skin areas.
2. Interpret test results and monitor appropriately for neurologic problems.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Cognitive Tests: Event-Related Potentials (ERPs)

Event-related potentials are used as objective measures of mental function in neurologic diseases that produce cognitive defects. These measurements use the method of auditory evoked response testing in which sound stimuli are transmitted through earphones. A rare tone is associated with a prominent endogenous P<sub>3</sub> component that reflects the differential cognitive processing of that tone. Although a systematic neurologic increase in P<sub>3</sub> component latency occurs as a function of increasing age in normal persons, in many instances of neurologic diseases associated with dementia, the latency of the P<sub>3</sub> component has been reported to exceed substantially the normal age-matched value.

This test is useful in evaluating persons with dementia or decreased mental functioning. It is also helpful in differentiating persons with real organic brain defects affecting cognitive function from those who are unable to interact with the examiner because of motor or language defects or those unwilling to cooperate because of problems such as depression or schizophrenia.

Reference Values

Normal No shift of P<sub>3</sub> components to longer latencies ERP: absolute latency of P<sub>3</sub> waveform P<sub>3</sub> wave mean and SD
294 ± 21 milliseconds

Procedure

1. Remember that this procedure is the same as that for auditory brain stem response.
2. Ask patients to count the occurrences of audible rare tones they hear through the earphones.
3. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications

An increased or abnormal P<sub>3</sub> latency is associated with neurologic diseases producing dementia such as the following:

1. Alzheimer's disease
2. Metabolic encephalopathy such as that associated with hypothyroidism or alcoholism with severe electrolyte disturbances
3. Brain tumor
4. Hydrocephalus

Interfering Factors

Latency of P<sub>3</sub> component normally increases with age.

Interventions

Pretest Patient Care

1. Explain the purpose and procedure of the test
2. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Interpret test results and monitor appropriately for neurologic disease.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Brain Mapping: Computed Tomography

Brain mapping uses transitional EEG data and specialized computer digitization to display the diagnostic information as a topographic map of the brain and spinal cord. The computer analyzes EEG signals for amplitude and distribution of alpha, beta, theta, and delta frequencies and displays the analysis as a color map. Specific and/or minute abnormalities are enhanced and allow comparison with normal data. This methodology is used for assessing cognitive function and for evaluating patients with migraine headaches, trauma, or episodes of vertigo or dizziness. Persons who lose periods of time and select patients with generalized seizures, dementia of organic origin, ischemic abnormalities, or certain psychiatric disorders are also candidates for this testing. With this procedure, it is possible to localize a specific area of the brain that may otherwise show up as a generalized area of deficit in the conventional EEG. Children or adults who demonstrate hyperactivity, dyslexia, dementia, or Alzheimer's disease may benefit from evaluation through brain mapping.

Reference Values

Normal Normal frequency signals and evoked responses presented as a color-coded map of electrical brain activity

Procedure

1. Ensure that the patient is rested and awake for the test so that no sleep signals appear as indicators of beta wave activity.
2. After the skin of the scalp is cleansed with an abrasive solution, place 42 electrodes at designated areas on the scalp and hold in place with adhesive or paste formulated for this purpose.
3. Place the patient in a recumbent position and instruct to keep the eyes closed and to refrain from any movement.
4. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications Abnormal brain maps can pinpoint the following conditions:

1. Areas of focal seizure discharge in persons who experience generalized seizures
2. Areas of focal irritation in persons with migraine
3. Areas of ischemia
4. Areas of dysfunction in states of dementia
5. Areas of possible brain abnormalities associated with schizophrenia or other psychotic states

Interfering Factors

1. Tranquilizers may alter results.
2. Unwashed hair or the use of hair preparations can interfere with electrode placement.
3. Eye and body movements cause changes in signals and wave patterns.

Interventions

Pretest Patient Care

1. Explain the test purpose and procedure. There are no known risks. Emphasize the fact that electrical impulses pass from the patient to the machine and not the opposite.
2. Tell patient that food and fluids can be taken before testing. However, no coffee, tea, or caffeinated drinks should be ingested for at least 8 hours before test.
3. Ensure that hair has been recently washed.
4. Ensure that tranquilizers are not taken before testing (check with physician). Other prescribed medications such as antihypertensives and insulin may be taken. If in doubt, contact the testing laboratory for guidelines.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Remove the conduction gel and encourage the patient to wash his or her hair. Provide supplies if possible.
2. Interpret test results and monitor appropriately for seizure activity and other neurologic manifestations.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Electromyography (EMG); Electroneurography; Electromyoneurogram (EMNG)

Electromyoneurography combines electromyography and electroneurography. These studies, done to detect neuromuscular abnormalities, measure nerve conduction and electrical properties of skeletal muscles. Together with evaluation of range of motion, motor power, sensory defects, and reflexes, these tests can differentiate between neuropathy and myopathy. The electroneuromyogram can define the site and cause of muscle disorders such as myasthenia gravis, muscular dystrophy, and myotonias; inflammatory muscle disorders such as polymyositis; and lesions that involve the motor neurons in the anterior horn of the spinal cord. EMG can also localize the site of peripheral nerve disorders such as radiculopathy and axonopathy. Skin and needle electrodes measure and record electrical activity. Electrical sound equivalents are amplified and recorded on tape for later studies.

Reference Values

Normal Normal EMG and EMNG

Procedure

1. Be aware that the test is done in a copper-lined room to screen out outside interference.
2. Remember that the patient may lie down or sit during the test.
3. Apply a surface disk or lead strap to the skin around the wrist or ankle to ground the patient. Choose the muscles and nerves examined according to the patient’s signs and symptoms, history, and physical condition (select nerves innervate specific muscles).
4. Encourage the patient to relax (massage certain muscles to get the patient to relax) or to contract certain muscles (eg, to point to toes) at specific times during the test.
5. Remember that testing is divided into two parts. The first test determines nerve conduction.
   a. Coat metal surface electrodes with electrode paste and firmly place over a specific nerve area. Pass electrical current (maximum, 100 mAmp for 1 msec) through the area to cause sensations, similar to shock from carpeting or static electricity or the equivalent of an AA battery, that are directly proportional to the time the current is applied. Patients with mild forms of neuromuscular disorders may feel mild discomfort, whereas those with polymyopathies may experience moderate discomfort.
   b. Read the amplitude wave on an oscilloscope and record on magnetic tape for later studies.
   c. Be aware that electrical current leaves no mark but can cause unusual sensations that are not usually considered unpleasant. How fast and how well a nerve transmits messages can be measured. Nerves in the face, arms, or legs are appropriate for testing in this way.
6. Remember that the second test determines muscle potential.
   a. Insert a monopolar electrode (a 1.25- to 7.5-cm long small-gauge needle), and incrementally advance into the muscle. Manipulate the needle without actually removing it to see if readings change, or place the needle in another muscle area.
   b. The electrode usually causes no pain unless the tip is near a terminal nerve. Ten or more needle insertions may be necessary. The needle electrode detects electricity normally present in muscle.
   c. Observe the oscilloscope for normal wave forms and listen for normal quiet sounds at rest. A “machine-gun popping” sound or a rattling sound like hail on a tin roof is normally heard when the patient contracts the muscle.
   d. If the patient complains of pain, remove the needle because the pain stimulus yields false results.
e. Total examining time is 45 to 60 minutes if testing is confined to a single extremity; testing may take up to 3 hours for more than one extremity. There is no completely “routine” EMG. The length of the test depends on the clinical problem.

7. Follow guidelines in Chapter 1 regarding safe, effective, informed intratest care.

**Clinical Implications**

1. Abnormal neuromuscular activity occurs in diseases or disturbances of striated muscle fibers or membranes in the following conditions:
   a. Muscle fiber disorders (eg, muscular dystrophy)
   b. Cell membrane hyperirritability; myotonia and myotic disorders (eg, polymyositis, hypocalcemia, thyrotoxicosis, tetanus, rabies)
   c. Myasthenia (muscle weakness states) caused by the following conditions:
      1. Myasthenia gravis
      2. Cancer due to nonpituitary adrenocorticotropic hormone (ACTH) secretion by the tumor
         a. Bronchial cancer
         b. Sarcoid
      3. Deficiencies
         a. Familial hypokalemia
         b. McArdle's phosphorylase
      4. Hyperadrenocorticism
      5. Acetylcholine blocking agents
         a. Curare
         b. Botulin
         c. Kanamycin
         d. Snake venom

2. Disorders or diseases of lower motor neurons
   a. Lesions involving motor neuron on anterior horn of spinal cord (myelopathy)
      1. Tumor
      2. Trauma
   b. Lesions involving the nerve root (radiculopathy)
      1. Guillain-Barré syndrome
      2. Entrapment of the nerve root
         a. Tumor
         b. Trauma
         c. Herniated disk
         d. Hypertrophic spurs
         e. Spinal stenosis
   c. Damage or disease to peripheral or axial nerves
      1. Entrapment of the nerve
         a. Carpal or tarsal tunnel syndrome
         b. Facial, ulnar, radial, or peroneal palsy
         c. Neuralgia paresthetica
      2. Endocrine
         a. Hypothyroidism
         b. Diabetes
      3. Toxic
         a. Heavy metals
         b. Solvents
         c. Antiamoebicides
         d. Chemotherapy
         e. Antibiotics
   d. Early peripheral nerve degeneration and regeneration.

**Interfering Factors**

1. Conduction can vary with age and normally decreases with increasing age.
2. Pain can yield false results.
3. Electrical activity from extraneous persons and objects can produce false results as a result of movement.
4. The test is ineffective in the presence of edema, hemorrhage, or thick subcutaneous fat.

**Interventions**

**Pretest Patient Care**

1. Explain the test purpose and procedure. There is a risk for hematoma if the patient is on anticoagulant therapy.
2. Be aware that sedation or analgesia may be ordered.
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
1. If the patient experiences pain, provide pain relief through appropriate interventions. Obtain an order for an analgesic if necessary.
2. Promote rest and relaxation.
3. Interpret test results and monitor appropriately for nerve and muscle disease. Provide assistance as necessary.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Clinical Alert**

1. When ordering the test, the more information that is known, the more precise the interpretation of findings will be.
2. Enzyme levels that reflect muscle activity (e.g., aspartate aminotransferase, lactate dehydrogenase, creatine phosphokinase) must be determined before actual testing because the EMG causes elevation of these enzymes for up to 10 days postprocedure.
3. Although rare, hematomas may form at needle insertion sites. Take measures, such as application of pressure to the site, to control bleeding. Notify the physician. Ascertain whether the patient is taking anticoagulants or aspirin-like drugs.

**Electronystagmogram (ENG)**

This study aids in the differential diagnosis of lesions in the brain stem and cerebellum. It can confirm the causes of unilateral hearing loss of unknown origin, vertigo, or ringing in the ears. Evaluation of the vestibular system and the muscles controlling eye movement is based on measurements of the nystagmus cycle. In health, the vestibular system maintains visual fixation during head movements by means of nystagmus, the involuntary back-and-forth eye movement caused by initiation of the vestibular-ocular reflex.

**Reference Values**

**Normal**

Vestibular-ocular reflex: normal nystagmus accompanying head turning is expected.

**Procedure**

1. Be aware that the test is usually done in a darkened room with the patient sitting or lying.
2. Remove any earwax before testing.
3. Tape five electrodes at designated positions around the eye.
4. During the study, ask the patient to look at different objects, to open and close his or her eyes, to change head position.
5. Toward the end of the test, gently blow air into each external ear canal, first on the affected side. Instill cold water, then warm water, into the ears during the test to record eye movement in response to various stimuli.
6. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

**Clinical Implications**

Prolonged nystagmus and postural instability following a head turn is abnormal and can be caused by lesions of the vestibular or ocular system, as in the following conditions:

1. Cerebellum disease
2. Brain stem lesion
3. Peripheral lesion occurring in elderly person; head trauma; middle ear disorders
4. Congenital disorders
5. Ménière’s disease

**Interfering Factors**

1. Test results are altered by the inability of the patient to cooperate, poor eyesight, blinking of the eyes, or poorly applied electrodes.
2. The patient’s anxiety or medications such as central nervous system depressants, stimulants, or antivertigo agents can cause false-positive test results.

**Interventions**

**Pretest Patient Care**

1. Explain the test purpose and procedure. No pain or known risks are associated with the test. The procedures to stimulate involuntary rapid eye movement are uncomfortable.
2. Have the patient remove makeup.
3. Have the patient abstain from all caffeinated and alcoholic beverages for at least 48 hours. Heavy meals should be avoided before testing.
4. Be aware that in most cases, medications such as tranquilizers, stimulants, or antivertigo agents should be withheld for 5 days before the test. If in doubt, consult the clinician who ordered the test.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Clinical Alert**

1. The test is contraindicated in persons who have pacemakers.
2. Water irrigation of the ear canal should not be done when there is a perforated eardrum. Instead, a finger cot may be inserted into the ear canal to protect the middle ear.

**Posttest Patient Aftercare**
1. Allow the patient to rest as necessary.
2. Be aware that if present, nausea, vertigo, and weakness may require treatment and medication. Check with the clinician who ordered the test.
3. Interpret test results and monitor appropriately for brain disease, which may manifest as loss of balance, or middle ear disease, which may cause spasmodic eye movement, vertigo, or hearing loss.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

HEART

Electrocardiography (ECG or EKG), With Brief Description of Vectorcardiogram

An ECG records the electrical impulses that stimulate the heart to contract. It also records dysfunctions that influence the conduction ability of the myocardium. The ECG is helpful in diagnosing and monitoring the origins of pathologic rhythms; myocardial ischemia; myocardial infarction; atrial and ventricular hypertrophy; atrial, atrioventricular, and ventricular conduction delays; and pericarditis. It can be helpful in diagnosing systemic diseases that affect the heart; determining cardiac drug effects (especially digitalis and antiarrhythmic agents); evaluating disturbances in electrolyte balance (especially potassium and calcium); and analyzing cardiac pacemaker or implanted defibrillator functions.

An ECG provides a continuous picture of electrical activity during a complete cycle. Heart cells are charged or polarized in the resting state, but they depolarize and contract when electrically stimulated. The intracellular body fluids are excellent conductors of electrical current and are an important component of this process. When the depolarization (stimulation) process sweeps in a wave across the cells of the myocardium, the electrical current generated is conducted to the body’s surface, where it is detected by special electrodes placed on the patient’s limbs and chest. An ECG tracing shows the voltage of the waves and the time duration of the waves and intervals. By studying the amplitude of the waves and measuring the duration of the waves and intervals, disorders of impulse formation and conduction can be diagnosed.

Reference Values
Normal
Normal positive and negative deflections in an ECG recording Normal cardiac cycle components (one normal cardiac cycle is represented by the P wave, QRS complex, and T wave; additionally, a U wave may be observed). This cycle is repeated continuously and rhythmically. The P wave indicates atrial depolarization; QRS complex indicates ventricular depolarization; T wave indicates ventricular repolarization/resting stage between beats; and U wave indicates nonspecific recovery after potentials.

![FIGURE 16.4 Commonly measured complex components. (Source: Smeltzer SC, Bare BG: Brunner and Suddarth's Textbook of Medical-Surgical Nursing, 8th ed. Philadelphia: Lippincott-Raven Publishers, 1996)](image)

Normal Waves
Capital letters refer to relatively large waves (>5 mm), and small letters refer to relatively small waves (<5 mm).

1. The P wave is normally upright; it represents atrial depolarization and indicates electrical activity associated with the original impulse that travels from the sinus node through the atrial sinus. If P waves are present; are of normal size, shape, and deflection; have normal conduction intervals to the ventricles; and demonstrate rhythmic timing variances between cardiac cycles, it can be assumed that they began in the sinoatrial node.
2. The T_a or T_p designation is used to differentiate atrial repolarization, which ordinarily is obscured by the QRS complex, from the more conventional T wave, which signifies ventricular repolarization (see number 8).
3. The Q(q) wave is the first downward/negative deflection in the QRS complex; it results from ventricular depolarization. The Q(q) wave may not always be apparent.
4. The R(r) wave is the first upright/positive deflection after the P wave (or in the QRS complex); it results from ventricular depolarization.
5. The S(s) wave is the downward/negative deflection that follows the R wave.
6. The Q and S waves are negative deflections that do not normally rise above the baseline.
7. The T wave is a deflection produced by ventricular repolarization. There is a pause after the QRS complex, and then a T wave appears. The T wave is a period of no cardiac activity before the ventricles are again stimulated. It represents the recovery phase after the ventricular contraction.
8. The U wave is a deflection (usually positive) following the T wave. It represents late ventricular repolarization of Purkinje’s fibers or the intraventricular papillary muscles. This wave may or may not be present on an ECG. If it appears, it may be abnormal, depending on its configuration.

Normal Intervals

1. The R-R interval (normally, 0.83 second at a heart rate of 72 beats/minute) is the distance between successive R waves. In normal rhythms, the interval, in seconds or fractions of seconds, between two successive R waves divided into 60 seconds provides the heart rate per minute.
2. The P-P interval (normally, 0.83 second at a heart rate of 72 beats/minute) will be the same as the R-R interval in
normal sinus rhythm. The responsiveness of the sinus node to physiologic activity (eg, exercise, rest, respiratory cycling) produces a rhythmic variance in P-P intervals.

3. The PR interval (0.16 second) measures conduction tone and includes the time it takes for atrial depolarization and normal conduction delay in the atrioventricular node to occur. It terminates with the onset of ventricular depolarization. It is the period from the start of the P wave to the beginning of the QRS complex. This interval represents the time it takes for the impulse to traverse the atria, proceed through the atrioventricular node, and reach the ventricles and initiate ventricular depolarization.

4. The QRS interval (normally, 0.12 second) represents ventricular depolarization time and tracks the electrical impulse as it travels from the atrioventricular node through the bundle branches to Purkinje's fibers and into the myocardial cells. Normal waves consist of an initial downward deflection (Q wave), a large upward deflection (R wave), and a second downward deflection (S wave). It is measured from the onset of the Q wave (or R if no Q is visible) to the termination of the S wave.

5. QT interval measures the duration of ventricular activation and recovery. It is measured from the beginning of the QRS complex to the end of the T wave. The QT interval varies with the heart rate, gender, and time of day. Normal QT interval is 350–430 milliseconds.

Normal Segments and Junctions

1. The PR segment is normally isoelectric and is the portion of the ECG tracing from the end of the P wave to the onset of the QRS complex.

2. The J junction (or J point) is the point at which the QRS complex ends and the ST segment begins.

3. The ST segment is that part of the ECG from the J point to the onset of the T wave. Elevation or depression is determined by comparing its location with the portion of the baseline between the end of the T wave and the beginning of the P wave or relating it to the PR segment. This segment represents the period between the completion of depolarization and onset of repolarization (ie, recovery) of the ventricular muscles.

4. The TP segment (~0.25 second) is the portion of the ECG record between the end of the T wave and the beginning of the next P wave. It is usually isoelectric.

Normal Voltage Measurements

1. Voltage from the top of the R wave to the bottom of the S wave is 1 mV. Voltage of the P wave is ~0.1 to 0.3 mV.

2. Negative deflection voltage is measured from the lower portion of the baseline to the nadir of the wave.

Recording the Electrical Impulses

1. Because cardiac electrical forces extend in several directions at the same time, a comprehensive view of heart activity is possible only if the flow of current in several different planes is recorded.

2. For a 12-lead ECG, 12 leads are simultaneously used to present this comprehensive picture:
   a. Limb leads (I, II, III, AVL, AVF, AVR) record events in the frontal plane of the heart.
   b. Chest leads (V1, V2, V3, V4, V5, and V6) record a horizontal view of the heart's electrical activity.

3. Occasionally, an esophageal lead, which is swallowed or placed in the esophagus, can supply additional information. This type of lead is frequently used during surgical procedures.


ECG Versus Vectorcardiogram

The vectorcardiogram, like the ECG, records the electrical forces of the heart. The major difference between these two methods is the way in which these forces are displayed. A vectorcardiogram records a three-dimensional display of the heart's electrical activity, whereas the ECG is a single-plane representation. The following are the three planes of the vectorcardiogram:

1. Frontal plane (combines the Y and X axes)
2. Sagittal plane (combines the Y and Z axes)
3. Horizontal plane (combines the X and Z axes)

Procedure

The following steps apply to both the ECG and the vectorcardiogram:

1. Have the patient assume a supine position; however, recordings can be taken during exercise.
2. Prepare the skin sites and, if necessary, shave, and place electrodes on the four extremities and on specific chest sites. Ensure that the right leg is the ground.
3. Remember that all 12 leads can be recorded simultaneously by newer ECG machines.
4. Remember that a rhythm strip is a 2-minute recording from a single lead, usually lead II. It is frequently used to
evaluate dysrhythmias.

5. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

<table>
<thead>
<tr>
<th><strong>Comparison of the ECG and Vectorcardiogram</strong></th>
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<tbody>
<tr>
<td><strong>ECG</strong></td>
<td><strong>Vectorcardiogram</strong></td>
</tr>
<tr>
<td>Records electrical forces as positive or negative deflections on a scale</td>
<td>Depicts electrical forces as vector * loops, which show the direction of electrical flow</td>
</tr>
<tr>
<td>Records activity in the frontal and horizontal planes</td>
<td>Records activity in the frontal, horizontal, and sagittal planes</td>
</tr>
<tr>
<td><em>The term vector indicates the directional flow of electrical activity.</em></td>
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</tbody>
</table>

**Clinical Implications**

1. **ECG**
   a. The ECG does not depict the actual mechanical state of the heart or functional status of the valves.
   b. An ECG may be normal in the presence of heart disease unless the pathologic process disturbs the electrical forces. It cannot predict future cardiac events.
   c. An ECG should be interpreted and treatment ordered within the context of a comprehensive clinical picture.
   d. ECG abnormalities are categorized according to five general areas:
      1. Heart rate
      2. Heart rhythm
      3. Axis or position of the heart
      4. Hypertrophy
      5. Infarction/ischemia
   e. Typical abnormalities include the following:
      1. Pathologic rhythms
      2. Conduction system disturbances
      3. Myocardial ischemia
      4. Myocardial infarction
      5. Hypertrophy of the heart
      6. Pulmonary infarction
      7. Altered potassium, calcium, and magnesium levels
      8. Pericarditis
      9. Effects of drugs
      10. Ventricular hypertrophy

2. **Vectorcardiogram**
   a. The vectorcardiogram is more sensitive than the ECG for diagnosing myocardial infarction; it is probably not any more specific.
   b. Vectorcardiography is more specific than the ECG in determining hypertrophy or ventricular dilatation.
   c. Differentiation of intraventricular conduction abnormalities is possible.


**Clinical Considerations**

1. Chest pain, if present, should be noted on the ECG strip.
2. The presence of a pacemaker and the use of a magnet in testing should be documented.
3. Marking the position on the chest wall in ink ensures a reproducible precordial lead placement.

**Interfering Factors**

1. Race: ST elevation with T-wave inversion is more common in African Americans but disappears with maximal exercise effort.
2. Food intake: high carbohydrate content is especially associated with an intracellular shift of potassium in association with intracellular glucose metabolism. Nondiagnostic ST depression and T-wave inversion are evident
Anxiety: episodic anxiety and hyperventilation are associated with prolonged PR interval, sinus tachycardia, and ST depression with or without T-wave inversion. This may be due to autonomic nervous system imbalances.

3. Deep respiration: the position of the heart in the chest shifts more vertically with deep inspiration and more horizontally with deep expiration.

4. Exercise/movement: strenuous exercise before the test can produce misleading results. Muscle twitching can also alter the tracing.

5. Position of heart within the thoracic cage: there may be an anatomic cardiac rotation in both horizontal and frontal planes.

6. Position of precordial leads: inaccurate placement of the bipolar chest leads and the transposition of right and left arm and left leg electrodes will affect test results. In normal persons, lead reversal produces the typical ECG findings of dextrocardia in frontal plane leads and can mimic a myocardial infarction pattern.


8. Age: at birth and during infancy, the right ventricle is hypertrophied because the fetal right ventricle performs more work than the left ventricle. T-wave inversion in leads V1–V3 persists into the second decade of life and into the third decade in black persons.

9. Gender: women exhibit slight ST-segment depression.

10. Chest configuration and dextrocardia: in this congenital anomaly in which the heart is transposed to the right side of the chest, the precordial leads must also be placed over the right side of the chest.

11. Severe drug overdose, especially with barbiturates, and many other medications can influence ECG configuration. Antiarrhythmics, antihistamines, and antibiotics can widen QT intervals.

12. Increases in Ca++: shortened QT; less frequently, prolonged PR interval and QRS complex

13. Decreases in Ca++: prolonged QT

a. Increased Ca++: shortened QT; less frequently, prolonged PR interval and QRS complex

b. Decreased Ca++: prolonged QT

c. Alterations in K+ may produce cardiac arrhythmias.

Interventions

Pretest Patient Care

1. Explain the test purpose, procedure (ECG is a graphic record of electric pulses associated with the contraction and relaxation of heart), and interfering factors. Emphasize that ECG is painless and does not deliver electrical current to the body. A resting ECG is no more than a 1-minute record of the heart’s electrical activity (the amount of voltage generated by the heart and the time required for that voltage to travel through the heart).

2. Have the patient completely relax to ensure a satisfactory tracing.

3. Be aware that ideally, the person should rest for 15 minutes before ECG recording. Have the patient avoid heavy meals and smoking for at least 30 minutes before the ECG, and longer if possible.

4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare

1. Recognize the limitations of an ECG. A normal ECG does not rule out coronary artery disease or areas of cardiac ischemia. Conversely, an abnormal ECG in and of itself does not always signify heart disease.

2. Interpret test results and counsel and monitor the patient appropriately. A resting ECG is usually normal in those patients who experience only angina. It can provide evidence of prior heart damage. The ECG is one diagnostic tool within a repertoire of diagnostic modalities and should be viewed as such. The presence or absence of heart disease should not be presumed solely on the basis of the ECG.

3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

1. When an ECG shows changes that indicate ischemia, injury, or infarction, these changes must be reported and acted on immediately. The goal of diagnosis and treatment is to increase myocardial blood supply and reduce oxygen demand.

a. When ECG changes represent stages of ischemia, injury, or necrosis and symptoms of possible myocardial infarction appear, the primary concern is balancing myocardial oxygen supply and demand as follows:

1. Nitroglycerin dilates blood vessels.

2. Narcotics relieve pain and anxiety.

3. Calcium channel blockers relieve coronary spasm.

4. Oxygen increases O2 supply available to the myocardium.

5. Beta-blocking drugs slow rapid heart rates.

6. Antiarrhythmic agents correct abnormal rhythms.

7. Frequent reassurances alleviate patient anxiety.

b. Monitoring for cardiac rhythm disturbances is an essential component of care. Potentially lethal dysrhythmias, especially ventricular tachyarrhythmias, require immediate intervention and may signal the need for possible cardiopulmonary resuscitation.

2. Serious diagnostic errors can be made if the ECG is not interpreted in the broader context of the patient's history, signs, and symptoms.

3. The electrical axis is not synonymous with the anatomic position of the heart.

Signal-Averaged Electrocardiogram (SAE)

The signal-averaged ECG (SAE) is a noninvasive tool for identifying patients at risk for malignant ventricular dysrhythmias, particularly after myocardial infarction. During the later phase of the QRS complex and ST segment, the myocardium produces high-frequency, low-amplitude signals termed late potentials. These late potentials correlate with
delayed activation of certain areas within the myocardium, a condition that predisposes to reentrant forms of ventricular tachycardia.

**Indications**
SAEs are performed to evaluate the etiology of ventricular dysrhythmias or as a precursor to electrophysiologic studies. Disorders that may produce regions of delayed myocardial conduction include myocardial infarction, nonischemic dilated cardiomyopathy, left ventricular aneurysm, and some forms of healed ventricular incisions (e.g., scar from tetralogy of Fallot surgical intervention).

**Reference Values**
*Normal*
Normal QRS complexes and ST segments

**Procedure**
1. Remember that the SAE, which is a modification of the conventional ECG, uses computerized techniques to provide signal averaging, amplification, and filtering of electrical potentials.
2. Place electrodes on the abdomen and anterior and posterior thorax. The signals received are converted to a digital signal. A typical QRS complex is used as a template against which subsequent cardiac cycles are compared. Typically, several hundred beats are averaged to analyze for late potentials.
3. Be aware that data collection usually takes about 20 minutes. Optimal recordings require that the patient be in a comfortable position and remain quiet, the proper application of electrodes, and elimination of interference from other electrical equipment.
4. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

**Clinical Implications**
1. SAE provides predictive values for potential ventricular tachycardias in patients who have a history of myocardial infarction or coronary artery disease.
2. Late potentials are stronger predictors of sudden death or sustained ventricular tachycardias than are ventricular dysrhythmias from a Holter monitor recording.
3. Evidence shows that late potentials associated with ventricular tachycardias are abolished following successful surgical intervention.
4. Patients who experience late potentials have a 17% incidence of sustained ventricular tachycardia or sudden death, compared with a 1% incidence in patients without late potentials. The incidence is even greater in the presence of decreased ejection fractions.
5. SAE may explain the cause of syncope subsequently identified as ventricular tachycardia during electrophysiologic study.

**Interfering Factors**
1. Increased time is required for recording beats in the presence of slow heart rates or frequent ventricular ectopics. Patient movement, talking, and restlessness also delay data procurement.
2. Bundle-branch block can interfere with impulse averaging.
3. SAE does not provide information about antiarrhythmic drug effectiveness.
4. Late potentials do not occur in every patient with ventricular tachycardia.
5. Ventricular pacing prolongs ventricular activation time and obscures late potentials. Conversely, atrial pacing, even at rapid rates, does not alter ventricular late potentials.

**Interventions**

**Pretest Patient Care**
1. Explain the test purpose and procedure.
2. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

**Posttest Patient Aftercare**
1. Interpret test outcome and monitor appropriately.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

**Cardiac Event Monitoring; ECG Continuous Monitoring; Holter Monitoring; 30-Day Event Monitoring; Implantable Monitor**

Cardiac event monitoring refers to continuous ECG recording of cardiac rhythms, unusual cardiac events, and patient activity. The patient wears a special monitor (Holter) using a loop magnetic tape recording for 24–48 hours or a memory loop battery-operated ECG recorder. These tracings are used to record onset and termination of rhythm disturbances and to diagnose the cause of dizziness, palpitations, fainting (syncope), lightheadedness, and unexplained fatigue. These procedures are also used to check pacemaker function and automatic implantable defibrillator function status and to trace drug and treatment effectiveness.

**Reference Values**
*Normal*
Normal tracings of cardiac ECG sinus rhythms and heart rate No hypoxic or ischemic ECG changes

**Procedure**
1. Holter, 24–48 hour ECG monitor
   a. Prepare the site and apply the leads. Areas may need to be shaved, cleansed with rubbing alcohol, and abraded with gauze.
   b. If and when the patient experiences symptoms, ask the patient to push an indicator marked to save the current ECG tracing. The tracings are transmitted by telephone for analysis.
2. 30-Day cardiac event ECG monitoring
   a. Prepare the site appropriately (e.g., shaving, cleansing).
   b. Apply two-channel electrodes, place leads, and connect to the monitor.
   c. Ask the patient to press a record marker when any symptoms (events) occur and also to keep a diary of symptoms.
3. Implantable monitor
   a. Surgically inserted just beneath the skin in the upper chest area
   b. Following a symptom (e.g., dizziness, fainting spell), a pager-sized device is placed over the implanted monitor to capture and save the data.
   c. Data are then analyzed by a physician or nurse.

Interfering Factors
1. Incomplete diary or event marker not pushed during symptoms
2. Mechanical ineffectiveness
3. Smoking, certain drugs

Clinical Implications
1. Abnormal tracings and record may indicate unsuspected disturbances, e.g., arrhythmias, friction, scratching, tachycardia (atrial and ventricular)
2. Bradycardia and bradycardia-tachycardia syndrome
3. Premature atrial and bradycardia-tachycardia syndrome
4. Heart blocks
5. Junctional rhythm
6. Flutter or fibrillation
7. Premature atrial or ventricular contractions
8. Hypoxic/ischemia changes

Interventions

Pretest Patient Care
1. Explain monitoring purpose and procedure. Holter monitor is usually worn for 24–48 hours and then removed. The loop recorders are usually worn for 1–2 weeks and up to 1 month. Implantable monitors can be used for several days up to several months.
2. If the patient experiences symptoms such as dizziness, or palpitations, ask the patient to push an indicator and record time of event in a diary.
3. Encourage patient to continue normal daily events; do not get recorder wet.
4. Instruct patient to avoid magnets, metal detectors, high-voltage environments, and electric blankets.
5. Be aware that an itching sensation under electrodes is common. Instruct patients not to adjust placement sites unless they call in and receive proper procedure.
6. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Aftercare
1. Remove recorder and chart time monitor is discontinued.
2. Clean electrode sites with mild soap and water and dry thoroughly.
3. Evaluate outcomes and counsel the patient appropriately about further testing and/or possible treatment.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Stress Test/Exercise Testing (Graded Exercise Tolerance Test), Submaximal Effort

This test measures the efficiency of the heart during a dynamic exercise stress period on a motor-driven treadmill or ergometer. It is valuable for diagnosing ischemic heart disease and investigating physiologic mechanisms underlying cardiac symptoms such as angina, dysrhythmias, inordinate blood pressure elevations, and functionally incompetent heart valves. Exercise testing can also measure functional capacity for work, sports, or participation in rehabilitation programs, and it can be a predictor or potential response to medical or surgical treatment. Additionally, upper limits of physiologically responsive pacemakers can be evaluated.

Systolic blood pressure normally increases with exercise, and diastolic pressure normally remains essentially unchanged. Stress exercise testing takes place under controlled conditions that include low temperatures (20°C) and lower humidity.

Reference Values
Normal Negative when the patient does not exhibit significant symptoms, arrhythmias, or other ECG abnormalities at 85% of maximum heart rate predicted for age and gender

Procedure There are many different types of stress tests. Most include the following steps:
1. Place recording electrodes on the patient's chest (see description of ECG) and attach to a monitor. Place a blood pressure recording device appropriately.
2. Be aware that as the patient walks on a motor-driven treadmill, or pedals an ergometer if walking is not possible, computerized ECG and heart monitoring devices record performance. The patient walks at progressively greater speeds and higher levels of elevation to increase both heart rate and workload.
3. Record the initial or resting ECG, heart rate, and blood pressure. Ask the patient to report any symptoms such as chest pain or shortness of breath experienced during the test. Normal persons are symptom free at submaximal
efforts; however, at peak or maximal efforts, symptoms expected in normal persons include exhaustion, fatigue, and sometimes nausea or dizziness.

4. Have the patient undergo stress testing in stages. Each stage consists of a predetermined treadmill speed (in miles or kilometers per hour) and a treadmill grade elevation (in percentage grade or degrees).

5. Monitor the ECG, heart rate, and blood pressure continually for abnormalities and any unusual symptoms such as intolerable dyspnea, chest pain, or severe cramping (claudication) in the legs.

6. Record vital signs, together with other abnormalities and complaints, at 1- to 3- minute intervals for 6 to 8 minutes posttest as the patient rests. The test is terminated if ECG abnormalities, fatigue, weakness, abnormal blood pressure changes, or other intolerable symptoms occur during the test.

7. Remember that common criteria for terminating a test include the following:
   a. Achieving maximum possible performance
   b. Emerging signs or symptoms that indicate an existing disease process
   c. Recording a predetermined endpoint, such as 85% of age-related maximal heart rate, arbitrary workload (one that raises heart rate to 150 beats/minute), or diagnostic ECG change

8. Be aware that total examination time is about 30 minutes; however, ask the patient to plan to be in the laboratory for 1 to 1.5 hours.

9. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications Abnormal responses to exercise testing include the following:

1. Alterations in blood pressure, such as:
   a. Failure of systolic pressure to rise
   b. Progressive fall in systolic pressure
   c. Elevation of diastolic blood pressure

2. Alterations in heart rate, such as:
   a. Tachycardia above that which is predetermined
   b. Brachycardia

3. Changes in ECG, such as:
   a. Depression or elevation of ST segments caused by ischemia
   b. Dysrhythmias, ventricular tachycardia, multifocal premature ventricular contractions, atrial tachycardia, second- or third-degree atrioventricular block
   c. Pacemaker failure to perform within set rate limits

4. Ventricular or supraventricular ectopics are considered abnormal responses not necessarily ischemic in origin.

5. Ischemic ST-segment depression > 0.2 mm or elevation > 1.0 mm is the most common abnormality. Men aged 40 to 59 years who develop ST depression during exercise that is not present at rest have five times the risk for overt coronary heart disease compared with men without this ST depression.

6. Unusual symptoms such as:
   a. Anginal pain
   b. Severe breathlessness
   c. Faintness, dizziness, lightheadedness, confusion
   d. Claudication, leg pain

7. Unusual signs such as:
   a. Cyanosis, pallor, skin mottling
   b. Cold sweats, piloerection
   c. Ataxia, glassy stare
   d. Gallop heart sounds
   e. Valvular regurgitation

Interfering Factors Common causes of false-positive exercise ECG responses include the following:

1. Left ventricular hypertrophy
2. Digitalis toxicity
3. ST-segment abnormality
4. Hypertension
5. Valvular heart disease
6. Left bundle-branch block
7. Anemia
8. Hypoxia
9. Vasoregulatory asthenia
10. Lown-Ganong-Levine syndrome
11. “Panic” or anxiety attack
12. Wolff-Parkinson-White syndrome

Interventions

Pretest Patient Care

1. Explain the test purpose and procedure. No food, coffee, or cigarettes are allowed for 2 hours before testing. Water may be taken.

2. Ensure that a legal consent form is signed by the patient or patient's designee.

3. Ask the patient to wear flat walking shoes or tennis shoes (no slippers). Men should wear gym shorts or light, loose-fitting trousers. Women should wear a bra, a short-sleeved blouse that buttons in front, and slacks, shorts, or pajama pants (no one-piece undergarments, pantyhose, or slips).

4. Be aware that certain medications should be withheld or discontinued before testing. Beta-adrenergic blocking agents (eg, propranolol) should have dosage reduced or be tapered gradually. The physician should write orders regarding management of the patient's drug regimen well before the test.
5. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results and monitor appropriately for abnormal responses to exercise. Report significant events or symptoms without delay.
2. Do not discharge the patient until acceptable levels for vital signs and ECG monitoring have been met.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

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**Clinical Alert**

Stress exercise testing can be risky for patients with recent onset of chest pain associated with significantly elevated blood pressures or with frequent attacks of angina. Testing may require a 4- to 6-week delay in these situations.

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**Cardiac Catheterization and Angiography (Angiocardiology, Coronary Arteriography)**

This procedure is performed to evaluate the coronary vessels and function of the left ventricle. This method is chosen to study and diagnose defects of the chambers of the heart, the heart valves, and certain blood vessels by means of inserting arterial venous catheters, which can carry contrast material into the right and left sides of the heart. As these catheters are introduced and advanced toward the heart, fluoroscopy and high-speed x-ray pictures projected onto monitors show actual heart function and motion. Injected contrast medium provides a visual definition of cardiac structures. Coronary artery patency and circulation is filmed as well. The patient’s heart rate, rhythm, and pressures are monitored continuously.

Coronary arteriograms are useful for evaluating abnormal stress tests, diagnosing heart disease, assessing the complications of a myocardial infarction, diagnosing congenital abnormalities, identifying cardiac structure and function, and measuring hemodynamic pressures within heart chambers and great vessels. They are used to measure cardiac output using contrast dilution, thermodilution, and Fick’s method and to obtain cardiac blood samples for measuring oxygen content and oxygen saturation.

Cardiac catheterization combined with angiography is indicated for patients who exhibit angina, chest pain, syncope, valve problems, ischemic heart disease, cholesteremia, symptoms with history of familial heart disease, abnormal resting or exercise ECGs, and recurring cardiac symptoms after revascularization. Other indications include young patients with a history of coronary insufficiency or ventricular aneurysm and patients who experience coronary neurosis and need assurance that their cardiac status is normal. This test can be performed during the acute stage of myocardial infarction, and if necessary, surgical intervention can be accomplished without significant delay. Although cardiac catheterization poses some risk, it is highly accurate diagnostic resource.

**Reference Values**

**Normal Cardiac Catheterization** Normal heart values, chamber size, and patent coronary arteries Normal wall and valve motion Normal cardiac output (CO): 4–8 L/minute Normal percentage of oxygen content (15–22 vol. %) and oxygen saturation (95%–100% of capacity, or 0.95–1.00)

**Normal Cardiac Volumes** End-diastolic volume (EDV): 50–90 mL/(body surface area)m² End-systolic volume (ESV): 25 mL/m² Stroke volume (SV): 45 ± 12 mL/m² Ejection fraction (EF): 0.67 ± 0.07

**Normal Hemodynamic Pressures (mm Hg)**

<table>
<thead>
<tr>
<th>Right atrium</th>
<th>Average Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>A wave</td>
<td>6 1–10</td>
</tr>
<tr>
<td>U wave</td>
<td>5</td>
</tr>
<tr>
<td>Mean</td>
<td>3 0–8</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Right ventricle</th>
<th>Average Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak systolic</td>
<td>25 15–30</td>
</tr>
<tr>
<td>End diastolic</td>
<td>4 1–7</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PAP</th>
<th>Average Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak systolic</td>
<td>25 15–30</td>
</tr>
<tr>
<td>End diastolic</td>
<td>9 3–12</td>
</tr>
<tr>
<td>Mean</td>
<td>15 9–19</td>
</tr>
<tr>
<td>PCWP</td>
<td>9 4–12</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Left atrium</th>
<th>Average Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>A wave</td>
<td>10 3–15</td>
</tr>
<tr>
<td>U wave</td>
<td>12 6–21</td>
</tr>
<tr>
<td>Mean</td>
<td>8 2–12</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Left ventricle</th>
<th>Average Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak systolic</td>
<td>130 100–140</td>
</tr>
<tr>
<td>End diastolic</td>
<td>8 3–12</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Complete aortic</th>
<th>Average Range</th>
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</thead>
<tbody>
<tr>
<td>Peak systolic</td>
<td>130 100–140</td>
</tr>
<tr>
<td>End diastolic</td>
<td>70 60–90</td>
</tr>
<tr>
<td>Mean</td>
<td>85 70–105</td>
</tr>
</tbody>
</table>
PAP, pulmonary artery pressure; PCWP, pulmonary capillary wedge pressure.

Procedure

1. Remember that the test is normally done in a special, darkened procedure room.
2. To decrease anxiety, explain the procedure and provide information about sensations the patient may experience.
   a. For right-heart catheterization, use the medial cubital, brachial, or femoral vein. Thread the catheter through the vena cava to the right atrium, through the tricuspid valve and right ventricle, to the pulmonary artery. Take pressure measurements and O₂ saturations from these areas as you manipulate the catheter.
   b. For left-heart catheterization procedure, heparinize the patient. Thread the catheter through the femoral or brachial artery and on through the aortic valve to the left ventricle. Again, take pressure readings. Introduction of contrast material, if done, provides data about left ventricular contractility, contour size, and presence of mitral regurgitation.
   c. Observe sterile surgical conditions. Prepare the skin with an antiseptic solution scrub. Inject a local anesthetic into the catheter insertion site area (eg, groin [femoral artery], antecubital [brachial artery]). Small incisions may be made to facilitate insertion. Once inserted, gently advance the catheters to the heart and great vessels.
3. If left-to-right shunt is suspected, obtain blood samples from the superior and inferior vena cava also.
4. Have the patient lie on a special x-ray table, and monitor the ECG continuously. Use intravenous sedation if necessary. During the procedure, the patient is placed in several different positions. The patient may be asked to exercise to evaluate heart changes associated with activity. Atrial pacing can also be done as part of the procedure in persons who cannot walk (eg, paraplegics) or use a treadmill. In these instances, there is a sequence of events that stress the heart followed by a rest period; then measurements are taken. The heart is paced again, followed by another rest period.
5. Be aware that sometimes the patient can watch the procedure on a television monitor if it happens to be positioned properly.
6. After x-ray films have been taken from all angles, remove the catheters, and apply manual pressure to the site for 20 to 30 minutes. Apply a sterile pressure bandage for several additional hours, if necessary. Some facilities no longer use pressure bandages. There are several devices on the market to close the access site (vascular closure devices) following the procedure. These devices can be separated into two categories: self-adsorbing sutures and hemostasis-promoting pads or patches. Less pressure and less time may be required for venous sites. Give protamine sulfate to reverse the effects of heparinization.
7. Reassure the patient frequently.
8. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Alert

Left atrial function and measurements are usually calculated from other measurements. If direct measurements are necessary, a transseptal approach must be done by advancing the catheter through the saphenous leg vein into the right atrium and then passing a needle through the catheter to puncture the atrial septum so that direct pressure readings may be obtained. The patient may be asked to exercise during the procedure to evaluate consistent changes; atrial pacing may be done during the procedure to incrementally stress and rest the heart for those patients unable to move normally (eg, paraplegic patients).

Clinical Implications

1. Abnormal results include the following:
   a. Altered hemodynamic pressures
   b. Injected contrast agent reveals altered ventricular structure and dynamics of occluded coronary arteries
   c. Blood gas analysis confirms cardiac, circulatory, or pulmonary problems
2. Abnormal hemodynamic pressures indicate the following conditions:
   a. Valve stenosis or insufficiency
   b. Left and/or right ventricular failure
   c. Idiopathic hypertrophic subaortic stenosis (IHS)
   d. Rheumatic fever sequelae
   e. Cardiomyopathies
3. Abnormal blood gas results indicate the following conditions:
   a. Congenital or acquired circulatory shunting
   b. Septal defects
   c. Other cardiac and pulmonary defects or pathology
4. When a contrast agent is injected into the ventricles, abnormalities of size, function, structure, ejection fractions, aneurysms, leaks, stenosis, and altered contractility can be detected.
5. When contrast is injected into coronary arteries, occluded vessels and circulatory function can be recorded.

Clinical Alert

1. Risk factors for complications following cardiac catheterization include >60 years of age, hypertension, peripheral vascular disease, and procedure done on an emergency basis or at same time as angioplasty. Risk factors for complications may be as high as 10% when more than three factors are present.
2. Complications associated with risk factors include myocardial infarction, cerebrovascular accidents, or death within 24 hours of procedure; hemorrhage requiring transfusion; pseudoaneurysm; fistula; or femoral thromboses.

Interventions

Pretest Patient Care
1. Explain the test purpose (determine whether arteries are obstructed and show evidence of lesions, grade the occlusions, and assess left ventricular function), procedure, benefits, and risks. A consent form must be signed before the examination. Always check for allergies, especially to iodine and contrast media. Extensive teaching may be necessary.
2. Have the patient fast for 6 to 8 hours before the procedure. Give routine, scheduled medications, eg, cardiac drugs or insulin, before the procedure unless directed otherwise. Discontinue anticoagulants at least 1 to 2 days before the procedure.
3. Give analgesics, sedatives, or tranquilizers before the procedure.
4. Ask the patient to void before the procedure.
5. The patient may wear dentures; have the patient remove jewelry and other accessories.
6. Instruct the patient regarding the need to perform deep breathing and coughing during the test, and inform them that they may feel certain sensations.
   a. Catheter insertion via antecubital or groin sites may produce significant pressure sensations when the sheath, through which the catheter is inserted and advanced, is introduced.
   b. A slight shock or “funny bone” sensation may be felt if the nerve adjacent to the artery is touched. A tiny “bump” in the neck may be felt as the catheter is inserted into the heart. Normally, pain is not felt.
   c. When the contrast agent is injected into the catheter, a pumping sensation with feelings of palpitations and hot flashes may last 30 to 60 seconds. Skin vessels vasodilate, and blood rises to the skin surface for a short time.
   d. Patients may experience nausea, vomiting, headache, and cough.
6. Angina may occur with exercise or with the contrast agent injection. Nitroglycerin or narcotics may be given.
7. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Be aware that bed rest is usually maintained for 6 hours after the test, based on the nature of the procedure, physician's protocols, and patient status. The patient is usually not permitted to raise his or her head more than 30 degrees during this time because greater angles put strain on the insertion site. Conversely, movement of the uninvolved extremities should be promoted.
2. Check vital signs frequently according to institution protocols. At the same time, check catheter insertion site for hematomas, swelling, bleeding, or bruits. Normal or other mechanical pressure to the catheter insertion site may be necessary if bleeding or hematoma develops. A bruised appearance around the site is normal. Swelling or lumps should be promptly reported to physician. Neurovascular checks should be done along with assessment of vital signs in bilateral extremities and results compared. Assess color, motion, sensation, capillary refill times, temperature, and pulse quality. Report significant changes immediately.
3. Administer prophylactic antibiotics as necessary.
4. Encourage fluid intake. Unless contraindicated, an intravenous infusion site may be maintained while the patient is on bed rest in the event that rapid intravenous access is needed.
5. Keep the affected extremity extended, not elevated or flexed. Immobilize the legs with sandbags if necessary. Apply ice packs and/or sandbags to the catheter site, if ordered; this pressure can be very painful. Prescribed analgesics can be administered for pain of hematomas or discomfort.
6. Sutures, if used, are removed per physician's instructions.
7. Interpret test results and monitor appropriately for cardiac, circulatory, neurovascular, and pulmonary problems.
8. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Clinical Alert**

1. This procedure is contraindicated in patients with gross cardiomegaly.
2. Complications include the following:
   a. Dysrhythmias
   b. Allergic reactions to contrast agent (evidenced by urticaria, pruritus, conjunctivitis, or anaphylaxis)
   c. Thrombophlebitis
   d. Insertion site infection
   e. Pneumothorax
   f. Hemopericardium
   g. Embolism
   h. Liver lacerations, especially in infants and children
   i. Excessive bleeding at the catheter site
3. Notify attending physician immediately if increased bleeding, hematoma, dramatic fall or elevation in blood pressure, or decreased peripheral circulation and abnormal or changed neurovascular findings are noted. Rapid treatment may prevent more severe complications.
4. The following equipment should always be available to treat complications of angiography:
   a. Resuscitation equipment
   b. DC defibrillator
   c. External pacemaker
   d. EEG monitor
   e. Emergency drugs
OTHER INVASIVE CARDIAC PROCEDURES

These procedures are performed by a cardiologist in the cardiac catheterization laboratory. Typically, the procedure is performed similar to a cardiac catheterization (ie, a sheath is inserted into a femoral, brachial, or radial artery, and a guide catheter is then advanced into the coronary arteries).

Percutaneous Transluminal Coronary Angioplasty (PTCA)

PTCA is a nonsurgical procedure of dilating significantly occluded arteries, thus allowing more blood flow and subsequently an increase in oxygen being delivered to the heart. It is primarily performed as an alternative to the medical or surgical management of coronary heart disease. This procedure can be done at the same time as an initial catheterization, electively at some time after the catheterization, or urgently during an acute MI.

The procedure involves passing a small catheter, tipped with a balloon, through the groin or brachial artery to the narrowed artery. The balloon is then inflated several times against the narrowed area in an attempt to reduce the occlusion and enlarge the inner lumen of the artery. During the procedure, nitroglycerin is frequently administered into the coronary artery to help dilate the artery and to prevent coronary spasms. Intravenous anticoagulation with heparin is also maintained throughout the procedure to prevent thrombus formation on the catheters and at the areas of vascular damage.

Coronary Artery Stent

A coronary artery stent is a coiled metal device permanently embedded into the coronary artery. The stent can be balloon inflated or self-expanding. A balloon inflated–type stent is fluoroscopically guided to the occluded artery and subsequently embedded into the artery upon inflation of the balloon. Self-expanding stents are covered by a retaining sheath, which is removed at the site of occlusion. Patients are maintained on antiplatelet agents for 4–6 weeks after the procedure.

Directional Artherectomy

Directional artherectomy is a technique in which a portion of the blockage is mechanically shaved off and removed from inside the artery. This procedure uses a balloon-tipped catheter with a special cutting blade on one side. As the catheter is placed against the stenotic lesion, the balloon is inflated at a low pressure on the opposite side of the artery to stabilize the catheter. The blade is then passed through the plaque (works best on noncalcified lesions).

Rotational Artherectomy

Rotational artherectomy (Rotablator, rotational ablation) is used when the blockages are long and hard. This system uses a high-speed, rotating, diamond-studded burr. When the burr is spun at a high speed (140,000–200,000 rpm), the plaque is pulverized, and the debris is then released into the bloodstream as microparticles. This procedure is particularly effective on heavily calcified lesions (Chart 16.2).

<table>
<thead>
<tr>
<th>Chart 16.2 Grading of Coronary Occlusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Normal, no decrease in lumen diameter</td>
</tr>
<tr>
<td>2. 25%: decrease in the lumen diameter of up to 25%</td>
</tr>
<tr>
<td>3. 50%: decrease in the lumen diameter of 26%–50%</td>
</tr>
<tr>
<td>4. 75%: decrease in the lumen diameter of 51%–75%</td>
</tr>
<tr>
<td>5. 90%: decrease in the lumen diameter of 76%–90%</td>
</tr>
<tr>
<td>6. 99%: hair-width lumen with &gt; 90% narrowing</td>
</tr>
<tr>
<td>7. 100%: total occlusion</td>
</tr>
</tbody>
</table>

Footnote

Source: The American Heart Association Ad Hoc Committee for Grading of Coronary Artery Disease.

Electrophysiology (EP) Studies; His Bundle Procedure

Electrophysiology studies are accomplished through an invasive test for diagnosis and treatment of ventricular and supraventricular arrhythmias. This is similar to cardiac catheterization, the difference being that EP studies measure cardiac electrical conduction system activity through solid electrode catheters instead of the open-lumen catheters used to measure circulatory system pressures. Chest electrode catheters are almost always inserted into veins because of the greater risk they pose in the arterial system (spasms, occlusion). Using fluoroscopy as a visual guide, the catheters are advanced into the right atrium and right ventricle. An x-ray monitor tracks the catheter location, and a physiologic monitor shows ECG rhythms as well as intracardiac catheter
electrograms. An EP study is highly useful for diagnosing diseases of the cardiac conduction system and provides indications for optimal treatment. In addition to measuring baseline values, the electrode catheters are used to pace the heart in an attempt to induce the same arrhythmia causing the problem. When the patient is taking antiarrhythmic drugs, the EP study can determine how well the medication is working by how easily the arrhythmia can be induced. This is in contrast to the trial-and-error method, in which there is no way to know that a particular drug is ineffective until that drug has failed to resolve the problem, frequently over a significant period of time. EP is indicated to differentiate disorders of impulse formation (supraventricular versus ventricular rhythms). EP studies also provide diagnostic insight into the etiology and mechanism of conduction disorders. EP studies are often part of the workup for syncpe, sick sinus syndrome, or tachyarrhythmias. Finally, EP studies are indicated for testing the effectiveness of antiarrhythmic drugs. Each antiarrhythmic drug has certain effects that must be anticipated during the loading phase (eg, hypotension with quinidine and procainamide, abdominal cramping with quinidine, venous pain with phenytoin). A state of “happy drunkenness” may also occur. Intravenous saline is normally used to support blood pressure in the event hypotension occurs.

Reference Values

Normal Normal EP/His bundle procedure Normal conduction intervals, refractory periods, and recovery times Controlled, induced arrhythmias

Procedure

1. Darken the room.
2. To decrease anxiety, keep the patient informed of what is being done as the procedure evolves.
3. Position the patient on an x-ray table and attach the ECG leads to specific locations.
4. Maintain sterile, aseptic surgical conditions. Usually one or two sites are chosen and prepared for catheter insertion (right and/or left antecubital area, right and/or left groin). The sites chosen depend on where in the heart the catheters have to be placed and the patency and size of the patient’s veins. Inject the insertion site with local anesthetic before catheter insertion.
5. As the catheters are advanced toward the desired location, record baseline information. Sometimes cardiac pacing may be necessary; for example, measuring sinus node recovery times requires pacing atrium until the sinus is fatigued and then measuring the time the sinus takes to recover.
6. After baseline values have been determined, use pacing to induce arrhythmias. If a sustained arrhythmia is induced, make an attempt to terminate the arrhythmia through pacing. Should the patient lose consciousness, use an external cardioverter-defibrillator to terminate the arrhythmia.
7. Hold a continuous, quiet conversation to assess the patient’s level of consciousness.
8. After the procedure, remove the catheters, and apply a sterile pressure bandage to the catheter insertion site. Manual pressure on the site may be necessary if bleeding occurs.

Clinical Implications

1. Abnormal EP results will reveal the following conditions:
   a. Conduction intervals longer or shorter than normal
   b. Refractory periods longer than normal
   c. Prolonged recovery times
   d. Induced dysrhythmia in a normal subject
2. Abnormal results indicate the following conditions:
   a. Long atrial His (AH) bundle intervals indicate disease in the atioventricular (AV) node if sympathetic and vagal influences on the AV node have been eliminated.
   b. Long ventricular His (VH) bundle intervals indicate disease in the His-Purkinje system.
   c. Prolonged sinus node recovery times indicate sinus node dysfunction such as sick sinus syndrome.
   d. Prolonged sinoatrial conduction times can indicate sinus exit block.
   e. A wide or split His bundle deflection indicates a His bundle lesion.
   f. Induction of a sustained ventricular and supraventricular tachycardia confirms the diagnosis of recurrent ventricular tachycardia (Chart 16.3).

Interventions

Pretest Patient Care

1. Explain the test purpose, procedure, benefits, and risks. Describing possible physical sensations that may be felt helps to reduce patient anxiety. These sensations may include the following:
   a. The sensation of a bug crawling in the arm and neck as the catheter is advanced
   b. Palpitations or racing heart during pacing
   c. Lightheadedness or dizziness (these must be reported when felt)
2. Obtain a signed consent form before the procedure.
3. Draw blood samples for potassium levels, and other drug levels if the effectiveness of a drug is to be determined.
4. Perform a standard 12-lead ECG before testing.
5. Ensure that nothing is consumed for at least 3 hours before testing.
6. Be aware that analgesics, sedatives, or tranquilizers are usually withheld before the procedure.
7. Ask the patient to void before the procedure is initiated.
8. Allow the patient to wear dentures.
9. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
Chart 16.3 Features of Interventional or Therapeutic EP Studies

1. Endocardial catheter ablation
2. Placement of implantable cardioverter-defibrillators for management of VT or VP
3. Radiofrequency (RF) transcatheter ablation has become the treatment of choice to obliterate pathways within the AV node and accessory pathways causing palpitations.
4. Intracardiac transcatheter ablation is used to destroy accessory bypass tracks, reentrant pathways, or pathways within the AV node. A special catheter is used to produce an injury to the target site(s) using RF energy.

Footnote


Posttest Patient Aftercare

1. Have the patient remain on flat bed rest for 4 to 8 hours postprocedure and do not allow flexion or bending of the extremity used for the catheter insertion because this may lead to bleeding or vascular occlusion. A pillow may be placed under the head.
2. Check vital signs, neurovascular status of extremity used, and insertion site for swelling, bleeding, hematoma, or bruise every 15 minutes for 4 hours, 30 minutes for 2 hours, and every hour for 2 hours postprocedure, or according to institutional protocols. Neurovascular checks include assessing for pulses, color, motion, sensation, temperature, and capillary refill times.
3. Keep the affected extremity extended, not elevated or flexed, to decrease discomfort and risk for bleeding. Prescribed analgesics can be administered.
4. Encourage range-of-motion exercise of uninvolved limbs.
5. If an electrode catheter is left in place for sequential studies, ensure that it is sutured in place and covered with sterile dressings. Care for the site using sterile, aseptic technique.
6. Interpret test results and monitor ECG and other parameters appropriately. Stress the importance of compliance with prescribed therapies including drugs.
7. Follow guidelines in Chapter 1 regarding safe, effective, informed posttest care.

Clinical Alert

1. Relative contraindications to EP: although an acute myocardial infarction may limit detailed and prolonged EP procedures, brief but clinically useful procedures can be performed in this situation.
2. Complications can include the following conditions:
   a. Rapid, dramatic hemorrhage at the catheter insertion site (apply manual pressure to the site and notify the physician immediately)
   b. Thrombosis at the puncture site; thromboembolism
   c. Phlebitis
   d. Hemopericardium
   e. Atrial fibrillation (usually transient)
   f. Ventricular fibrillation or ventricular ectopy
3. Notify the attending physician of bleeding, hypotension, altered neurovascular status, decrease in distal perfusion, or life-threatening arrhythmias. Be aware of drug studies performed and monitor for effects of that drug. Have cardiopulmonary resuscitation equipment and drugs readily available for emergency use.

Transesophageal Echocardiography (TEE)

This test permits optimal ultrasonic visualization of the heart when traditional transthoracic (noninvasive) echocardiography fails or proves inconclusive. A miniaturized high-frequency ultrasound transducer is mounted on an endoscope and coupled with an ultrasound instrument to display and record ultrasound images from the heart. Endoscope controls allow remote manipulation of the transducer tip. Various images of heart anatomy can be displayed by rotating the tip of the instrument and by varying the depth of insertion into the esophagus. Indications for TEE include the following:

1. To assess function of prosthetic valves, diagnose endocarditis, evaluate valvular regurgitation and congenital abnormalities, and examine the aorta for dissecting aneurysms
2. To monitor left ventricular wall motion intraoperatively
3. To measure ejection fraction in selected patients
4. Situations in which a transthoracic echocardiogram has not been satisfactory (eg, obesity, chest wall trauma, chronic obstructive pulmonary disease)
5. When results of traditional transthoracic echocardiography do not agree or correlate with other clinical findings

Reference Values

Normal Normal position, size, and function of heart valves and heart chambers

Procedure

1. Explain test purpose, procedure, benefits, and risks.
2. Apply a topical anesthetic to the pharynx. Insert a bite block into the mouth to reduce the risk for damage to the teeth and other oral structures as well as the endoscope itself (see Chapter 12).
3. Have the patient assume a left lateral decubitus position before the lubricated endoscopic instrument is inserted to a depth of 30 to 50 cm. The patient may be asked to swallow so that the scope advances more easily.
4. Remember that manipulation of the ultrasound transducer allows a number of image planes to be visualized.
5. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Clinical Implications Abnormal TEE findings may reveal the following conditions:
Heart valve diseases
2. Pericardial effusion
3. Congenital heart disease
4. Endocarditis
5. Intracardiac tumors or thrombi
6. Left ventricular dysfunction

**Interventions**

**Pretest Patient Care**

1. Explain test purpose, procedure, benefits, and risks.
2. Ensure that the patient fasts from food and fluids at least 8 hours before the procedure to reduce the risk for aspiration. Premedications such as analgesics or sedatives may be ordered. Prescribed oral medications may be taken with small sips of water (see Appendix C for sedation and analgesia precautions).
3. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Interpret test results. Monitor vital signs and level of consciousness (if sedated). Ensure patent airway at all times.
2. Position patient on his or her side if sedated to prevent risk for aspiration.
3. Evaluate return of swallow, cough, and gag reflexes before introducing food or fluids orally.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Clinical Alert**

Swallowing reflexes may be diminished for several hours because of topical anesthetic effects. Ingesting food or fluids may result in aspiration if these reflexes are not intact. Obtain signed, witnessed, informed consent.

**OTHER ORGANS AND BODY FUNCTIONS**

Magnetic Resonance Imaging (MRI); Magnetic Resonance Angiography (MRA); Magnetic Resonance Spectroscopy (MRS)

Magnetic resonance (MR) is a diagnostic modality that employs a superconducting magnet and radiofrequency (RF) signals to cause hydrogen nuclei to emit their own signal; computers use these signals to construct detailed, sectional images of the body. Unlike computed tomography (CT), no ionizing radiation is used. Additionally, the ability of magnetic resonance to discern anatomy is most closely linked to the molecular nature of tissue. For example, MR spectroscopy provides information about the chemical composition of tissue and is commonly used to evaluate brain function. Special techniques primarily based on the magnetic reactions of hydrogen nuclei can influence the MR signal to enhance certain types of tissue (eg, fat is accentuated in T1-weighted images, cerebrospinal fluid and other pure fluids are highlighted in T2-weighted images). Computer reconstruction techniques allow images to be produced in any plane as well as in the three-dimensional views.

During the procedure, the patient lies on specially designed couch, which is moved into a gantry. “Closed” systems, typically of higher magnetic strength, are the most commonly used magnets and can range from 0.5 to 3.0 Tesla (international unit of magnetic flux density). “Open” magnets have a more spacious gantry and are often used when claustrophobia is a problem or for a pediatric scanning. Older “open” magnets employed low-field magnets (3.2 Tesla). Recently, high-field open magnets (0.7–1.0 Tesla) have become available. Generally speaking, a higher Tesla strength magnet is associated with improved image quality. For certain procedures, surface coils are placed over the body area to be imaged. During the test, loud, rhythmic knocking sounds are produced; less noise is associated with the open-design scanner. To relieve patient anxiety and the potential for claustrophobia, some laboratories provide music for relaxation. Two-way communication systems and pulse oximeters are commonly used to monitor patient responses to the procedure. Magnetic resonance applications are continually evolving and improving. In general, the most common MR applications include the following:

1. **MR of the brain** provides exquisite visualization of the soft tissue structures of the brain. Some laboratories perform neurofunctional imaging, which maps the brain’s response to the stimuli. Although bony anatomy is seen using MRI, CT is the test of choice to evaluate bone lesions and fractures.
2. **MR of the spine** provides excellent views of the spinal cord and subarachnoid space without intrathecal contrast injection.
3. **MR of the musculoskeletal system** accurately demonstrates fat, muscles, tendons, ligaments, nerves, blood vessels, and bone marrow. If the anatomic region of interest is a small area, a surface coil, which produces the RF signal, is placed directly on the skin overlying the part to be examined. Dynamic studies of the joint in motion can be performed on open scanners.
4. **MR of the heart** (cardiac MRI) allows visualization of the structures of the heart, including valves and coronary vessels. Image acquisition is synchronized to the ECG—a process known as “gating”—to help eliminate motion artifacts. Functional studies can evaluate cardiac wall motion in response to exercise.
5. **MR of the abdomen and pelvis** visualized soft tissue organs, particularly the liver, pancreas, spleen, adrenals, kidneys, blood vessels, and reproductive organs. This is the preferred method for staging uterine, cervical, and vulvar carcinoma as well as prostate cancer.
6. **MR of the breast** is a promising new technique capable of producing exquisitely detailed analysis of complex breast lesions.
7. **MR angiography** provides both anatomic and hemodynamic information in two-dimensional and three-dimensional...
representations (likened to noninvasive angiography). MR angiography is becoming more common and is used to evaluate known vascular lesions and is finding greater utility in evaluating stroke.

Magnetic resonance spectroscopy uses a conventional MR scanner to detect chemicals in all body tissues to evaluate tumors, muscle disease, or ischemic heart disease; to differentiate causes of coma; to rule out Alzheimer's disease; to monitor cancer treatment; to differentiate the diagnosis of multiple sclerosis, human immunodeficiency virus (HIV) infection, and adrenoleukodystrophy; to prepare for temporal lobe epilepsy surgery; and to assess the extent of stroke and head injury.

Intravenous MR contrast agents, all primarily containing water-soluble gadolinium complex (most commonly gadolinium-50-DTPA or DOTA) or other metals such as manganese (Mn-DPDP) and iron (Mion, USP10) are often used in evaluating the central nervous system. These agents have been approved as safe for patients, including those < 2 years of age, and are available in oral, intravenous (most common) and inhalation formulations. Gadolinium presents with very low toxicity and fewer side effects than traditional x-ray contrast agents because of its rapid renal clearance. Other agents used include gadodiamide (nonionic) and gadopentetate, which are used for body scanning. MR contrast agents have lower toxicity and fewer side effects than x-ray contrast agents. However, because these MR contrast agents are primarily excreted via the kidneys, renal failure is a contraindication for use. Other potential contraindications include pregnancy, allergies or asthma, anemia, hypotension, epilepsy, and sickle cell disease.

Clinical Alert

1. Adverse effects, although rare, of Gd-DOTA include vomiting, sensations of local warmth or coldness, headache, dizziness, urticaria, paresthesias, unusual mouth sensations, and respiratory problems.
2. MR contrast agents allow for better basic contrast and tissue signals; most abnormal tissues show regions of increased T\textsubscript{1} and T\textsubscript{2} (relaxation time, RF signals) regardless of the nature of tumors, edema, hemorrhage, inflammation, and necrosis.

Reference Values

**Normal** Soft tissue structures: normal brain, spinal cord, subarachnoid spaces, fat, muscles, tendons, ligaments, nerves, blood vessels, marrow of limbs and joints, heart, abdomen, and pelvis Blood vessels: normal size, anatomy, and hemodynamics

Procedure

1. Have the patient lie supine on a movable examination couch after a thorough medical history is obtained.
2. Be aware that sedation may be necessary if the patient is claustrophobic or restless. Earplugs with music are another option. A two-way communication system between the patient and the operator allows continual monitoring and vocal feedback and somewhat reduces the patient's sense of isolation. Many MR laboratories routinely use a pulse oximeter to monitor the patient's arterial oxygenation during the study.
3. For examining many superficial structures (eg, knee, neck, shoulder, breast), apply a surface coil over the skin. Obtain improved images of the prostate or reproductive organs by using a transrectal coil.
4. Once the patient is positioned and instructed to remain still, move the couch into the scanner.
5. In some instances, inject a noniodinated contrast into a vein for better anatomic visualization. For abdominal or pelvic scans, administer glucagon to reduce bowel peristalsis.
6. Be aware that examination time varies and averages between 30 and 90 minutes.
7. Follow guidelines in Chapter 1 for **safe, effective, informed intratest care**.

NOTE

The closed-gantry design is narrow and may upset some individuals. Reassure patients that there is sufficient air to breathe and that they will be monitored and given voice contact during the entire procedure.
Clinical Alert

Usually, no special dietary restrictions or preparations are necessary before MRI, unless conscious sedation is to be used. However, numerous safety factors must be considered.

1. Absolute contraindications to MRI include the following conditions:
   a. Implanted devices, including pacemakers, automatic cardiac defibrillators, cochlear implants, certain prosthetic devices (consult with MR laboratory for specific information), implanted drug infusion pumps, neurostimulators, bone growth stimulators, cardioverters, certain intrauterine contraceptive devices, and metal artificial heart valves. Most new surgical implants are MRI compatible; however, most labs will not scan a patient with compatible implants until approximately 2 weeks after placement. The exact brand, style, and serial number of the device are needed in order for the MRI department to verify compatibility.
   b. Internal metallic objects such as bullets or shrapnel and certain surgical clips, pins, plates, screws, metal sutures, or wire mesh
2. MRI is generally not advised for pregnant patients (increase in amniotic fluid temperature may be harmful) or individuals with epilepsy. All patients having an MRI need to remove hearing aids, dentures, jewelry, hair pins, wigs, hairpieces, and other accessories.
3. Patients unable to remain still and those who are claustrophobic may require intravenous conscious sedation before MRI.
4. Certain types of eye makeup and permanent eye liners that contain metallic fragments sometimes cause discomfort during MRI. Assess for these cosmetic enhancements.
5. A thorough patient history is mandatory before any MR study. Commonly, radiology services perform conventional x-ray imaging to confirm or rule out the presence of metallic fragments before MR imaging. This is particularly important for metal or foundry workers who may have tiny metallic fragments in their eyes.
6. Common metallic equipment (eg, scissors, oxygen tanks, electronic devices) can become lethal projectiles when exposed to the strong magnetic fields. Therefore, a thorough screening of all patients, visitors, and staff before entering the scan room is mandatory.
7. Local burns from ECG leads, other wires, and surface coils have been reported. It is imperative that the patient describe any burning sensation to the technologist during the procedure.

Advantages of Open MRI

1. May not need to sedate the claustrophobic patient
2. Suitable for the extremely obese patient
3. Enhances patient comfort—because of the low magnetic field, another person may stay with the patient (especially useful with children or confused patients)
4. Kinematic studies of joints (eg, shoulders) are possible
5. Improved accessibility to the patient allows open MRI to be used as a guide for interventional and select surgical procedures (eg, biopsies)
6. The open head coil features a unique mirror that allows the patient to see outside the magnet during the procedure.
7. Less noise

Clinical Alert

Some open MR imaging systems use only a fraction of the traditional high-field magnets (eg, 0.2–0.3 Tesla compared with 1.0–1.5 Tesla). This results in a slimmer profile and much less intimidating appearance for the magnet. Although extremely appealing in certain instances, the open-design magnet is currently not the best choice for all MR imaging, and careful consideration to magnetic field strength should be given. Certain types of studies can only be performed with a high-field magnet. Some scans performed on an open-design low-field magnet must be repeated.

Interfering Factors

1. Respiratory motion causes severe artifacts with abdominal and thoracic imaging.
2. Morbidly obese persons may not fit into the gantry opening or surface coil configurations.

Clinical Implications

1. MRI and MRS of the brain demonstrate the following conditions:
   a. White matter disease (eg, multiple sclerosis)
   b. Infectious disorders affecting the brain (eg, toxoplasmosis in acquired immunodeficiency syndrome [AIDS], vasculitis, tuberculosis)
   c. Neoplasms (primary and metastatic brain tumors, pituitary adenomas)
   d. Ischemias, cerebrovascular accident
   e. Aneurysms, hemorrhage
   f. Hydrocephalus
   g. Vascular abnormalities (aneurysm, angiomas)
   h. Congenital CNS defects (Chiari malformation, Dandy-Walker syndrome)
2. MRI and MRS of the spine demonstrate the following conditions:
   a. Disk herniation or degeneration
   b. Neoplasm (primary and metastases)
   c. Inflammatory disease
   d. Demyelinating disease
   e. Congenital abnormalities (eg, tethered cord, spinal dysraphism)
3. MRI of the heart demonstrates the following conditions:
   a. Abnormal chamber size or myocardial thickness
   b. Cardiac tumors
c. Congenital heart disorders
d. Pericarditis
e. Graft patency
f. Thrombic disorders
g. Aortic dissection or aneurysm
h. Cardiac ischemia
i. Anomalous pulmonary venous connection

4. MRI and MRS of the limbs, joints, and soft tissue demonstrate the following conditions:
   a. Neoplasms of soft tissue and bone
   b. Ligament or tendon damage
   c. Osteonecrosis, occult fracture
   d. Bone marrow disorders
   e. Muscle fatigue
   f. Changes in blood flow
      1. Atherosclerosis
      2. Aneurysm
      3. Thrombus
      4. Embolism
      5. Bypass grafts
      6. Endocarditis
      7. Shunt placement

5. MRI of the abdomen and pelvis demonstrates the following conditions:
   a. Neoplasms (especially useful in staging tumors)
   b. Retroperitoneal structures
   c. Status of renal transplants

6. MRI angiography demonstrates the following conditions:
   a. Aneurysms
   b. Stenosis or occlusions
   c. Graft patency
   d. Vascular malformations

FIGURE 16.8 Open MRI. (Courtesy: General Electric Medical Systems, Waukesha, WI, USA)

FIGURE 16.9 Closed MRI. (Courtesy: General Electric Medical Systems, Waukesha, WI, USA.)

Interventions

Pretest Patient Care

1. Explain the test purpose, procedure, benefits, and risks. Safety concerns for the patient and staff during MRI procedures are based on interaction of strong magnetic fields with body tissues and metallic objects. These potential hazards are mainly due to projectiles (metallic objects can be displaced, giving rise to potentially dangerous projectiles); torquing of metallic objects (implanted surgical clips and other metallic structures or implants can be torqued or twisted within the body when exposed to strong magnetic fields); local heating (exposure to RF pulses can cause heating of tissues or metallic objects within the patient's body; for this reason, pregnant women are not routinely scanned because an increase in the temperature of the amniotic fluid or fetus may be harmful); interference with electromechanical implants (electronic device implants are at risk for damage from both magnetic fields and the RF pulses; consequently, patients with cardiac pacemakers, implanted drug infusion pumps, cochlear implants and similar devices should not be exposed to MR procedures); and allergic reactions to MR contrast agents.

2. Assess for contraindications to testing. Obtain a relevant history regarding any implanted devices such as heart valves, surgical and aneurysm clips, plates, internal orthopedic screws and rods, and pacemakers, among other objects.

3. Ensure that the following materials are removed before the procedure: removable dental bridges and oral appliances, credit cards, keys, hair clips, shoes, belts, jewelry, clothing with metal fasteners, wigs, hairpieces, and removable prosthesis.
4. Remember that claustrophobic feelings can be avoided if the patient keeps his or her eyes closed during the test. Recommend that the patient not eat a large meal within 1 hour of testing to reduce physiologic demands and possible emesis while in the scanner.
5. Encourage the patient to relax and instruct him or her to remain as motionless as possible during testing. Reassure the patient that this is a painless procedure.
6. Ask patients having blood flow testing to abstain from alcohol, nicotine, caffeine, and prescription drugs for iron. The patient should fast for 2 hours before testing to avoid unexpected blood vessel vasoconstrictions or dilation. No smoking is permitted before the test. Promote rest in the supine position for 10 minutes before the test.
7. Be aware that fasting or drinking only clear liquids may be necessary for several hours before an abdominal pelvic MR.
8. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Clinical Alert

In the event of respiratory/cardiac arrest, the patient must be removed from the scanning room before resuscitation. Most general hospital equipment (eg, oxygen tanks, intravenous pumps, monitors) are not permitted in the MR suite.

Posttest Patient Aftercare

1. Interpret test results. Counsel and monitor appropriately for side effects of the MR contrast agent. Common side effects include coldness at the injection site, dizziness, and headache. Treatment is usually not needed unless symptoms are bothersome or prolonged. Rare side effects include convulsions, irregular or rapid heart rate, itching and watery eyes, skin rash or hives, facial swelling, thickening of tongue, fatigue or weakness, wheezing, chest tightness, and difficulty breathing. Alert the physician if any of these occur and initiate treatment as indicated.
2. Assess the contrast dye injection site for signs of inflammation, bruising, irritation, or infection.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Special Pediatric Considerations for MR Testing

Pediatric cautions related to MR testing include the following considerations:

1. Age, ability to understand and cooperate, physical condition, and reasons for testing
2. MRI body imaging: most of the adult guidelines apply. Sedatives, tranquilizers, or modified restraints may be necessary if the child is uncooperative or fearful.
3. MRI for blood flow studies in extremities: simple restraints may be used to restrict motion of arms or legs. No tranquilizers or sedatives may be used because blood flow will be affected.
4. Also see Cautions, pages 1060–1061, 1064.

SLEEP STUDIES

Excessive daytime sleepiness (hypersomnolence) is a classic symptom of inadequate nocturnal sleep, which manifests itself pathologically in various ways. Typically, much of the daytime sleepiness in today's society is a result of irregular sleep patterns and times (eg, shift workers), lack of adequate sleep, poor nutrition, and certain medications. Sleep disorders are grouped into four major categories:

1. Dyssomnias
2. Parasomnias
3. Medical-psychiatric
4. Others

The dyssomnias are sleep disorders associated with too little or too much sleep as a result of problems initiating or maintaining sleep states or exhibiting excessive sleepiness states. Examples include sleep apnea (an intrinsic sleep disorder), periodic limb movement disorder, narcolepsy, and restless-leg syndrome. Parasomnias include arousal disorders, sleep-wake transition disorders, nightmares, sleep paralysis, and other rapid eye movement (REM) disorders. Dementia, Parkinson's disease, anxiety, and mood and panic disorders are the most common forms or medical-psychiatric sleep disorders. The "others" category includes "short" and "long" sleepers, pregnancy-associated sleep disorder, and sleep choking syndrome. These disorders are diagnosed using polysomnography methodology (eg, EEG, EMG, EOG).

A “short sleeper,” also referred to as a “healthy” hyposomniac, sleeps substantially less in a 24-hour period than is expected (sleep duration of < 5 hours in a 24-hour period before age 60 years). A “longer sleeper,” also referred to as a "healthy" hypersomniac, consistently sleeps more in a 24-hour period than is expected (sleep duration of > 10 hours in a 24-hour period). People with “sleep choking syndrome” awaken suddenly with a feeling of shortness of breath and a choking sensation. The etiology of this disorder is unknown, but it is more prevalent in early to middle adulthood in persons with obsessive-compulsive anxiety disorders.

The solution to the problem relates to reversing pathologic sleep patterns to more normal status by means of various interventions.

Use of Tests

Sleep studies, or polysomnography (PSG), can be divided into two types: full PSG, or 16-channel recording, and screening PSG, or 4-channel recording. Full PSG can be used to diagnose any of the previously described sleep disorders, whereas the 4-channel limited PSG is reserved for sleep disorders involving breathing (eg, sleep apnea).
Sleep Studies

<table>
<thead>
<tr>
<th>Term</th>
<th>EEG Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sleep onset</td>
<td>Transition from wakefulness to sleepfulness; usually takes at least 10 minutes (ie, nREM stage I)</td>
</tr>
<tr>
<td>Stage I nREM</td>
<td>Occurs at sleep onset, consists of low-voltage EEG with mainly theta and alpha activity; 4%–5% of sleep</td>
</tr>
<tr>
<td>Stage II nREM</td>
<td>Follows stage I; low-voltage EEG with sleep spindles and K complexes; 45%–55% of sleep</td>
</tr>
<tr>
<td>Stage III nREM</td>
<td>Consists of 20%–50% high-amplitude delta waves, referred to as delta or slow wave sleep; 4%–6% of sleep</td>
</tr>
<tr>
<td>Stage IV nREM</td>
<td>Consists of &gt;50% of high-amplitude delta waves and is also called slow wave sleep; 12%–15% of sleep</td>
</tr>
<tr>
<td>Stage REM</td>
<td>Low-voltage, mixed frequency, nonalpha activity with rapid eye movements, called paradoxical sleep; 20%–25% of sleep</td>
</tr>
<tr>
<td>Sleep offset</td>
<td>Transition from sleepfulness to wakefulness, alpha and beta activity, also called awakening</td>
</tr>
</tbody>
</table>

*Sleep staging is done in 30-second epochs.

Classification of Tests

The full PSG includes the following tests:

1. Electroencephalogram (EEG): at least 2 channels are recorded to determine sleep onset, sleep stages, and sleep offset.
2. Electro-oculogram (EOG): documents both slow rolling and rapid eye movements seen at sleep onset and in REM sleep, respectively.
3. Electromyogram (EMG): the chin EMG is used as a criterion for REM sleep; the leg EMG is used to evaluate periodic leg movements or leg jerks.
5. Chest impedance: monitors respiratory effort by use of cardiopneumotachographs, strain gauges, or piezoelectric crystal belts.
6. Airflow monitors: thermistors or thermocouples are used to monitor oral/nasal airflow.
8. Pulse oximetry (SpO$_2$): continuous monitoring of arterial oxygen saturation by noninvasive means.
9. Snoring sensor: microphone placed just below the jaw and lateral to the trachea.
10. pH meter: pH probe placed in the lower third of the esophagus transnasally to monitor episodes of gastric reflux
11. Audio/video recordings: document restless sleep, sleep walking, sleep talking, and night terrors, among other conditions

The 4-channel limited PSG includes the following tests:

1. Electrocardiogram (ECG)
2. Chest impedance
3. Airflow monitoring
4. Pulse oximetry

**Polysomnography (PSG)** The PSG determines underlying sleep disorder pathology, provides qualitative and quantitative measurements associated with the disorder, and provides information upon which to base the proper course of treatment. PSG is indicated for persons complaining of daytime sleepiness, fatigue, inability to stay on task, falling asleep at inappropriate times, insomnia, nocturnal awakenings, waking with gasping or choking feelings, witnessed sleep-related apneas, abnormal snoring patterns, and any other unexplained symptoms associated with disruption of normal sleeping patterns that have persisted for 6 to 12 months.

**Reference Values**

**Normal**

| Electroencephalogram (EEG): normal sleep onset time, sleep stages, and sleep offset (going from sleepfulness to wakefulness [ie, awakening]) | Airflow monitors: evidence of sustained airflow throughout the night Electro-oculogram (EOG): normal slow, rolling movements at sleep onset; rapid eye movement during REM sleep Capnography end-tidal CO$_2$ (ETCO$_2$): normocapnic (35–45 mm Hg during the awake state, increasing a couple of mm Hg during sleep) | Electromyogram (EMG): absence of periodic leg movements or jerks Pulse oximetry (SpO$_2$): >90% Snoring sensor: absence of abnormal patterns of snoring Electrocardiogram (EEG): absence of rhythmic disturbances, bradycardias, or tachycardias Audio/video recordings: absence of restless sleep, sleep walking, sleep talking, and night terrors, among other conditions Chest impedance: evidence of sustained respiratory effort throughout night Respiratory disturbance index (RDI): adults <5 apneas/hypopneas per hour (after age 60, <10 apneas/hypopneas per hour) Oxygen desaturation index (ODI): adults <5 times per hour (SpO$_2$ < 90%) |

**Procedure**

1. Instruct the patient to keep a sleep log for 1 to 2 weeks before the polysomnogram (PSG).
2. Remind patient that on the day of the study, caffeinated beverages, alcohol, and sedatives are not permitted.
3. Be aware that extra time is needed to set up and attach equipment to the patient. Typically, the PSG is recorded during the patient's normal sleep time; however, partial or extended periods of sleep deprivation may be necessary if seizure activity is suspected.
4. Be aware that the sleep technologist records the patient’s history and factors such as age, height, weight, current medications, visual problems, and history of seizures, head injuries, headaches, or strokes. The sleep log is reviewed, and a bedtime questionnaire is completed. The patient wears normal bedtime attire.
5. Use the following list to identify the monitoring equipment used:
   a. Two sets of scalp electrodes to monitor sleep stages (EEG)
   b. One electrode to the outer canthus of each eye (EOG)
   c. One electrode to the chin (submental)
   d. Electrodes to the legs (anterior tibialis; EMG)
   e. ECG leads for heart rhythms and rates
   f. Impedance monitor (respiratory effort)
   g. Oral/nasal thermistor between nose and upper lip (air flow)
   h. Pulse oximeter (SpO₂; O₂ sensor)

6. After application, interface all electrode leads with a “jack box,” which contains the preamplifiers and impedance meter. From the jack box, signals are sent through additional amplifiers and filters and finally to a multichannel recorder or polygraph. The polygraph can provide a hard copy recording of all channels and signals that can be computer processed and displayed on a monitor. Electrode connections are subsequently tested for integrity and adjustments made before the patient retires.

7. Be aware that during the recording, both audio and infrared camera video recordings are made.

8. Provide a bedside commode because the leads are relatively short.

9. When the test is completed and equipment removed from the patient, ask the patient to complete another questionnaire; score related to the patient's sleep experience during the test.

10. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

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**NOTE**

If seizures are a factor, up to 16 additional scalp electrodes are applied according to the International 10-20 System of Electrode Placement. The International 10-20 System of Electrode Placement is the conventional system (established in 1958) used to identify and place scalp surface electrodes for the recording of brain electrical potentials. The 10-20 System nomenclature is used because most of the electrodes are placed either 10% or 20% between specific skull landmarks (eg, the nasion or inion) or in relation to the circumference of the head.

**Clinical Alert**

A home sleep study is an alternative for patients who have trouble falling asleep in a laboratory. Sensors are applied in the clinic, and the patient is shown how to attach the mobile monitoring unit.

**Interfering Factors**

1. Caffeinated beverages and alcohol can delay sleep onset or exacerbate some types of sleep disorders.
2. Sedatives (hypnotics) shorten sleep onset and reduce nocturnal awakenings, which may skew the results of the PSG.
3. Changes in daily routine on the day of the sleep study may cause false-positive or false-negative results.
4. During the PSG, environmental noise, lights, and temperature may have an adverse effect on the patient's ability to fall asleep.

**Clinical Implications**

1. Abnormal EEG recordings indicate problems with either sleep architecture (eg, sleep onset, stages, offset) or seizure disorders.
2. Abnormal leg EMG is consistent with movement disorders (eg, restless-leg syndrome, nocturnal myoclonus, leg jerks).
3. A respiratory sleep index (RDI) > 5 indicates sleep-disordered breathing. Obstructive sleep apnea (OSA) is characterized by absence of airflow for > 10 seconds despite continued respiratory effort (eg, thoracic breathing or snoring accompanied by periods of apnea). Central sleep apnea (CSA) is characterized by absence of both airflow and respiratory effort; airflow ceases because respiratory effort is absent. Mixed sleep apnea (MSA) generally begins as a central apnea and becomes obstructive apnea. Sleep apnea has been linked with cardiac arrest, strokes, pulmonary hypertension, brain stem lesions, and head trauma.
4. An oxygen desaturation index (ODI) > 5 is associated with oxygen desaturation, which generally occurs with an apneic event but can also occur with hypoventilation.

**Interventions**

**Pretest Patient Care**

1. Explain test purpose and procedure. These tests are done when signs and symptoms have persisted for at least 6 to 12 months. Caution the patient not to change his or her daily routine the day before the test.
2. Reassure the patient that lead wires, monitors, and sensors will not interfere with changes of position during sleep.
3. Record the patient's age, height, weight, and gender. A brief history and before- and after-bedtime questionnaires are taken.
4. Have the patient prepare for sleep at the normal time according to routine and discontinue any medications used to help with sleep.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Have patient resume usual activities and routines.
2. Interpret test outcomes and monitor appropriately. If test results indicate OSA, explain possible need for further treatment.
3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Sleepiness Tests; Multiple Sleep Latency Test (MSLT); Maintenance of Wakefulness Test (MWT)**

The multiple sleep latency test (MSLT) is used as an objective measure of excessive daytime sleepiness and determines its severity. Typically, the MSLT is administered the morning following sleep study. An alternative to this test is the maintenance of wakefulness test (MWT), which measures the ability of an individual to stay awake rather than to fall asleep. Both the MSLT and MWT are used to diagnose narcolepsy and to evaluate the effectiveness of pharmacologic interventions in the treatment of daytime hypersomnia. Indications for these tests include falling asleep at inappropriate times, daytime hypersomnia, suspected narcolepsy, and evaluation of drug effectiveness in treating various sleep disorders. The MSLT is an objective measure of a patient's sleepiness and is done to evaluate the severity of daytime sleepiness, to diagnose narcolepsy or falling asleep at inappropriate times, and to evaluate effectiveness of drug therapy for daytime hypersomnia. The MSLT is administered after a sleep study to rule out any sleep-related pathology that might affect the results and to assess the quality of sleep. An alternative to the MSLT is the MWT, which measures the ability of a person to stay awake rather than to fall asleep. The MSLT includes the following tests:

1. EEG: at least 2 channels are recorded to determine sleep onset, sleep stages, and sleep offset.
2. EOG: to document both slow and rapid eye movements present at sleep onset and during REM sleep, respectively.
3. EMG: the chin EMG is used as a criterion for REM sleep.
4. ECG: to monitor heart rate and rhythm.

The MWT includes the following tests:

1. Electromyogram (EMG): the chin EMG is used as a criterion for REM sleep.
2. Electrocardiogram (ECG): to monitor heart rate and rhythm.

**Reference Values**

**Normal**
- MSLT: average sleep latency is 10 to 20 minutes
- MWT: Average sleep latency on the 40-minute test is 35 minutes; Average sleep latency on the 20-minute test is 18 minutes

**Procedure**

1. Remember that typically, the MSLT or MWT is administered the morning following a sleep study. Following the sleep study, have the patient dress, eat (avoiding caffeine), and report back to the sleep laboratory.
2. Reapply the electrodes if necessary.

3. Be aware that the first nap (for the MSLT) or first session (for the MWT) will begin 1.5 to 2 hours after morning awakening, with a minimum of four additional naps or sessions at 2-hour intervals throughout the day.
4. Terminate the nap or session after 20 minutes for the MSLT, or after 20 to 40 minutes for the MWT. With the MSLT test, if the patient falls asleep, continue the recording for 15 minutes after sleep onset.
5. Instruct the patient to allow himself or herself to fall asleep or not to resist the urge to fall asleep for the MSLT.
...whereas for the MWT, instruct the patient to resist the urge to sleep or to attempt to remain awake.
6. Between the naps or sessions, ensure that the patient remains awake and encourage moving around.
7. Following the testing, disconnect all equipment and discharge the patient.
8. Have the technologist score the MSLT or MWT in conjunction with the PSG test results.
9. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

**NOTE**

The term “nap” indicates a short intentional or unintentional episode of subjective sleep taken during habitual wakefulness, whereas the term “falling asleep” or “sleep onset” is defined objectively by electroencephalographic recordings (EEG) (ie, stage 1 of nREM sleep).

**Interfering Factors** Caffeinated beverages can delay sleep, whereas sedatives (hypnotics) shorten sleep onset. Additionally, sleep deprivation may result in a false-positive MSLT result. During naps, environmental noise, lights, and temperature can have an adverse effect on the patient's ability to fall asleep.

**Clinical Implications**

1. An average sleep onset of 6 to 9 minutes in the MSLT is considered a “gray area” diagnostically because these tests are done in a laboratory setting and not in the patient's home environment. Reevaluation may be necessary if the patient complains and symptoms persist.
2. An average sleep onset < 5 minutes and two or more REM periods in the five to six naps during the MSLT is diagnostic for narcolepsy. This indicates a disturbance of the normal sleep architecture pattern, although the REM periods are not unlike nocturnal REM periods. These REM episodes, however, occur prematurely in the cycle and are termed sleep-onset REMs (SOREMs).

**Interventions**

**Pretest Patient Care**

1. Explain MSLT or MWT purpose and procedure. Remind the patient not to change daily routines the day of testing.
2. Reassure the patient that lead wires, monitors, and sensors will not interfere with sleep.
3. Record the patient’s age, height, weight, and gender.
4. Remind patient that no alcohol or caffeinated beverages should be consumed the day of the test.
5. Administer standard sleep questionnaires or scales (eg, Epworth Scale, Stanford Scale) and evaluate (see Appendix H for examples).
6. Follow guidelines in Chapter 1 regarding safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Explain test outcome and possible need for follow-up testing.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Nasal Continuous Positive Airway Pressure (nCPAP) Titration** Following the diagnosis of OSA, this test is done before treatment is begun. The nCPAP machine supplies air under pressure, acting as a pneumatic splint that keeps the upper airway open during sleep. The pressure required depends on the severity of the OSA and can vary; therefore, the patient is typically required to return to the sleep laboratory on a second night to repeat the sleep study (PSG) while wearing an nCPAP mask. Positive airway pressures are increased until the apneas “break.” This procedure is referred to as nCPAP titration. Under some circumstances (eg, severe sleep apnea), titration can be done on the same night as the PSG. In that case, it is termed a “split-night study.” The nCPAP machine provides continuous positive pressure during both inspiration and expiration. Conversely, bilevel positive airway pressure (nBiPAP) uses two separate pressures: one during inspiration and a lower pressure during expiration. Conversely, bilevel positive airway pressure (nBiPAP) uses two separate pressures: one during inspiration and a lower pressure during expiration. In cases in which nCPAP is not well tolerated, nBiPAP may be a better alternative. An nCPAP unit may be used in the home and preset to the test pressures that ameliorated the apneas.

**Reference Values**

**Normal**

- Electroencephalogram (EEG): normal time to sleep onset, sleep stages, and sleep offset
- Chest impedance
- Sleep log

**Procedure**

1. On the day of the titration, instruct the patient to avoid caffeinated beverages, alcohol, and sedatives, and to keep a sleep log.
2. Allow sufficient time before testing to attach the patient to the monitoring devices and other equipment, including the nCPAP machine. A brief orientation to nCPAP should take place before the actual day of titration to relieve the patient's anxiety.
3. Remember that the sleep technologist takes a brief patient history. The sleep log is reviewed, and a bedtime questionnaire is completed (see Appendix H). The patient then prepares for sleep.
4. Have the technologist apply the electrodes, monitors, sensors, and microphone, and interface these with the other electronic devices (see Polysomnography [PSG], page 1067).
5. Fit the patient with an nCPAP mask and ensure that it can be easily removed in case of discomfort, shortness of breath, or claustrophobia.
6. Provide a bedside commode because the leads are relatively short.
7. Adjust CPAP pressures throughout the sleep period, beginning with 3 to 5 cm H₂O and increasing in 2.5-cm H₂O increments until the apneas “break.” Time increments can vary from 15 minutes to 2 hours per pressure setting. Decisions are based on protocols being used, severity of sleep apnea, and patient tolerance for testing. If nBiPAP is being performed, inspiratory and expiratory pressures are adjusted separately, keeping the inspiratory pressure at least 2 to 4 cm H₂O above the expiratory pressure.

8. After the test, remove the equipment and have the patient complete another questionnaire, which the sleep technologist evaluates and scores.

9. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

Interfering Factors

1. Caffeinated beverages and alcohol can delay sleep onset or exacerbate OSA, which may interfere with determining optimal pressure settings.

2. Changes in the patient's daily routine on the day of titration can alter results.

3. Patients with a deviated nasal septum or chronic sinusitis may have problems tolerating the nCPAP. The use of nCPAP is contraindicated in persons with severe bullous emphysema or chronic perforated tympanic membrane.

4. Skin irritations from tight-fitting masks (especially on the bridge of the nose), nasal congestion, and headaches are occasional complaints with the use of nCPAP.

5. The benefit of nCPAP to patients with central sleep apnea has not been well documented.

Clinical Implications

1. An RDI > 5 indicates OSA, which is characterized by the absence of airflow for > 10 seconds in the presence of continued respiratory effort. nCPAP used in treating OSA has been shown to be clinically beneficial.

2. Following even short-term nCPAP use, there is documented evidence of rapid symptomatic improvement, with restoration of nocturnal sleep and subsequent improvement in lessening of daytime sleepiness and improving quality of life.

Interventions

Pretest Patient Care

1. Explain test purpose and nCPAP titration procedure.

2. Reassure patients that the mask can easily be removed if anxiety or claustrophobia develops.

3. Record the patient's age, height, weight, and gender. A brief history is taken, and before- and after-bedtime questionnaires are filled out.

4. Have the patient prepare for sleep at the normal time in the usual manner.
5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

**Posttest Patient Aftercare**

1. Explain test outcome and possible need for follow-up testing and treatment. Depending on the test outcome, an nCPAP unit may be ordered for home use.
2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

**Gastric Analysis (Tube Gastric Analysis); Gastric Fluids**

This test examines stomach contents for abnormal substances and also measures gastric acidity. It aids in diagnosing ulcers, obstructions, pernicious anemia, or carcinoma of the stomach. It can determine the cause of gastrointestinal bleeding as well as the effectiveness of medical or surgical therapies. Examinations of gastric washings (eg, tuberculosis studies) can identify mycobacterial infection when previous sputum tests have been negative.

**Reference Values**

**Normal**

Fluid: clear or opalescent; no food, blood, drugs, or bile present in sample. pH: 1.5–3.5 Culture: negative for mycobacterial organisms

Fasting specimen total acidity: <2 mEq or <2 mmol/L

Basal acid output (BAO) without stimulation: 0–5 mmol/hour or 0–5 mmol/hour

Maximal acid output (MAO) or normal secretory ability when using a gastric stimulant such as histamine or betazole hydrochloride intramuscularly or pentagastrin subcutaneously: 10–20 mEq or 10–20 mmol/hour

BAO-to-MAO ratio: 1:2.5 to 1:5

**Procedure**

1. Collect fasting gastric analysis specimens during endoscopy (see Chapter 12) or through a nasogastric (NG) tube inserted for the test. Follow NG tube institutional protocols.
2. Aspirate initial gastric acid through the NG tube with a syringe, test for pH, and discard. If no acid is present, reposition the NG tube and obtain another specimen.
3. Remember that specimens are normally collected via continuous intermittent low suction over 1 to 2 hours at 15-minute intervals, depending on the type of gastric stimulant given. Each specimen is placed in a separate specimen cup and labeled BAO or MAO, along with patient's name, date, and time collected.
4. Remove the NG tube after all specimens are collected.
5. Document the date and time; type of procedure; type and size of tubes used; number of specimens collected; appearance, consistency, and measured volumes of gastric fluid obtained; the patient's response to testing; complications; interventions; and other pertinent information.
6. Follow guidelines in Chapter 1 for safe, effective, informed intratest care.

**Clinical Implications**

1. Decreased levels of gastric acid (hyposecretion and hypochlorhydria) occur in the following conditions:
   a. Pernicious anemia
   b. Gastric malignancy
   c. Atrophic gastritis
   d. Adrenal insufficiency
   e. Vitiligo
   f. Rheumatoid arthritis
   g. Thyroid toxicity
   h. Chronic renal failure
   i. Postvagotomy
2. Increased levels of gastric acid (hypersecretion and hyperchlorhydria) occur in the following conditions:
   a. Peptic or duodenal ulcer
   b. Zollinger-Ellison syndrome
   c. Hyperplasia and hyperfunction of antral gastric cells
   d. Post–small intestine resection

**Interfering Factors**

1. Lubricants or barium from previous tests present in sample affects the result.
2. Medications such as antacids or histamine blockers, foods, and smoking alter gastric secretions.
3. Gastric secretions are altered in patients with diabetes who use insulin or in those who have had surgical vagotomy.
4. Elderly patients have lower levels of gastric hydrochloric acid.

**Interventions**

**Pretest Patient Care**

1. Assess for contraindications to the procedure, including carcinoid syndrome, congestive heart failure, recent myocardial infarction, or hypertension. The use of histamine may exacerbate these conditions.
2. Explain test purpose and procedure. Inform patient that there may be some discomfort and possibly a gagging sensation when the nasogastric tube is inserted and that a gastric stimulant may be injected. Devise a method of communication for the patient before insertion of NG tube (eg, raise index finger to indicate “wait” before proceeding). Explain that panting, mouth breathing, and swallowing facilitate tube insertion.
3. Record baseline vital signs. Remove dentures before test.
4. Have the patient fast from food, fluids, smoking, and gum chewing for at least 8 to 12 hours before testing.
5. Restrict or withhold anticholinergic agents, cholinergic agents, adrenergic blockers, antacids, steroids, alcohol, and caffeine for at least 24 hours before testing. Check with the clinician well before the procedure.
6. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
Clinical Alert

1. If histamine is injected, the patient may experience flushing, dizziness, headache, faintness, and numbness of the extremities and abdomen during or immediately after testing. These symptoms must be reported immediately. Have epinephrine easily available.
2. Specimens for acid-fast bacillus (AFB) and tuberculosis (TB) cultures must be warm and should be taken to the laboratory immediately. Laboratory personnel should be alerted.

Posttest Patient Aftercare

1. Monitor vital signs. Observe for possible drug side effects, gastrointestinal bleeding, or respiratory distress (gastrointestinal bleeding may signal perforation).
2. Provide nasal and oral care after tube removal. Allow patient to rest. Provide food or fluids as tolerated and ordered. If a local anesthetic was used on the throat, assess for return of gag and swallow reflexes before allowing patient to drink or eat (usually 2 hours postadministration.)
3. Interpret test outcomes. Counsel patient regarding possible lifestyle alterations such as smoking cessation, restricted alcohol intake, dietary changes, stress reduction, medication, medical treatment, or surgical intervention.
4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

DNA Typing or Fingerprinting

Deoxyribonucleic acid (DNA) is a complex, high-molecular-weight protein composed of deoxyribose, phosphoric acid, and four bases (adenine, guanine, thymine, cytosine). These six substances are arranged in two long chains that twist around each other to form a double helix. The complementary components on each of these two chains link together between the chains. The nucleic acid component is present in the cell nuclei chromosomes and forms the chemical foundation for heredity. It carries the genetic material for every living organism except RNA viruses. DNA provides the actual code for individual genetic characteristics through a specific sequence, or “blueprint,” that is unique to that person alone. DNA testing is used to establish identity (ie, military and disaster casualties), to determine parentage (ie, infant abductions), and during immigration disputes and criminal investigations (ie, murder, sexual abuse, rape) through a process termed restriction fragment length polymorphism (RFLP). This process allows evaluation of different DNA tissue samples from several sources to determine matching patterns, similar to comparing bar codes. Polymerase chain reactions (PCR) techniques have become widely popular and have advantages over previous techniques. Less DNA is required, and typing can often be done with partially degraded DNA.

Reference Values

Normal Specific and unique to each person Each person has a unique DNA profile.

Procedure

1. Remember that DNA can be extracted from dried whole blood or any tissue that contains nucleated cells (eg, skin, saliva, hair shafts, urine, semen.) More cellularly dense tissues produce more DNA. Bone and teeth are the most stable sources for postmortem DNA.
2. Collect venous blood samples in a yellow-topped (ACI) tube or lavender-topped (EDTA) tube. For tissue samples, 0.1–1.0 gram of tissue is obtained and placed in a plastic freezing bag. Freeze tissue samples, and keep frozen until shipped in dry ice. Do not freeze blood samples.
3. Process DNA samples until DNA fragments can be visually represented on x-ray film. These films are called autoradiographs or autorads. At this point, the fragments somewhat resemble bar codes.
4. Compare the autorads for matching or nonmatching characteristics among several samples. If a match between two or more different autorads is found, there exists a high probability that the different samples come from the same source person.

Clinical Alert

1. The DNA samples are collected and stored using great care with pristine equipment to prevent contamination and preserve specimen that may have crucial legal implications.
2. Samples of DNA can be stored for indefinite periods.

Clinical Implications

1. Identity is confirmed when there are matching patterns in certain areas of the autorads. Adhere to caution when collecting and storing DNA specimens to prevent contamination and to preserve the specimens, which may have crucial legal implication.
2. In parentage studies, even though each person has a unique DNA profile, matching characteristics in certain areas of autorads that come from two different individuals can indicate a parent–child relationship.
3. In criminal cases, matching DNA characteristics associated with tissue samples retrieved from both victim and suspect may establish the suspect's presence at the crime scene. A nonmatch definitively disproves that the different samples came from the same person.

Interfering Factors

1. Insufficient amount of DNA
2. DNA tissue sample deterioration/degradation
3. Lack of material database to conduct effective sample comparison

Interventions

Pretest Patient Care

1. Explain the DNA test purpose and procedure with the concerned individual and family members, being mindful of privacy and confidentiality.
2. Be aware that there must be no transfusion 90 days before testing.

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<tr>
<td>1. Communicate results in confidence.</td>
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<td>2. DNA specimen collection identification, packaging, and storage are extremely important. In forensic studies, the issue of chain of custody arises.</td>
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Posttest Patient Aftercare

1. Counsel appropriate persons about the meaning of test results.
2. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

POSTMORTEM TESTS

It is the basic civil right of deceased human beings to have competent medical investigations of their deaths. This is particularly true in our 21st century environment of social ills, drug use, crime, violence, bioterrorism, and genetic relationships to disease and mortality among all sociocultural and political classes. Any death has potential civil, legal, criminal, or economic implications for the deceased, the family and significant others, and society globally.

As with establishing a medical diagnosis for a living person, the medical model is similarly executed in developing a postmortem diagnosis. A history is taken, consisting of medical history, risk factors, and death interpretation; a physical examination is performed (ie, autopsy); and laboratory tests are interpreted (tissue, organ, postmortem blood, and other body fluids such as vitreous fluids, urine, bile, CSF, or gastric contents).

Death Investigation

All deaths, whether from a natural sequence of events, during medical treatment, in unexplained circumstances, or criminally related, need to be investigated regarding cause and manner of death so that the legal death certificate may be accurately completed, signed, and recorded. Deaths can be defined as natural or medical-legal:

**Natural death** is the cessation of cardiorespiratory function due to a medical disease process (eg, metastatic cancer, cerebrovascular accident) or natural progression of life events (ie, “old age”).

**Medical-legal death** results from some “unnatural” (unexpected, unusual, or suspicious) event such as homicide, suicide, or accident; this situation is specifically governed by legal statutes and requires that a coroner, medical examiner, and law enforcement officials be involved.

Death Interpretation

The process of postmortem examination and interpretation follows a certain defined protocol. First, the history portion of the investigation is obtained. In natural death without autopsy, the medical history, diagnostic tests before death, clinical record, and knowledge about lifestyle can provide reasonably sufficient information to arrive at conclusions regarding cause and manner of death. Only autopsy, however, can definitively confirm these suspicions and initial conclusions.

Medical-legal death investigation first focuses on the death scene. Interviews lay the groundwork for the investigation. Evaluation of the site of death and other physical evidence involves a detailed examination of blood stains, disrupted environment, position of body, and signs of struggle or injury manifested by fluids leaking from body orifices, eyes, ears, nose, or mouth. Color changes and rigidity can be beneficial in establishing time frames for events. Interviews of known witnesses are a part of the investigation.

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<td>Postmortem examination falls under the domain of the physician pathologist. In criminal cases, forensic pathologists who have specialized knowledge, skills, and state-of-the-art investigative techniques at their disposal should perform the autopsy.</td>
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The second part of the death interpretation process involves the autopsy itself. The autopsy consists of a detailed comprehensive physical examination of both external and internal body features of tissues, organs, and fluids.

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<td>In the case of a medical-legal autopsy, samples of organs and specimen sections (slides) should be retained for 3 to 5 years pending outcomes of litigation and legal system appeals. Photographs should accompany reports and should be archived.</td>
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Time of Death

Although time of death is usually not a major issue, determination of time of death is important in both natural deaths (for insurance and other death benefits) and unnatural deaths, either witnessed or when body parts have been naturally or intentionally altered so that an individual's distinguishing features are altered. Time of death estimate is based on presence of the following:
1. **Rigor mortis.** Stiffening of body as pH changes and lack of adenosine triphosphate in muscles occur; rigidity appears anywhere from instantly to 6–12 hours after death.
2. **Livor mortis.** Red-purple color caused by settling of blood in dependent body parts due to gravity; onset immediate, sometimes beginning before death (maximum in 8–12 hours).
3. **Algor mortis.** Cooling of body; body temperature is subject to interpretation, based on cocaine use, presence of infection or fever before death, death scene, heat absorption, amount and type of clothing, size of body, activity just before death, and decomposition.
4. **Decomposition.** Processes occur as a result of chemical breakdown of cells and organs due to intracellular enzymes and putrefaction due to bacterial action.
5. **Gastric emptying.** Food in stomach; digestion and stomach emptying varies in both life and death.
6. **Chemical changes.** Potassium in vitreous fluid of the eye; as the time since death becomes greater, the concentration of potassium increases.
7. **Insect activity.** Flies and other insects are associated with decomposed bodies; any attempt to fix time of death using insect evidence should be done only with the aid of an entomologist.

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<tr>
<td>1. Time of death is expressed as an estimate of the time range during which death could have occurred and varies according to cooling of body, color change, decomposition of stomach contents, clinical changes, and insect activity.</td>
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<tr>
<td>2. There is no single accurate marker of time of death.</td>
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When the time interval between actual death and the initial death investigation is months or years, body changes may be quite variable and can include saponification of subcutaneous tissue (changes due to prolonged exposure to moisture taking several months); mummification (drying process due to lack of moisture; occurs quickly in hot, dry climate, exposure to air, dying of thirst); and skeletonization (takes months to years). Examination of bones may yield general knowledge of deceased (eg, estimate of age, stature, race, gender).

Physical evidence such as mail delivery, telephone calls answered and unanswered, “last contact times” can help the death investigator estimate time of death.

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<tr>
<th>Clinical Alert</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. If the HIV status of decedent is not directly related to the cause of death or an exposure to another, the decedent's HIV status need not be recorded or reported.</td>
</tr>
<tr>
<td>2. Some examiners may be reluctant to include HIV-positive information in public record autopsy reports out of compassion for the decedent's family. However, certain states require that every autopsy done through the medical examiner's office include an HIV test. As with other issues of confidentiality, the applicable laws must be obeyed; however, the pathologist must make every effort to protect the patient's and family's right to privacy to the degree that the law will allow. Consideration of “exposure” of the decedent's blood/ saliva/body fluids to other health care workers, including EMTs, must be given.</td>
</tr>
</tbody>
</table>

Cancer is reported when the cancer is first discovered or confirmed upon autopsy.

The third step in death interpretation includes collection of blood, bile, urine, and ocular fluids (if available) for analysis. Blood toxicology studies must be performed at a state-certified laboratory. Blood samples are the only life/death determinants; urine samples provide information about levels of substances excreted, but they do not provide blood level values of these same substances.

Typical blood toxicology screens include the following tests:

1. Alcohol screens determine levels of various alcohols.
2. Acid-neutral screens detect barbiturates and salicylates.
3. Basic screens detect tranquilizers, synthetic narcotics, local anesthetics, antihistamines, antidepressants, and alkaloids.
4. Higher volatile screens use gas chromatography to detect substances such as toluene, benzene, trichloroethane, and trichloroethylene.
5. Cannabis screens detect the presence of cannabis (marijuana).

All of these screens serve a purpose in cases that involve accidents and work-related deaths; they also serve to rule out foul play in otherwise natural deaths.

After the life events and scene investigation, the autopsy, and the laboratory tests are completed, data from all sources are integrated, scrutinized, and analyzed. Findings and conclusions are then documented and certified on the death certificate, which then becomes a matter of public record. Findings may then be shared with the decedent's immediate family, significant other, or legal authorities, or may become part of a court deposition process.

**Autopsy** An autopsy is an investigation of the cause and manner of death by direct examination of the body. **Cause of death** is the disease or injury that, through its physiologic effects, results in the actual death of the individual. **Manner of death** is the type of event that led to death and is categorized as natural, homicidal, suicidal, accidental, pending, or
undetermined. Interpretation of physical findings results in setting forth an opinion regarding the probable cause of death. Before autopsy, as much pertinent information as possible is gathered about the deceased. Available medical records are reviewed thoroughly. In cases of medical-legal death investigation, not only the medical and social background but also the terminal events and circumstances of death, including the environment, presence of drugs and alcohol, and exact condition and position of the body, are thoroughly investigated (scene investigation of physical evidence and personal contacts).

Reference Values

**Normal**

External and internal findings: within normal limits or demonstrate significant pathology related to cause of death. Gross and microscopic findings: within normal limits or abnormalities related to cause of death. No drugs, alcohol, or other metabolites present in laboratory or toxicology screens.

**Procedure**

1. Observe standard precautions throughout. Identify the body and tag (usually on the great toe) with the decedent's name (if available), gender, age, and a number. The body is then weighed and measured.
2. Photograph the head and chest and mark with an identification number. This step occurs in nonhospital, nonclinical institution deaths (eg, deaths at home, work, school, industry, roadway, whenever foul play is suspected, and whenever a 911 call results in law enforcement officers at the death scene). Identification marking of bodies is particularly important in mass casualty disasters such as plane crashes and building disasters.
3. Describe and record information about clothing and valuables. When the body is found by someone other than a family member (eg, at work, highway or airplane accidents), these items are removed, inventoried, and given to family or law enforcement agency.
4. Take fingerprints only in criminal cases or in unidentified victims.
5. Cleanse the body. In trauma cases or unusual death findings, the face is photographed again if blood, dirt, and other materials were present initially.
6. Perform an external examination on the entire body. Record the location and description of all identifying marks, scars, tattoos, incisions, injuries, deformities, and other significant findings on a body diagram.
7. When foul play is suspected, photograph all injuries from at least two views: one showing the location of the injury on the body and the other providing a close-up view of the injury. Take and record measurements of wounds.
8. Be aware that in some instances, x-ray films may be necessary to verify gross anatomic deformities, injuries (cervical spine and skull fracture), or pathologies that may provide clues regarding the cause of death. Radiography tracks the trajectory of bullets and other projectiles through the entire body or just through a specific area and may also be performed on exhumed organs or decomposed bodies. In some instances, bodies may be completely unrecognizable (eg, putrefied beyond recognition). X-ray studies can determine age and can establish a victim's identity by comparing bone and dental detail to previous x-ray films and dental x-rays of the victim.
9. Ensure that the autopsy proceeds in an orderly manner, observing standard precautions.
10. Document descriptions of color and distinguishing features of the hair and eyes and appearance of the nose, ears, mouth, teeth, face, head, neck, genitalia, torso, and extremities in detail. Examine the front, side, and back of the body thoroughly. Injuries, wounds, bruises, contusions, and lacerations are described, mapped, measured, and detailed. Include descriptions of size, depth, location, and presence of foreign objects or materials at or near the injured areas, as well as fluids draining from body orifices and wounds, in the report.
11. Be aware that internal examination includes a complete head and pelvic dissection with removal of all organs from the skull, neck, abdomen, and pelvis. Specific organs are subjected to gross examination that includes measurement of size and actual weight. Once this is done, take organ sections and prepare for microscopic slides to be examined later. Save the slides for evidence. Virtually any part of the body can be microscopically examined. The brain and the neck organs are always removed and examined. As part of this examination, the dura matter is removed to permit visualization of the skull and calvarium to determine injury or natural pathology.
12. Withdraw blood and fluid specimens by syringe from the heart, aorta, eyes (vitreous fluid), gallbladder (bile), and bladder (urine). Refrigerate these until examined; they can be saved for an indefinite period. In the instance of trauma, blood samples can be retrieved from the pulmonary trunk or the chest. If clots are present and syringe sampling cannot be done, pericardial tapping is an alternative method of procuring a blood sample.
13. Be aware that sometimes it is necessary to collect specimens for culture. Most internal organs of previously uninfected persons remain sterile for about 20 hours after death.
14. Return the organs to the body after examination is completed. Organs and tissues may be donated for transplant procedures.
15. Following completion of autopsy and possibly organ/tissue procurement, release the body immediately to the funeral home for burial or cremation per family wishes. If there are legal questions, the body may be stored in a frozen state for some time (eg, months).
Clinical Alert

Evidence, which may include clothing, a toothbrush, eyeglasses, and other personal (physical evidence) items, should be transported to the proper designated laboratory as quickly as possible, and refrigerated or frozen if delayed longer than a week. Examples of unusual evidence material that has yielded successful DNA profiles include:

From hands: automobile armrest, baseball cap brim, bottle cap, chocolate bar (the handled end), dime, doorknob, electrical cord, ignition switch, keys, pen, seat belt buckle, shoelaces, gun (trigger, stock, handgrips), knife handle, hammer, wrench, screwdriver, ice pick, and saws

From mouth and nose: bite mark, chicken wing, envelope, glass rim, lipstick, food item, ski coat collar, telephone receiver, welding goggles

From eyes: contact lens fragments in vacuum cleaner, tears on a tissue, eyeglasses

From the body in general: burned remains, hair comb, automobile head rest, razor, shirt, underarms, socks, urine in snow

Appendix L describes methods for collecting for evidentiary specimens. DNA extracted from the victim or related evidence is compared with samples taken from a suspect. The odds of a match are then determined by looking at nine different chromosome pairs and examining microscopic sequences of DNA. The odds of two people sharing the exact sequence on one chromosome pair is determined; then it is multiplied by the odds of two people sharing the exact sequence on two pairs, three pairs, and so on to nine pairs. The odds reach awesome proportions because those nine sections of DNA are highly variable among individuals. Scientists can be more certain when they are eliminating suspects. If analysts find a variation on any of those nine DNA locations, they can say with accuracy that the two samples could not have come from the same person.

Protocols for Retrieving Postmortem Cultures

1. Follow standard precautions. See Appendix A.
2. Use sterile instruments and gloves when obtaining specimens for culture.
3. Cleanse the area with a povidone-iodine 5-minute scrub followed by a 70% alcohol 5-minute scrub.
4. For sample collection, either aspirate body fluid samples and transfer them to a sterile tube or swab the area with sterile swabs.
   a. Obtain blood culture specimens from the right ventricle of the heart.
   b. Collect peritoneal fluid immediately after entering the peritoneal cavity.
   c. Collect bladder urine directly from the bladder with a syringe and needle.
   d. Sample pericardial or pleural cavity fluid on a swab or with a syringe and needle.
5. Sear the external surface of an abscess to dryness with a red hot spatula; collect pus via syringe and needle (if possible) or use a swab.

Clinical Alert

1. If an external examination or other noninvasive studies are done in lieu of an autopsy, collect blood from the subclavian vessel and vitreous humor from the eyes.
2. Do not use plastic envelopes for storing biologic samples such as tissue and hair or foreign objects such as bullets. Plastic captures moisture and promotes bacterial/fungal growth. Instead, place objects in clean paper envelopes or brown paper evidence bags. Inventory and label each item properly; store appropriately.

Special Procedures

1. Gunshot wound procedure
   a. Remember that there is a mandatory x-ray of all gunshot wounds, including entrance and exit; locate bullet and fragments.
   b. Photograph entry and exit; cleanse wound; repeat photograph.
   c. Obtain samples of gunpowder residue on hands (close-range only wounds).
2. Blunt force injury procedure
   a. X-ray affected areas. X-ray hands and forearms for “defense wounds.”
   b. Photograph the wounds in original condition (after cleansing). Use rape kit if rape is suspected (both male and female) or when the nature of the injury suggests uncontrolled rage (hammer, axe, stab wounds) or other blunt force (beatings, clubbings).
3. Sharp wound procedure
   a. X-ray wound sites. Photograph wounds in original condition (after cleaning), after approximating wound margins.
   b. Check for “defense wounds” on hands and arms. Trace wounds on clear plastic sheet (optional). Save and photograph severed cartilage.
4. Drug overdose procedure
   a. Photograph evidence suggesting drug abuse, such as injection sites on body; presence of drug paraphernalia or drugs, and drug residue on lips, face, teeth, oral cavity, tongue, nose, or hands.
   b. Assess mouth area and body for bite marks and “fall” injuries (suggests seizure activity associated with drug ingestion). Check lymph nodes, spleen, liver (abnormal in intravenous drug abuser).
5. Special battery procedure. If sexual assault is suspected, perform the following in order:
   a. With body supine (face up), obtain scalp and pubic hair, oral samples, semen from inner thighs.
   b. With body prone (face down), obtain anal specimen first, then vaginal specimen.
   c. Collect fingernail evidence.
   d. Collect 25 pubic hairs from entire vulvar area.
e. Collect 25 head hairs from affected area.
f. Obtain oral, anal, cervical, and other specimens suspected of containing semen.

6. Child abuse/sudden infant death syndrome (SIDS) procedure
a. X-ray and photograph entire body.
b. Perform external examination of conjunctival petechiae, fingertip bruises, torso and shoulders (front/back), frenulum, and back; posterior thighs and buttocks may be incised from buckles or other sharp objects.
c. Perform internal examination for hematomas (due to direct injury): if present and no evidence of head trauma, remove eyes and examine retina (shows characteristic signs in presence of sudden infant death syndrome).
d. Document recent or healed fractures and estimated time of injury.
e. Re-examine and re-photograph the following day to delineate bruises not previously evident.

NOTE
Copper- and aluminum-jacket bullets remain in the body. Aluminum jackets are difficult to visualize on x-ray films, especially when lodged in bone.

Clinical Alert
1. Observe standard precautions during these procedures. Risk for disease transmission, hepatitis, TB, and HIV exposure is high.
2. Sketch, measure, and mark wounds on diagram in both inches and centimeters (U.S. residents relate to inches more accurately). Projectile (bullet) caliber is estimated as small, medium, or large.
3. Procure toxicologic specimens if indicated. Specimens for toxicologic analysis include the following:
   a. All ocular fluid from both eyes
   b. Blood—sodium fluoride preservative (50 mL)
   c. Blood—sodium fluoride preservative (retainer tube) (10 mL)
   d. Liver (3 g)
   e. Bile (10 mL)
   f. Urine (50 mL)
   g. Stomach and small bowel contents
4. Store specimens and fragments in paper envelopes or bags; never use plastic, which allows bacteria, mold, and fungus to grow.
5. Route specimens and reports to appropriate department or individual.

Metabolic Autopsy The reported frequency of sudden infant death syndrome (SIDS) is approximately 1 in 1000 live births (25% of all reported deaths in the first year of life in industrialized nations). A specific medical diagnostic cause of death in such cases does not result even following a complete and thorough autopsy. Growing research initiated in the mid-1980s suggests that metabolic derangement is implicated in SIDS deaths. A clinical picture of a child with lethargy, vomiting and/or fasting, and previous sibling death, coupled with postmortem determination of diffuse fatty changes in the liver as determined from tandem mass spectrometry analysis of blood and bile specimens, is implicated in such cases of sudden unexpected childhood death. This genetic error in metabolism is hypothesized to be related to medium-chain acyl-CoA dehydrogenase deficiency and disorders of mitochondrial fatty acid beta-oxidation. The required sophisticated laboratory analyses are limited to highly specialized laboratories in major diagnostic centers. However, routine blood and bile specimens collected at autopsy as dried spots on filter paper along with frozen sections of liver and skin biopsy specimens can be submitted to such a reference laboratory for the required mass spectrometry analysis.

Benefits of Metabolic Autopsy
1. A medically plausible cause of death when an infant dies unexpectedly
2. Family opportunity to prevent morbidity and mortality in siblings
3. Opportunity for genetic counseling and prenatal diagnosis
4. Rule out child abuse or neglect

Clinical Implications Causes of death are categorized as natural and unnatural.

1. Natural causes
   a. Cardiovascular
      1. Myocardial infarction
      2. Ventricular tachycardias
      3. Fibrillation (cardiac arrhythmias)
      4. Hypertensive cardiovascular disease
      5. Cardiomyopathies
      6. Strenuous activity during extremes of heat or cold weather
      7. Drug use
      8. Anorexia nervosa
   b. Brain
      1. Cerebrovascular accident
      2. Poorly controlled epilepsy or seizure disorders complicated by cardiac arrest related to \( \text{O}_2 \) deprivation
      3. Brain hemorrhage
      4. Primary brain tumors
      5. Aneurysms
      6. Head trauma
   c. Respiratory
      1. Epiglottitis
      2. Pulmonary thrombosis/embolus
      3. Status asthmaticus
4. Aspiration of food/gastric contents/blood
5. Cavernous tuberculosis
6. Premature birth
7. Fulminating pneumonia
8. Chest trauma
d. Gastrointestinal
1. GI bleed
2. Trauma
3. Peritonitis
4. Massive splenic enlargement or rupture
5. Ingested caustic substances
6. Liver or pancreatic diseases
7. Diabetes mellitus in the presence of diabetic coma (diagnosed by elevated glucose in vitreous of eye)
e. Other
1. Tubal pregnancy rupture leading to massive hemorrhage
2. HIV infection
3. Chronic illness in bedridden persons with septic decubitus ulcers
4. Malnutrition
5. Dehydration
6. Environmental causes (eg, Legionnaire's disease, hantavirus)
7. Communicable disease exposure such as influenza and meningitis

2. Unnatural causes
a. Trauma
1. Body wounds
2. Cuts
3. Lacerations
4. Traumatic amputations
5. Self-inflicted and self-defense wounds
6. Asphyxia
7. Motor vehicle/cycle accidents
8. Airplane crashes
b. Other
1. Sudden infant death syndrome (SIDS), which is the unexpected death of an apparently healthy infant. Postmortem examination may not reveal the cause of death. Neonaticide refers to the deliberate killing of an infant within 24 hours of birth; infanticide is murder of a child.
2. Fire or smoke inhalation
3. Drowning
4. Electrocution
5. Hyperthermia (heat)
6. Hypothermia (cold)
7. Embolism
8. Homicide may be associated with rape, criminal abortion, drug overdose, drug abuse, and drug-related or alcohol-related deaths.

**Interventions**

1. Family preparation for autopsy
   a. Explain rationale for postdeath procedures. Concern and respect for the deceased and significant others can reduce anxiety and objections to or misinterpretations of after-death testing. Obtain a signed, witnessed consent form for autopsy. (Consent is not required for coroner/medical examiner investigative autopsies.)
   b. Consider cultural habits and practices. Human responses and practices surrounding the death of a loved one vary among societies, religions, cultures, and races. In this light, postmortem examination may be offensive to some groups. Use elders, clergy, and social workers to assist in explanations.
   c. Assure the family that nothing will be done without their permission except where required by law.
   d. If fear of mutilation and delay in release of body for burial are concerns, provide clear and concise information to help with decision making. In the case of religious dilemmas, facilitate counsel and communication between clergy and other appropriate individuals or agencies.
   e. Conflict can occur when statutory authority is at odds with family wishes. Explanations and empathic dialogue are essential.

2. Family aftercare after autopsy
   a. Interpret postmortem test results and counsel families appropriately about organs procured for donation (as appropriate).
   b. Indicate need for follow-up if genetic or familial pathologic findings are discovered (eg, young adult with acute myocardial infarction, cardiomyopathy, SIDS, hereditary diseases).
   c. Autopsies are frequently mandatory procedures, especially in sudden, suspicious, or unexplained deaths.
   d. Investigation of accident- and work-related deaths serve the following functions:
      1. Serve as quality-control indicators to confirm predeath diagnoses and to assess effectiveness of drug therapy, diagnostic procedures, surgical techniques, gene therapy, and other diagnostic and treatment modalities
      2. Identify, track, and monitor disease prevalence, incidence, trends, or association with certain lifestyle, environmental, or occupational or safety influences (Occupational Safety and Health Administration [OSHA])
   3. Information gathered from autopsy findings provides a framework for developing better and more sophisticated treatments for disease and illness control or eradication.
   3. Consent from family is required unless autopsy is ordered by the coroner or medical examiner.

4. Should family members be undecided regarding autopsy, they wish to consider it in the following circumstances:
   a. When no firm medical diagnosis has been established
b. If there are questions about an unexpected or mysterious death due to apparent natural causes

c. If there are hereditary, genetic, or contagious diseases

d. When the cause of death could affect insurance settlements and other legal matters

e. When death occurs in the presence of unexpected medical or obstetric complications

f. During the use of experimental drug therapies

g. If death is a result of certain dental, invasive, surgical, or diagnostic procedures

h. When the death does not come under the jurisdiction of the medical examiner, hospital death with inconclusive medical diagnosis before death, Alzheimer's death, brain studies

Organs for Donation

Organs and tissues donated by the deceased or the deceased’s family for transplant are procured before, during, or immediately following autopsy. A special consent form must be signed by a responsible adult and witnessed by a professional. Life-saving organs (eg, kidneys, lungs, heart, pancreas, liver, intestines) are harvested before autopsy while the deceased is still on “life support” following declaration of “brain death.” Other tissues and organs are harvested simultaneously with autopsy procedures or after autopsy: eyes, bones, connective tissues, joints, ligaments, tendons, cartilage, heart valves, and veins. A request for organ donation can be made in a hospital or to a medical examiner (in most states). The request and donation report are documented in the deceased person’s medical record or coroner/medical examiner’s report.

Indications for Organ Donor Testing Before Transplantation

1. Determine exposure to infectious disease (eg, viral hepatitis B and C, possibly TT virus in the future, HIV-1, HIV-2, human T-cell lymphotrophic virus [HTLV-I and -II], cytomegalovirus [CMV], syphilis, other sexually transmitted diseases) to exclude the organ.

2. Match donor and recipient.

3. Establish blood type compatibility, ABO and Rh, for all transplant donors and recipients.

4. Test postmortem specimens for hepatitis B and HIV.

5. Cadaver donor blood specimens are usually obtained before death, except in post-mortem specimens for corneal transplantation, which may not be procured until days after death.

Incidence and Frequency of Blood Group and Rh Type

<table>
<thead>
<tr>
<th>Blood Group and Rh Type</th>
<th>Incidence Frequency of Occurrence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>O positive</td>
<td>1 of 3</td>
</tr>
<tr>
<td>O negative</td>
<td>1 of 15</td>
</tr>
<tr>
<td>A positive</td>
<td>1 of 3</td>
</tr>
<tr>
<td>A negative</td>
<td>1 of 16</td>
</tr>
<tr>
<td>B positive</td>
<td>1 of 12</td>
</tr>
<tr>
<td>B negative</td>
<td>1 of 67</td>
</tr>
<tr>
<td>AB positive</td>
<td>1 of 29</td>
</tr>
<tr>
<td>AB negative</td>
<td>1 of 167</td>
</tr>
</tbody>
</table>

Procedure

1. Observe standard precautions.

2. Obtain common blood and urine samples for initial testing and evaluation of the potential organ donor.

Clinical Implications

1. Victims of poisoning may be organ donors, especially in cases determined by brain death status.

2. Examples of toxins found in organs to be transplanted are included in Table 16.1.

Table 16.1 Toxins Found in Transplanted Organs

<table>
<thead>
<tr>
<th>Toxin</th>
<th>Organs Transplanted</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetaminophen</td>
<td>Heart, cornea, kidney</td>
</tr>
<tr>
<td>Carbon monoxide</td>
<td>Heart, liver, kidney, pancreas</td>
</tr>
<tr>
<td>Barbiturate</td>
<td>Liver, heart, kidney</td>
</tr>
<tr>
<td>Ethanol</td>
<td>Kidney</td>
</tr>
<tr>
<td>Cocaine</td>
<td>Liver, kidney</td>
</tr>
<tr>
<td>Cyanide</td>
<td>Cornea, skin, bone, heart, liver, kidney, pancreas</td>
</tr>
<tr>
<td>Methaqualone</td>
<td>Liver</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>Kidney, heart, liver</td>
</tr>
<tr>
<td>Tricyclic antidepressants</td>
<td>Kidney</td>
</tr>
<tr>
<td>Methanol</td>
<td>Kidney</td>
</tr>
<tr>
<td>Insulin</td>
<td>Kidney, heart, pancreas (islet cells)</td>
</tr>
</tbody>
</table>

3. Guidelines published for optimal time to transplantation from a poisoned organ donor for certain selected toxins are included in Table 16.2.

Table 16.2 Organ Transplantation Guidelines From a Poisoned Organ Donor
Toxin | Optimal Time for Transplant
---|---
Ethylene glycol, methanol, ethanol (ETOH) | Acidosis corrected and plasma/serum ethylene glycol <1.5 mg/L (<24 µmol/L)
Blood methanol <1.5 µg/mL (<47 µmol/L)
Plasma/serum ethanol <20 mg/dL (<0.34 mmol/L) | Shock corrected and serum cyanide level <0.2 mg/mL (<4.6 µmol/L)

**Interventions**

**Pretest Preparation**

1. Discuss potential tissue and organ donation with grieving family members.
2. Explain the rationale for autopsy, testing, procurement, and mortician procedures after death and for donation of tissue or organ. Concern, sensitivity, and respect for the deceased and significant others can reduce anxiety and objections to or misinterpretation of after-death testing. Organ donation may be offensive to some people.
3. Obtain a signed, witnessed consent form for the autopsy and organ/tissue donation.
4. Determine suitability for eye, organ, and tissue donation. Criteria include communicable diseases (HIV, hepatitis, infection, age).
5. Be sensitive. Allow family members to have as much time as needed to be present with their loved one.
6. Obtain a signed, informed consent form for organ/tissue donation. (Consent not required for coroner/medical examiner investigation autopsies.)

**Intratest Care**

1. Prepare the body for tissue donation after consent has been given by applying saline drops to the eyes, elevating the head, paper-taping the eyes closed, placing eye pads over the brow, and placing the body in a refrigerated morgue when possible.
2. Type human leukocyte antigen (HLA) on lymphocytes and determine compatibility before kidney and pancreas transplantation. A strong reaction in compatibility testing predicts rapid transplant rejection and is a contraindication.

**Posttest Aftercare for Organ Donation** Counsel family about organ and tissue procured for donor transplantation.

**BIBLIOGRAPHY**

- American Sleep Disorders Association: The International Classifications of Sleep Disorders, Rochester, New York, ASDA, 1998
- Fritzel J: Cerebral angiography. AJN 98(9): 19–16, September 1998
- Fritzel J: Transesophageal echocardiography. AJN 97(9): 17–18, September 1997
- Fryback DG: Medical care use and costs for adults with sleep apnea. AHRQ grant HS08281, University of Wisconsin, Madison, WI, 7/1/94–12/31/96
Jackson JL, Pettit T: Complications from cardiac catheterization: Analysis of a military population, Milit Med 165, April 2000


Montes P: Managing outpatient cardiac catheterization. AJN 97(8): 34–37, August 1997


The autopsy as an outcome and performance measure. (Online.) Available at www.ahrq.gov/clinic/epcsums/autopsum.htm


Appendix A Standard/Universal Precautions

SAFE Practice

COMMON CATEGORIES OF BODY SUBSTANCES, SECRETIONS, AND FLUIDS

NOTE

STANDARD PRECAUTION GUIDELINES AND PRACTICES FOR SPECIFIC SITUATIONS

Personal Protection Equipment

Disposal of Medical Wastes

Placement of Warning Tags and Signs

General Environmental Cautions

In Case of Exposure to Human Immunodeficiency Virus or Hepatitis B Virus

Handwashing Protocol

The term standard precautions refers to a system designed to reduce the risk of transmission of microorganisms from both recognized and unrecognized sources of infections. Standard precautions direct safe practice and are designed to protect health care workers, patients, and others from exposure to blood-borne pathogens or other potentially infectious materials from any body fluid or unfixed human tissue from any person, living or dead.

Revised guidelines are based upon new information about infectious disease patterns and modes of transmission. The guidelines, designed to be more user friendly, contain two tiers of precautions. Tier one, Standard Precautions, is designed to control nosocomial infections and reduce the risk of transmission of both known and suspected infections. Tier two, used in addition to standard precautions, includes airborne, droplet, and contact precautions to prevent the spread of known or suspected transmissible and virulent pathogens.

SAFE Practice

When handling specimens and performing or assisting with diagnostic procedures, it is important for all health care workers to protect and always take care of themselves first. Presume that all patients have hepatitis B, human immunodeficiency virus (HIV), hepatitis C, or other potential pathogens, and practice standard precautions consistently. Use special care when collecting, handling, packaging, transporting, storing, and receiving specimens. Initial observations and specimen handling are to be performed under a laminar flow hood, and protective clothing, which includes, but is not limited to, gloves, gowns, face masks or shields, and eye protection. These same precautions prevail in the performance of invasive diagnostic procedures. Follow Chapter 1 guidelines for safe, effective, informed pretest, intratest, and posttest care.

COMMON CATEGORIES OF BODY SUBSTANCES, SECRETIONS, AND FLUIDS

1. Blood and blood products
2. Urine
3. Vaginal secretions
4. Saliva
5. Pericardial fluid
6. Peritoneal fluid
7. Pleural fluid
8. Cerebrospinal fluid
9. Gastric fluid
10. Respiratory secretions
11. Semen
12. Synovial fluid
13. Vomitus
14. Wound or ulcer drainage
15. Ascites
16. Amniotic fluid
17. Sweat

NOTE

Standard precautions should also be used when handling amputated limbs and during removal of body parts (surgery, autopsy, or donation).

STANDARD PRECAUTION GUIDELINES AND PRACTICES FOR SPECIFIC SITUATIONS

Personal Protection Equipment

1. Take appropriate barrier precautions when exposure of skin and mucous membranes to blood, blood droplets, or other body fluids is anticipated.
2. Use protective equipment devices to protect eyes, face, head, extremities, air passages, and clothing. This
equipment must always be used during invasive procedures. Ensure proper fit.

Gloves

1. Wear gloves when collecting and handling specimens; touching blood, urine, other body fluids, mucous membranes, or nonintact skin; or performing vascular access procedures or other invasive procedures.
2. Wear gloves when handling items or surfaces soiled with blood, urine, or body fluids.
3. Mandate wearing of gloves when the health care worker's skin is cut, abraded, or chapped during examination of a patient's oropharynx, gastrointestinal or genitourinary tract, nonintact or abraded skin, or active bleeding wounds; and when cleaning specimen containers or engaged in decontaminating procedures.
4. **Possible exceptions to use of gloves:**
   a. When gloves impede palpation of veins for venipuncture (eg, neonates, morbidly obese patients)
   b. In a life-threatening situation where delay could be fatal (wash hands and wear gloves as soon as possible)
5. **Disposable gloves must be changed:**
   a. When moving between patients
   b. When moving from a contaminated to a cleaner site on a patient or on an environmental surface
   c. When gloves are torn or punctured or their barrier function is compromised (do so as soon as feasible)

**Clinical Alert**

Gloves, barrier gowns, aprons, and masks are worn only at the site of use. They are disposed of appropriately at the site of use.

Gowns, Masks, and Eye Protection

1. Wearing of gowns, aprons, and/or fluid impervious lab coats to cover all exposed skin is necessary whenever there is a potential for splashing onto clothing.
2. Gowns or aprons may not be hung and reused.
3. Wear masks correctly situated over nose and chin and tied at the crown of the head and the nape of the neck. Do not hang the mask around the neck. Change the mask when it becomes moist.
4. Wear mask, face shields, and goggles (or prescription glasses with side shields) when contamination of eye, nose, or mouth from fluid is most likely to occur.
5. Shoe covers should be worn in areas where contamination might occur (eg, operating room obstetrics or emergency department). These are disposed of at the site of care.
6. Provide masks, ambu-bags, or other ventilation devices as part of emergency resuscitation equipment kept in strategic locations.

Disposal of Medical Wastes

1. Pour fluids "low and slow" to prevent splash, spray, or aerosol effect.
2. Take precautions to prevent injuries caused by needles, lancets, scalpels, and other sharp instruments and devices during and after procedures and when disposing of used needles. Do not recap needles under normal circumstances.
3. Dispose of all disposable sharp instruments in specially designed, puncture-resistant containers. Do not recap, bend, break by hand, or remove needles from disposable syringes. Use forceps or cut intravenous tubing if necessary. Use care when transferring “sharps” to another person. Use forceps or put the “sharp” in a receptacle.

**Figure A.1** Needle safety device. (1) Attach any brand of needle. (2) Remove cap and draw patient. (3) Press sheath on flat surface. (4) Snap closed and dispose. (Source: MarketLab Inc., Kentwood, Michigan, USA)

4. Place and transport specimens in leakproof receptacles with solid, tight-fitting covers. Cap ports of containers. Before transport, specimens must be placed in a tightly sealed bag marked with a “biohazard” tag. Biohazard symbols warn of biologic hazards and must be displayed in the presence of these hazardous biologic agents or locations.
5. Place soiled linens and similar items in leakproof bags before transport.

Placement of Warning Tags and Signs

1. Properly place warning tags to prevent accidental injury or illness to clinicians who are exposed to equipment or procedures that are hazardous, unexpected, or unusual.
2. Require warning tags to contain a signal word or symbol, such as "Biohazard" or "Biochemical Material," along with the major message, such as "Blood Banking Specimen Inside." All specimens are placed in biohazard bags.

**General Environmental Cautions**

1. Use approved antimicrobial soaps between care of individual patients.
2. Wash hands immediately after removing gloves.
3. Wash hands and other skin surfaces immediately and thoroughly if contaminated with blood or other body fluids.
4. Consider saliva when blood is visible to be potentially infectious, even though it has not been implicated in HIV transmission.
5. Transmission of acquired immunodeficiency syndrome (AIDS) is possible from stool specimens, especially if there is a possibility of blood existing in the stool.
6. Health care workers with open skin lesions or skin conditions should not engage in direct care until the condition clears up or does not present a risk to the patient.
7. Development of an HIV infection during pregnancy may put the fetus at risk for infection.

**In Case of Exposure to Human Immunodeficiency Virus or Hepatitis B Virus**

1. Identify, obtain consent, and test source of exposure immediately for evidence of HIV, HBV, and HCV. If the patient refuses consent, a nonconsenting form must be signed (see Appendix H). If nonconsenting testing is done on the source, the exposed staff member must also have testing.
2. Advise the HIV-negative worker to seek medical evaluation of any acute febrile illness that occurs within 12 weeks after exposure to HIV and be retested at 6 weeks, 12 weeks, and 6 months after exposure.
3. Vaccine is available at no cost to health care workers to prevent hepatitis B infection. There is no vaccine for HIV or hepatitis C.

**Handwashing Protocols**

Unless the situation is a true emergency, hands must always be washed:

1. Before and after care activities that involve direct contact
2. Before surgical or obstetric procedures
3. Before and after endoscopy
4. Before and after invasive procedures
5. Before direct contact with an immunocompromised patient
6. After contact with body fluids or tissues or with soiled equipment, supplies, or surfaces
7. After direct contact with patients in isolation units

**Protocols for First Responders** Examples of protocols for suspected serious infectious disease with signs and symptoms classified as high risk (eg, skin rash or skin involvement and high fever); high fever prodrome, as in possible smallpox (classic lesions); yellow fever (jaundice); and plague (buboes) are stated as follows: when any of these other serious infectious diseases cannot be ruled out and there is uncertain diagnosis or no diagnosis, the patient is classified as high risk. Institute airborne and contact precautions, report immediately, and notify appropriate health department. If rash is present, obtain a dermatology consult and collect specimens by specially trained personnel. Testing is done at the Centers for Disease Control and Prevention (CDC). Some of these diseases may be result of possible bioterrorism (see Chapter 7) and can only be diagnosed by the febrile stage and classic signs. A diagnosis of smallpox (a serious disease that kills 30% of infected people) is based upon tests for variola virus and recognition of the febrile stage, classic smallpox lesions, and lesions in same stage of development.

**Clinical Alert**

1. All first responders and response teams need to be vaccinated before exposure to smallpox, within 3 days to a week after exposure.
2. Anyone directly exposed and those at risk for exposure should be vaccinated.

For additional information, go to the CDC website (www.cdc.gov/smallpox).

**Bibliography**


The rise in incidence of latex allergy may be attributed not only to increased use of latex products in patient care (especially since standard/universal precautions were mandated), but also to the manner in which raw latex was collected and aged. Allergic reactions are caused by latex proteins retained in the finished products, which can show great variations in latex allergen levels. The greatest environmental hazard exposure is produced by latex gloves and the powder from these gloves that becomes airborne.

The U.S. Food and Drug Administration now requires that all medical devices containing natural rubber latex that may directly or indirectly contact the patient display the following statement: “THIS PRODUCT CONTAINS NATURAL RUBBER LATEX.”

As allergy to latex products becomes more prevalent, both in the health care setting and in the general environment, it becomes necessary for agencies to institute specific guidelines and protocols to maximize latex-safe environments for patients and for health care personnel.

Persons at greatest risk for latex or rubber allergy include:

1. Health care workers (an estimated 17% are affected) compared to 1% to 3% of the general population
2. Persons with spina bifida, spinal cord injury, myelodysplasia, or urogenital anomalies (up to 73% are affected)
3. Individuals with a personal or family history of allergies (including hay fever, bee stings, asthma, pet dander, and food or drug allergies)
4. Persons with a history of multiple surgeries
5. Persons with occupational exposure (eg, rubber industry workers; 10% of those handling or manufacturing rubber are affected)
6. Persons with atopic dermatitis or eczema
7. Persons with intraoperative anaphylaxis (for unknown reason)

Increased or continued exposure increases sensitivity to latex allergens and worsens allergic reactions. Patients and health care workers can become sensitized to latex through repeated skin or mucous membrane contact or by inhaling aerosolized glove allergens.

Persons with latex allergies are more likely to react to certain foods that contain cross-reactive proteins, especially bananas, avocados, chestnuts, almonds, kiwi fruit, raw potato, tomato, peach, plum, cherry, melons, celery, apple, pear, and papaya. Latex allergy often begins with a rash on the hands (from gloves). Besides latex allergies, other glove-associated reactions may occur.

### LATEX ALLERGY PRECAUTIONS TO PROTECT THE PATIENT

Strategies and protocols include the following:

1. Identify allergic patients (those with a history of problems related to catheters, tubes, drains, household items, condoms, latex gloves, balloons, toys, and so on); allergy testing (see Chapter 8) may be desirable. Communicate and document data appropriately.

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Signs and Symptoms</th>
<th>Causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Irritant contact dermatitis (nonallergic irritation)</td>
<td>Dry, crusty, hard bumps, sores, and horizontal cracks on skin may manifest as itchy dermatitis on the back of the hands under the gloves.</td>
<td>Handwashing, insufficient rinsing, scrubs, antiseptics, glove occlusion, glove powder Exposure to chemicals used in latex manufacturing, mostly thiurams</td>
</tr>
<tr>
<td>Delayed-type hypersensitivity; allergic contact dermatitis; chemical allergy</td>
<td>Red, raised, palpable area with bumps, sores, and horizontal cracks may extend up the forearm. Occurs after a sensitization period. Appears several hours after glove contact and may persist for many days.</td>
<td></td>
</tr>
</tbody>
</table>
Immediate-type hypersensitivity; latex allergy; protein allergy

Wheal-and-flare response or itchy redness on the skin under the glove. Occurs within minutes; fades away rapidly after removal of the glove. In chronic form may mimic irritant and allergic contact dermatitis. Symptoms can include facial swelling; generalized rashes; nasal, sinus, and eye symptoms; asthma; and respiratory distress. In rare cases, anaphylactic shock may occur and is life-threatening. Generalized hives, bronchospasm, hypotension, extreme facial edema and laryngeal edema, and tachycardia may occur.

Exposure to proteins in latex on glove surface and/or bound to powder and suspended in the air, settled on objects, or transferred by touch

American Nurses Association latex allergy work place information series, Washington, DC, 1996.

2. Sensitive persons should carry autoinjectable epinephrine (Epi-Pen), nonlatex gloves, and emergency medical instructions; should wear a medical alert bracelet; should avoid all forms of latex; and should alert clinicians, family, friends, and employers of the diagnosis and need to avoid latex.

3. Never wear powdered latex gloves when caring for a sensitized patient.

4. Avoid contact of latex with tissue (eg, wounds, mucous membranes, vaginal skin). Practice proper handwashing.

5. Use latex-free products. Examples include:
   a. Gloves
   b. Endotracheal tubes
   c. Suction and wound drainage tubes and reservoir systems
   d. Catheters
   e. Blood pressure cuffs
   f. Stethoscopes
   g. Temperature probe covers, tape, dressings, Ace wraps
   h. Monitoring equipment and supplies (leads, pulse oximeter probes, and cables)
      i. Remove rubber stoppers from vials before withdrawing or reconstituting contents. Rinse syringes with sterile water or saline before use.
      j. Remove latex ports from intravenous tubing and replace with stopcocks or nonlatex plugs. Tape ports shut if no other alternative is available. Replace ports on intravenous therapy bags with nonlatex ports.
   k. Keep resuscitation equipment and emergency supplies and medications readily accessible at all times in the event that anaphylaxis occurs. (Caution: Some resuscitation supplies and equipment may contain latex.)
      l. Instruct the patient about latex-containing supplies, both medical and nonmedical, that could pose problems (see lists).

Medical Supply Items That Frequently Contain Latex

<table>
<thead>
<tr>
<th>Anesthesia equipment/ET tubes, airways</th>
<th>Home and Community Items That Frequently Contain Latex</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bandages/tapes</td>
<td>Appliance Cords</td>
</tr>
<tr>
<td>Bed protectors</td>
<td>Appliques (clothing); Spandex</td>
</tr>
<tr>
<td>Blood pressure tubing/cuffs</td>
<td>Art supplies (paint, glue, rubber bands, erasers, ink)</td>
</tr>
<tr>
<td>Bulb syringes</td>
<td>Balloons/toys/water toys and equipment</td>
</tr>
<tr>
<td>Catheters (many and varied types)</td>
<td>Balls (tennis, Koosh)</td>
</tr>
<tr>
<td>Dressings/elastic wraps</td>
<td>Carpet backing/rubber floors/cushions</td>
</tr>
<tr>
<td>G-tubes/drainas</td>
<td>Condoms/diaphragms</td>
</tr>
<tr>
<td>IV access (Y-sites, tourniquets, adapters, etc.)</td>
<td>Crutch accessories (tips/grips)</td>
</tr>
<tr>
<td>OR masks, hats, shoe covers</td>
<td>Dental braces, chewing gum</td>
</tr>
<tr>
<td>Oxygen masks/cannula/resuscitation devices</td>
<td>Diapers/incontinence products</td>
</tr>
<tr>
<td>Suction equipment</td>
<td>Elastic in socks, underwear, etc.</td>
</tr>
<tr>
<td>Reflex hammers, syringes</td>
<td>Feeding nipples/ pacifiers</td>
</tr>
<tr>
<td></td>
<td>Handles on garden/sporting equipment</td>
</tr>
<tr>
<td></td>
<td>Kitchen gloves</td>
</tr>
<tr>
<td></td>
<td>Tires, hoses</td>
</tr>
</tbody>
</table>

Note: These lists are not all-inclusive. If latex content is unknown, checking with the manufacturer or supplier before use is strongly advised.
NOTE
Assembling and maintaining a cart with latex-free supplies and equipment may be desirable to facilitate safe patient care.

NOTE
If latex-free blood pressure cuffs and stethoscopes are not available, shield the patient's arm with stockinette and apply the cuff over it. Small-diameter (finger-sized) stockinette can be used to cover stethoscope tubing, leads, and so on.

Clinical Alert
Symptoms of anaphylaxis include a dangerous drop in blood pressure, dyspnea, flushed facial appearance, swelling (of throat, tongue, and nose), a feeling of impending doom, and loss of consciousness.

STRATEGIES AND PRECAUTIONS TO REDUCE THE RISK OF LATEX ALLERGIES FOR HEALTH CARE PERSONNEL

Latex sensitivity is a health hazard for health care workers. It is one problem with many causes. Consequently, workplace practices to reduce the incidence of exposure are absolutely necessary to maintain a safe environment for clinicians. Early identification and treatment of latex allergy are important. Allergists who specialize in treating latex allergy may recommend patch testing with glove chemical sensitizers and latex allergy testing by serum or skin prick tests. Blood tests are not as sensitive or as accurate as the skin tests.

Clinical Alert
Currently, no commercially prepared latex skin test extract is available, and office-prepared skin test extract from gloves can vary in latex protein content.

Ways to reduce the risk of latex allergies for health care workers include the following:

1. Use latex-free gloves (powder-free gloves low in protein and chemical allergens) whenever possible and keep exposure to latex at a minimum.
2. Wear gloves that are appropriate for the task; remove gloves at least hourly to air and dry hands.
3. Wash, rinse, and dry hands thoroughly after removal of gloves or between glove changes.
4. Use a pH-balanced soap and avoid cutaneous contact with damaging chemicals.
5. Apply nonsensitizing products (outside of the workplace) to restore the skin's lipid barrier.
6. Wear synthetic gloves or cotton liners with latex work gloves for wet work, if possible.
7. Seek early medical diagnosis to prevent further allergy complications.
8. Advocate for and promote purchase of latex-free products that are of comparable function and quality.
9. Observe all latex allergy precautions that apply to patients. Natural latex is found in many consumer products, such as condoms, balloons, tires, rubber toys, nipples, and pacifiers.

Clinical Alert
Simply using powder-free gloves will not solve the problem.

Clinical Alert
Protocols for management of an allergic reaction:

1. Airway maintenance
2. Administration of oxygen
3. Volume expansion (intravenous lactated Ringer's or normal saline solution)
4. Epinephrine IV
5. Steroids (orally or IV)
6. Diphenhydramine (orally or IV)
7. Aminophylline IV

MANDATES AND STRATEGIES FOR EMPLOYEES REGARDING LATEX OR RUBBER ALLERGY

1. Include latex allergy information as part of new-employee orientation and conduct in-service education training on this subject.
2. Occupational Safety and Health Administration “Right To Know” laws require employers to inform employees of potentially dangerous substances in the workplace on an annual basis.
3. Make available current latex allergy information in newsletters; latex allergy should be on the agenda of risk management committees.
4. Make alternative products available.
5. Establish protocols and procedures related to latex allergy to ensure a safe practice environment.
6. Protect latex-allergic workers from being required to work in latex-contaminated areas.

NOTE

In March 1999, the U.S. House of Representatives conducted a hearing to examine latex allergy recommendations of OSHA, CDC, and the FDA (Food and Drug Administration).
Appendix C Sedation and Analgesia Precautions

INTERVENTIONS FOR ADULT PATIENTS RECEIVING MODERATE SEDATION AND ANALGESIA

Preadministration Patient Preparation

1. Explain the purpose of moderate sedation/analgesia before administering the medications. It is most commonly used for these diagnostic procedures: biopsies, bronchoscopy, ERCP, colonoscopy, gastroscopy, angiogram, cardiac catheterization, EP studies, and cystoscopy. Medications may be administered intravenously or by mouth.

2. Assess the patient’s health status, history of chronic or acute conditions, drug allergies, current medications and potential drug interactions, previous diagnostic test results, level of understanding, orientation, mental status, and ability to cooperate with the procedure. Screen and identify patients who are at high risk for development of complications: the very young, the very old, and those with heart, lung, liver, kidney, or CNS disease, marked obesity, sleep apnea, pregnancy, or drug or alcohol abuse. Patients presenting for moderate sedation/analgesia should undergo a focused physical examination including vital signs, auscultation of the heart, lungs, and evaluation of the airway.

3. Explain the process and procedure and what the patient may experience (feel sleepy, relaxed, no anxiety). Use a calm, caring manner. Controversy exists about fasting time frames; but there is an agreement that preprocedure fasting decreases risks during moderate sedation. Check your agency policy. For adults, no food or liquid (for 2 to 6 hours) to allow for gastric emptying. For infants under 6 months, fast 4 to 6 hours (this includes milk, formula, and breast milk); clear liquids, 2 hours.

4. Before beginning the procedure, establish an intravenous line and keep it open with the ordered intravenous solution. Monitor patency of the line.

5. Monitor pulmonary ventilation (exhaled carbon dioxide) and apply pulse oximeter sensor (oxygenation) especially if the patient is unable to be observed during moderate sedation. Monitor electrocardiogram, pulse oximetry, and patient’s vital signs should be documented (pre-, intra-, and post-procedure).

6. Provide a safe and caring environment. A designated individual, other than the practitioner performing the procedure, should be present to monitor the patient throughout the procedure. In anticipation of emergency situations, have resuscitation equipment and supplies of appropriate size readily available (oxygen therapy, IV fluid, reversal agents, and vasopressors).

7. Follow Chapter 1 guidelines for safe, effective, informed pretest care.

Intratest Patient Care

1. Continuously assess pain or discomfort and sedation levels at frequent established intervals.

2. Administer sedation and analgesics as ordered, often in incremental doses. In pediatrics, oral Versed, ketamine, or chloral hydrate may be administered in chocolate syrup.

3. Recognize physiologic effects of agents used for moderate sedation. These medications include the following, among others:
   a. Meperidine hydrochloride (Demerol)
b. Diazepam hydrochloride (Valium)
c. Midazolam hydrochloride (Versed)
d. Lorazepam (Ativan)
e. Droperidol (Inapsine) (Check with your pharmacy or institutional policy regarding use of this drug.)
f. Fentanyl citrate (Sublimaze)
g. Morphine sulfate

4. Monitor the intravenous site for infiltration and the general effects of the medication as well as local analgesia site. Local anesthesia and sedation may cause adverse reactions.
5. Assess level of consciousness—responses of patients to commands during the procedure serves as a guide to their level of consciousness. If reflex withdrawal from painful stimulation is the only response, the patient is likely to be deeply sedated, approaching the state of general anesthesia.
6. Monitor pulmonary ventilation by auscultation of breath and observation of spontaneous respiration. Automated apnea monitoring (detection of exhaled CO\textsubscript{2}) may be used, but is not a substitute for monitoring ventilatory function.
7. Be aware that detecting changes in heart rate and blood pressure for hemodynamics reduces risk of CV collapse and hypopnea.
8. Use oximetry to detect hypoxemia and decrease adverse outcomes such as cardiac arrest and death.
9. Anticipate and monitor for potential complications. Arrhythmias should be promptly reported and treated if necessary. Many of these medications are respiratory depressants, mandating frequent respiratory assessments. If oxygen saturation drops below acceptable levels (=90%), sedation may need to be held or reversed. Have intravenous reversal agents such as naloxone (Narcan) and flumazenil (Romazicon) readily available. Supplemental oxygen therapy may be necessary until oxygen saturation levels, vital signs, neurologic response, and cardiac rhythms are at acceptable levels.
10. Respond to emergencies rapidly and appropriately during administration of, or recovery from, moderate sedation and analgesia.
11. Document carefully and completely all observations, including medications and dosages. Record unexpected outcomes and follow-up care.
12. Follow Chapter 1 guidelines regarding safe, effective, informed intratest care.

Postadministration Patient Aftercare

1. Monitor vital signs, electrocardiogram, pulse oximetry, ventilation, neurologic signs, level of consciousness, and patient response to verbal commands according to established guidelines.
2. Monitor the patient after the procedure until the patient is stable and reactive to preprocedure levels.
3. Provide both verbal and written posttest instructions. Moderate sedation may not completely wear off for several hours. Patients should not:
   a. Drive or operate power machinery or tools for at least 24 hours.
   b. Consume alcoholic beverages or make legal decisions for 24 hours.
   c. Smoke—if the patient is a smoker, emphasize the risks of smoking in the postsedation state (ie, falling asleep).
   d. Take tranquilizers, pain medications, or other medications that may interact with drugs used for sedation without first contacting the physician.
4. Provide instructions for posttest care and the need for contacting the physician if any unexpected outcomes should occur.
5. Evaluate the patient for readiness for discharge. Patients should be alert and oriented, or if altered mental status was initially present, should have returned to baseline. Vital signs should be stable and within acceptable limits. Provide a safe transport or discharge in the presence of a responsible adult.
6. Allow sufficient time (up to 2 hours) to elapse following the last administration of reversal agents to ensure that patients do not become re-sedated after reversal effects have worn off.
7. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

Clinical Alert

Record ventilatory/oxygen status and hemodynamics before the procedure begins, after administration of sedative and analgesia, upon completion of procedure, during initial recovery, and at time of discharge.

The primary drugs used for moderate sedation and analgesia are benzodiazepams and opiates, which are CNS depressants. Opiates are also used for sedation, as are some tranquilizers (droperidol [Inapsine]); for pain relief, fentanyl (Sublimaze), morphine, and meperidine are used. (See Chart C.1). Combinations of drugs may be more effective than single agents in some instances. Agents must then be appropriately reduced, and there is a greater need to monitor respiratory function. Intravenous sedative and analgesic drugs are to be given in small incremental doses. When drug is administered orally, rectally, intranasally, intramuscularly, or subcutaneously, allow time for drug absorption before giving another dose.
<table>
<thead>
<tr>
<th><strong>Chart C.1 Examples of Sedatives and Analgesic Drugs Used</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Diazepam (Valium)</strong></td>
</tr>
<tr>
<td>CNS depressant, amnesic, lacks analgesia. Duration: 2–8 h.</td>
</tr>
<tr>
<td><strong>Dosage guidelines:</strong> 2–5 mg to maximum of 10 mg. No more than 5 mg/min. Give additional doses in 2.5 mg increments. Wait 3 min before redosing. Onset: 5–10 min; IV, 15–30 min. Reduce dose by 1/3 when an opiate is being used concomitantly.</td>
</tr>
<tr>
<td><strong>Precautions:</strong> Increased effects if taking CNS depressants, alcohol, cimetidine, or disulfiram. Avoid in patients with renal disease.</td>
</tr>
</tbody>
</table>

| **Midazolam (Versed)** |
| CNS depressant; three to four times as potent as diazepam. Provides sedation, amnesia, and decreases anxiety. Lacks analgesia. Duration: 60–90 min. |
| **IV dosage guidelines:** Initial dose: 0.1–0.2 mg/kg. 0.5–1.0 mg given slowly over at least 2–3 min; not to exceed 4 mg. Wait at least 2 minutes before redosing. Give in small increments after initial dose. Onset: 1–2 min (IV), 10–20 min (IM), 10–15 min (intranasally), 10–30 min (orally), and 10–30 min (rectally). Decrease dose if given with narcotics (by 25%–30% in healthy adult, by 55%–60% in elderly or debilitated). Duration: 30–60 min (IV), 1–2 h (IM), 45–60 min (intranasally), and 40–90 min (orally or rectally). |
| **Precautions:** Watch for respiratory depression, especially in children. Contraindicated in patients with narrow-angle glaucoma. |

| **Lorazepam (Ativan)** |
| CNS depressant; lacks analgesia. Duration: 4–6 h. |
| **Dosage guidelines:** 0.5–2 mg IV, given slowly; maximum 4 mg. Onset: 5–10 min. Decrease dose in elderly. |
| **Precautions:** Increased effects with MAOIs, barbiturates, narcotics, hypnotics, tricyclic antidepressants, alcohol; decreased effects with oral anticoagulants and heparin. Use with great caution in children. Owing to anticholinergic actions, use with caution in patients with asthma, narrow-angle glaucoma, prostatic hypertrophy, or bladder neck outlet obstruction. |

| **Meperidine (Demerol)** |
| Opiate narcotic analgesic sedative; 60–80 mg meperidine = 10 mg morphine. Duration: 2–4 h. |
| **IV dosage guidelines:** 10 mg IV. Give increments slowly; 25–50 mg IV over 2 min; repeat at 5-min intervals; 10–15 mg maximum; 150 mg over total period of procedure. Decrease dose in elderly or debilitated; use caution with renal disease. |
| **Precautions:** Contraindicated if patient has had a MAOI in last 14 days; may precipitate severe and irreversible reaction and death; decrease dose if given with other narcotic, barbiturate, tranquilizer, tricyclic antidepressant, or sedative. Use with caution in patients with supraventricular tachycardia; may cause increased ventricular response. |

| **Droperidol (Inapsine)** |
| Major tranquilizer; no analgesic properties. Produces cognitive dissociation—a sense of detachment; antiemetic. Duration: Varies over several hours. |
| **IV dosage guidelines:** 0.625–1.25 mg; decrease dose in elderly. Onset: 3–10 min. Peak action: 30 min. |
| **Precautions:** Potentiates narcotics and other CNS depressants. Produces mild alpha-adrenergic block. |

| **Fentanyl (Sublimaze)** |
| Opiate narcotic analgesic sedative; much more powerful than morphine. Duration 30–60 min. |
| **IV dosage guidelines:** 1.0–2.0 mcg/kg. 25–50 mg IV at 5-min. intervals, titrating time to patient response. May also be given by transdermal patch, or as a “lollipop” for children. Onset of sedation: 1–2 min; onset of analgesia: may not be noted for several minutes. Maximum dosage: 500 mg/5 h. |
| **Precautions:** Potentiates narcotics and other CNS and respiratory depressants. Produces mild adrenergic block. Reversal with Narcan: Rapid IV administration may cause chest wall rigidity; treat with chemical paralytics, intubation, and ventilatory support. |

| **Morphine** |
Reversal with Narcan: Rapid IV administration may cause chest wall rigidity; treat with chemical paralytics, intubation, and ventilatory support.

**Morphine**

Opiate narcotic analgesic. Duration: 1–3 h.

**IV dosage guidelines:** 2–15 mg IV over a 5-min period; maximum of 20 mg over total procedure time.

**Precautions:** Causes analgesia and respiratory depression; check respiratory status.

**Ketamine (Ketalar)**

A PCP derivative used in pediatrics: analgesic, sedative, and amnestic. Duration: 10–30 min (IV), 60–90 min (IM).

**Dosage guidelines:** Rapid onset with both IV and IM (1 min IV and 5–6 min IM); longer if given orally or rectally.

**Precautions:** Causes copious saliva production and airway secretions (treated with atropine or Robinul). Does not cause respiratory depression. Associated with nightmares (rare in children) and not with oral or rectal routes.

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**Clinical Alert**

Ketamine is known as a “club drug” that is used unknowingly by the victim, who is later sexually assaulted.

**Sufentanil (Sufenta)**

Analgesic more potent than fentanyl; used in pediatrics. Duration: 1–2 h.

**Dosage guidelines:** Onset: 5–15 min, intranasally; may be administered with Versed.

**Precautions:** Reversal with Narcan. Precautions same as for fentanyl (Sublimaze).

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**Bibliography**


Normal or reference values for laboratory results are reported in conventional units, SI units, or both. The SI system uses seven dimensionally independent units of measurement to provide logical and consistent measurements. For example, SI concentrations are written as amount per volume (moles or millimoles per liter) rather than as mass per volume (grams, milligrams, or millequivalents per deciliter, 100 milliliters, or liter). Universal values may differ between systems or may be the same. For example, chloride is the same in both systems: 95–105 mEq/L (conventional units) and 95–105 mmol/L (SI units). Clinical laboratory data may be reported in either conventional units or SI units, or both. Examples of conversion of data from the two systems are included (Table D.1). To convert SI units to conventional U.S. units, divide by the factor; to convert conventional U.S. units to SI units, multiply by the factor.

Table D.1 Examples of Conversions to Systéme International (SI) Units

<table>
<thead>
<tr>
<th>Component</th>
<th>System</th>
<th>Present Reference Intervals</th>
<th>Present Unit</th>
<th>Conversion Factor</th>
<th>SI Reference Intervals</th>
<th>SI Unit Symbol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alanine aminotransferase (ALT)</td>
<td>Serum</td>
<td>5–40</td>
<td>U/L</td>
<td>1.00</td>
<td>5–40</td>
<td>U/L</td>
</tr>
<tr>
<td>Albumin</td>
<td>Serum</td>
<td>3.9–5.0</td>
<td>g/dL</td>
<td>10</td>
<td>39–50</td>
<td>g/L</td>
</tr>
<tr>
<td>Alkaline phosphatase</td>
<td>Serum</td>
<td>35–110</td>
<td>U/L</td>
<td>0.01667</td>
<td>0.6–1.8</td>
<td>µkat/L</td>
</tr>
<tr>
<td>Aspartate aminotransferase (AST)</td>
<td>Serum</td>
<td>5–40</td>
<td>U/L</td>
<td>0.01667</td>
<td>0.08–0.67</td>
<td>µkat/L</td>
</tr>
<tr>
<td>Bilirubin</td>
<td>Serum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Direct</td>
<td>mg/dL</td>
<td>0–0.2</td>
<td></td>
<td>17.10</td>
<td>0–4</td>
<td>µmol/L</td>
</tr>
<tr>
<td>Total</td>
<td>mg/dL</td>
<td>0.1–1.2</td>
<td></td>
<td>17.10</td>
<td>2–20</td>
<td>µmol/L</td>
</tr>
<tr>
<td>Calcium</td>
<td>Serum</td>
<td>8.6–10.3</td>
<td>mg/dL</td>
<td>0.2495</td>
<td>2.15–2.57</td>
<td>mmol/L</td>
</tr>
<tr>
<td>Carbon dioxide, total</td>
<td>Serum</td>
<td>22–30</td>
<td>mEq/L</td>
<td>1.00</td>
<td>22–30</td>
<td>mmol/L</td>
</tr>
<tr>
<td>Chloride</td>
<td>Serum</td>
<td>98–108</td>
<td>mEq/L</td>
<td>1.00</td>
<td>98–108</td>
<td>mmol/L</td>
</tr>
<tr>
<td>Cholesterol</td>
<td>Serum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age &lt;29 yr</td>
<td>mg/dL</td>
<td>&lt;200</td>
<td></td>
<td>0.02586</td>
<td>&lt;5.15</td>
<td>mmol/L</td>
</tr>
<tr>
<td>30–39 yr</td>
<td>mg/dL</td>
<td>&lt;225</td>
<td></td>
<td>0.02586</td>
<td>&lt;5.80</td>
<td>mmol/L</td>
</tr>
<tr>
<td>40–49 yr</td>
<td>mg/dL</td>
<td>&lt;245</td>
<td></td>
<td>0.02586</td>
<td>&lt;6.35</td>
<td>mmol/L</td>
</tr>
<tr>
<td>&gt;50 yr</td>
<td>mg/dL</td>
<td>&lt;265</td>
<td></td>
<td>0.02586</td>
<td>&lt;6.85</td>
<td>mmol/L</td>
</tr>
<tr>
<td>Complete blood count</td>
<td>Blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hematocrit</td>
<td>Blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Men</td>
<td>%</td>
<td>42–52</td>
<td>0.01</td>
<td>0.42–0.52</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Women</td>
<td>%</td>
<td>37–47</td>
<td>0.01</td>
<td>0.37–0.47</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Red cell count</td>
<td>Blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Men</td>
<td>/mm$^3$</td>
<td>4.6–6.2 × 10$^6$</td>
<td>10$^6$</td>
<td>4.6–6.2 × 10$^{12}$/L</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Women</td>
<td>/mm$^3$</td>
<td>4.2–5.4 × 10$^6$</td>
<td>10$^6$</td>
<td>4.2–5.4 × 10$^{12}$/L</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White cell count</td>
<td>/mm$^3$</td>
<td>4.5–11.0 × 10$^3$</td>
<td>10$^6$</td>
<td>4.5–11.0 × 10$^{9}$/L</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Platelet count</td>
<td>/mm$^3$</td>
<td>150–300 × 10$^3$</td>
<td>10$^6$</td>
<td>150–300 × 10$^{9}$/L</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cortisol</td>
<td>Serum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8 AM</td>
<td>µ/dL</td>
<td>5–25</td>
<td>27.59</td>
<td>140–690</td>
<td>nmol/L</td>
<td></td>
</tr>
<tr>
<td>8 PM</td>
<td>µg/dL</td>
<td>3–13</td>
<td>27.59</td>
<td>80–360</td>
<td>nmol/L</td>
<td></td>
</tr>
<tr>
<td>Creatinine kinase</td>
<td>Serum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High CK group (black men)</td>
<td>U/L</td>
<td>50–250</td>
<td>1.00</td>
<td>50–520</td>
<td>U/L</td>
<td></td>
</tr>
<tr>
<td>Intermediate CK group (nonblack men, black women)</td>
<td>U/L</td>
<td>35–345</td>
<td>1.00</td>
<td>35–345</td>
<td>U/L</td>
<td></td>
</tr>
<tr>
<td>Low CK group (nonblack women)</td>
<td>U/L</td>
<td>25–145</td>
<td>1.00</td>
<td>25–145</td>
<td>U/L</td>
<td></td>
</tr>
<tr>
<td>Creatinine kinase isoenzyme, MB fraction</td>
<td>%</td>
<td>&gt;5</td>
<td>0.01</td>
<td>&gt;0.05</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Creatinine</td>
<td>mg/dL</td>
<td>0.4–1.3</td>
<td>88.40</td>
<td>35–115</td>
<td>µmol/L</td>
<td></td>
</tr>
</tbody>
</table>
### Men
- DIGOXIN, THERAPEUTIC: Serum, 0.4–1.1 mg/dL (35–97 µmol/L)
- GLUCOSE: Serum, 62–115 mg/dL (3.4–6.1 mmol/L)
- IRON-BINDING CAPACITY: Serum, 230–410 µg/dL (41–73 µmol/L)
- MAGNESIUM: Serum, 1.2–1.9 mEq/L (0.50–0.78 mmol/L)
- PHOSPHATE (PHOSPHORUS, INORGANIC): Serum, 2.3–4.1 mg/dL (0.75–1.35 mmol/L)
- POTASSIUM: Serum, 3.7–5.1 mEq/L (1.00–3.7–5.1 mmol/L)
- UROCALCIC ACID (URATE): Serum, 6.5–8.3 g/dL (0.5–8.3 g/L)
- SODIUM: Serum, 134–142 mEq/L (134–142 mmol/L)
- THEOPHYLLINE, THERAPEUTIC: Serum, 5.0–5.25 mg/dL (5.25–11.6 µmol/L)
- THYROID-STIMULATING HORMONE (TSH): Serum, 0.5–2.0 mIU/mL (0.6–2.6 nmol/L)
- TRIGLYCERIDES: Serum, 50–200 mg/dL (0.55–1.85 mmol/L)
- UREA NITROGEN: Serum, 6–25 mg/dL (2.1–8.9 mmol/L)
- VITAMIN B₁₂: Serum, 250–1000 pg/mL (0.7378–180–700 pmol/L)

### Women
- DIGOXIN, THERAPEUTIC: Serum, 0.7–1.3 mg/dL (35–97 µmol/L)
- GLUCOSE: Serum, 88.40–140 mg/dL (4.91–7.66 mmol/L)
- IRON-BINDING CAPACITY: Serum, 230–410 µg/dL (41–73 µmol/L)
- MAGNESIUM: Serum, 1.2–1.9 mEq/L (0.50–0.78 mmol/L)
- PHOSPHATE (PHOSPHORUS, INORGANIC): Serum, 2.3–4.1 mg/dL (0.75–1.35 mmol/L)
- POTASSIUM: Serum, 3.7–5.1 mEq/L (1.00–3.7–5.1 mmol/L)
- UROCALCIC ACID (URATE): Serum, 6.5–8.3 g/dL (0.5–8.3 g/L)
- SODIUM: Serum, 134–142 mEq/L (134–142 mmol/L)
- THEOPHYLLINE, THERAPEUTIC: Serum, 5.0–5.25 mg/dL (5.25–11.6 µmol/L)
- THYROID-STIMULATING HORMONE (TSH): Serum, 0.5–2.0 mIU/mL (0.6–2.6 nmol/L)
- TRIGLYCERIDES: Serum, 50–200 mg/dL (0.55–1.85 mmol/L)
- UREA NITROGEN: Serum, 6–25 mg/dL (2.1–8.9 mmol/L)
- VITAMIN B₁₂: Serum, 250–1000 pg/mL (0.7378–180–700 pmol/L)

### Digoxin, therapeutic
- Serum, 0.5–2.0 ng/mL (1.281 ng/mL)
- PLASMA, 0.5–2.0 ng/mL (1.281 ng/mL)

### Erythrocyte indices
- BLOOD, MEAN CORPUSCULAR VOLUME (MCV): 80–100 microns
- BLOOD, MEAN CORPUSCULAR HEMOGLOBIN (MCH): 27–31 pg
- BLOOD, MEAN CORPUSCULAR HEMOGLOBIN CONCENTRATION (MCHC): 32–36 %

### Ferritin
- SERUM, MEN: 29–438 ng/mL (29–438 µg/L)
- SERUM, WOMEN: 9–219 ng/mL (9–219 µg/L)

### Folate
- SERUM: 2.5–20.0 ng/mL (6–46 nmol/L)

### Follicle-stimulating hormone (FSH)
- SERUM, CHILDREN: 12 or < mIU/mL (12 or < IU/L)
- SERUM, MEN: 2.0–10.0 mIU/mL (2.0–10.0 IU/L)
- SERUM, WOMEN, FOLLICULAR: 3.2–9.0 mIU/mL (3.2–9.0 IU/L)
- SERUM, WOMEN, MIDCYCLE: 3.2–9.0 mIU/mL (3.2–9.0 IU/L)
- SERUM, WOMEN, LUTEAL: 2.0–6.2 mIU/mL (2.0–6.2 IU/L)

### Gases, arterial
- BLOOD, PO₂: 80–95 mm Hg (10.7–12.7 kPa)
- BLOOD, PCO₂: 37–43 mm Hg (4.9–5.7 kPa)

### Glucose
- SERUM: 62–110 mg/dL (3.4–6.1 mmol/L)

### Iron
- SERUM: 50–160 µg/dL (9–29 µmol/L)

### Iron-binding capacity
- SERUM: 230–410 µg/dL (41–73 µmol/L)

### Lactic dehydrogenase
- SERUM: 120–300 U/L (120–300 U/L)

### Luteinizing hormone
- SERUM, MEN: 4.9–15.0 mIU/mL (4.9–15.0 IU/L)
- SERUM, WOMEN, FOLLICULAR: 5.0–25 mIU/mL (5.0–25 IU/L)
- SERUM, WOMEN, LUTEAL: 3.1–13 mIU/mL (3.1–13 IU/L)

### Magnesium
- SERUM: 1.2–1.9 mEq/L (0.4114 mEq/L)

### Osmolality
- SERUM: 278–300 mOsm/kg (278–300 mmol/kg)

### Phenobarbital, therapeutic
- SERUM: 15–40 µg/mL (65–175 µmol/L)

### Phenyltoin, therapeutic
- SERUM: 120–300 U/L (120–300 U/L)

### Phosphate (phosphorus, inorganic)
- SERUM: 2.3–4.1 mg/dL (0.75–1.35 mmol/L)

### Procalcitonin
- SERUM: 0.2–0.8 ng/mL (0.3–1.4 pmol/L)

### Protein, total
- SERUM: 6.5–8.3 g/dL (65–83 g/L)

### Sodium
- SERUM: 134–142 mEq/L (134–142 mmol/L)

### Theophylline, therapeutic
- SERUM: 5.0–5.25 mg/dL (5.25–11.6 µmol/L)

### Thyroid-stimulating hormone (TSH)
- SERUM: 0.5–2.0 mIU/mL (0.6–2.6 nmol/L)

### Thyroxine
- SERUM: 4.5–13.2 µg/dL (12.87–58–170 nmol/L)

### T₃-uptake ratio
- SERUM: 0.88–1.19

### Triiodothyronine (T₃)
- SERUM: 70–235 ng/mL (1.1–3.6 nmol/L)

### Triglycerides
- SERUM: 50–200 mg/dL (0.55–2.25 mmol/L)

### Uric acid
- SERUM: 2.9–8.5 mg/dL (170–510 µmol/L)

### Urea nitrogen
- SERUM: 6–25 mg/dL (2.1–8.9 mmol/L)

### Vitamin B₁₂
- SERUM: 250–1000 pg/mL (0.7378–180–740 pmol/L)

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**Example:**

To convert a digoxin level of 0.6 nmol/L (SI units), **divide** by the factor 1.281 to obtain conventional units of 0.5 ng/dL.

To convert a Ca **²** value of 8.6 mg/dL (conventional units), **multiply** by the factor 0.2495 to obtain the SI units of 2.15 mmol/L.

Appendix E Guidelines for Specimen Transport and Storage

Clinical Alert

Routines for collection and handling of specimens and reporting of specific patient information vary depending on agency protocols, the clinical setting, and specialty laboratory requirements. The primary objectives in the transport of diagnostic specimens are to maintain the sample as near to its original state as possible with minimum deterioration and to minimize hazards to specimen handlers. Specimens should be collected and transported as quickly as possible (a 2-hour time limit is recommended). For urine transport, a small amount of boric acid may be used; a holding or transport medium can be used for most other specimen types. Follow carefully the instructions for handling and transport of specimens provided on the kit or by the manufacturer, or by the laboratory that has provided these collection kits.

1. When the patient delivers the specimen directly, provide a biohazard bag and include clearly written directions about the specific handling precautions, storage conditions, and specific directions for locating the physical facility.

2. Specimens may be mailed or transported to specialty laboratories located in other cities or distant areas. To avoid delays in specimen analysis, it is important to follow specific instructions for collection, packaging, labeling, and transporting of specimens. Some specimens must be received in the laboratory within an exact time frame, under specified storage conditions. Regulatory agencies (e.g., the Department of Transportation [DOT], or the International Air Transport Association [IATA]) require training to ensure that samples are properly packaged. The DOT requires training every 3 years, whereas IATA requires every 2 years.

   a. When packaging a specimen for shipping to a specialty laboratory, place the specimen in a securely closed, watertight container (such as a test tube, vial, or other primary container), then enclose the entire primary container in a second, durable, watertight container (secondary container). Each set of primary and secondary containers should then be enclosed in a sturdy, strong outer shipping container (Fig. E.1).

   FIGURE E.1 Proper technique for packaging of biologically hazardous materials. (CDC Laboratory Manual, DHEW publication No. [CDC] 74-8272, Atlanta, Centers for Disease Control, 1974.)

   b. Follow appropriate labeling for etiologic agents and biomedical materials (Fig. E.2 and Fig. E.3). If the package becomes damaged or leaks, the carrier is required, by federal regulations, to isolate the package and notify the Biohazards Control Office, Centers for Disease Control and Prevention, in Atlanta, Georgia. The carrier must also notify the sender that (improper) packaging not meeting regulatory requirements can cause a significant delay in specimen analysis, reporting of results, and medical diagnosis and treatment of the patient’s problem. Examples of specialty laboratory requirements for transporting, packaging, and mailing of specific specimens are shown in the table.

   FIGURE E.2 Etiologic agents logo and “notice to carrier” that must be affixed to the outside of any package containing potentially hazardous and infectious biologic materials. Refer to packaging instructions in the event additional paperwork is required to accompany the package.
Specimen
Blood for trace metals
Observe contamination control in sample collection—for example, most blood tubes are contaminated with trace metals, and all plastic syringes with black rubber seals contain aluminum, varying amounts of zinc, and all heavy metals (lead, mercury, cadmium, nickel, chromium, and others). The trace metal sample should be collected first—once the needle has punctured the rubber stopper, it is contaminated and should not be used for trace metal collection. Use alcohol swabs to cleanse sets; avoid iodine-containing disinfectants, use only stainless steel phlebotomy needles. Blood for serum testing of trace elements should be collected in a royal-blue top (sodium heparin anticoagulant) trace element blood collection tube. After collecting and centrifuging, place in a 5-mL, metal-free, screw-capped polypropylene vial; transfer 5 mL using sterile polypropylene pipette. Cap vial tightly, attach specimen label, and send to lab cool or frozen. All specimens stored >48 hours should be frozen and sent on dry ice. (Keep specimen cool with frozen coolant April–October, refrigerated coolant November–March.)

Blood for photosensitive analysis
Avoid exposure to any type of light (artificial or sunlight) for any length of time. These specimens need aluminum foil wrap or brown plastic vial. Specimens for vitamin A, vitamin B_{12}, ß-carotene, porphyrins, vitamin D, and bilirubin are examples of substances that need to be protected from light.

Routine urinalysis, random, midstream
Preferred transport container is a yellow plastic screw-top tube that contains a tablet that preserves any formed element (crystals, casts, or cells) and prevents alteration of chemical constituents caused by bacterial overgrowth. Pour urine into tube, cap tube securely, and invert to dissolve the tablet.

Urine culture
Use a culture and sensitivity (C&S) transport kit containing a sterile plastic tube and transfer device for collection. This tube contains a special urine maintenance formula that prevents rapid multiplication of the bacteria in the urine. Pour the urine specimen into the tube and seal properly.

Urine for calcium, magnesium, and oxalate
Use acid-washed plastic containers for collection and transport of specimen. If urine pH is >4, the results may be inaccurate. Do not collect urine in metal-based containers such as metal bed pans or urinals.

Stool
Use a special 1000-mL container, such as Nalgene, for total sample collection and a 100-mL white polypropylene container for a portion of a large sample (aliquot) for feces collection. Each container should have a similar label affixed before it is given to the patient. When the container is given to the patient, provide the following instructions: test to be done, specimen requirements, diet requirements, collection and storage of specimen; two 1000-mL Nalgene containers provided for timed collection and one-100 mL container for a random collection specimen; information on how to obtain additional containers if necessary, and do not fill any container more than ¾ full (indicated line on label). At the time that the patient returns the container to the clinic, the health care worker fills in the label with the correct information. If “Other” is checked, enter duration on line on label. If more than one container is sent, be sure to indicate total number sent on the line.

Stool, homogenized
For a homogenized (blended) specimen, the required mailed specimen is a 80-mL portion of homogenized feces. Homogenize and weigh according to laboratory protocol. Pour the homogenate into the container as soon as possible (to avoid settling). On the request form, indicate specimen total weight and amount of water added. Include length of period of collection on request form, also. Send the homogenized specimen at the preferred transport temperature listed in agency specimen requirements protocol. This test is usually performed at a special reference laboratory.

Infectious substance
A biohazard ("Etiologic Agent") label must be affixed (or preprinted on bags) to all patient specimens for transport. Body fluids have been recognized by the Centers for Disease Control and Prevention as being directly linked to the transmission of HIV (AIDS) and hepatitis B virus (HBV). Standard precautions apply to these fluids and include special handling requirements of blood, semen, blood products, vaginal secretions, cerebrospinal fluid, synovial fluid, pleural fluid, peritoneal fluid, pericardial fluid, amniotic fluid, and concentrated HIV and HBV. Also, a "Biohazard" label must be affixed to all microbiology specimens, including anaerobic and aerobic bacteria, mycobacteria, fungi, and yeast. The specimen must be sent on an agar slant tube in a special transport container (a pure culture, actively growing); do not send on culture plates. The outer shipping container of all etiologic agents transported via interstate traffic must be labeled as illustrated in Figure E.2.
Specimens requiring exceptional handling

- Clearly and accurately label each specimen with patient's full name, sex, birth date, unique identification number, time and date of specimen collection, name of practitioner ordering specimen, and signature of person collecting specimens. The test order form and sample should be checked for a match and transported in a single package.
- If a delay of >4 days before specimen examination is expected, freezing of the specimen is preferred. Place the specimen in a plastic vial (not glass); the container should not be more than ¾ full, to allow for expansion when frozen. Store in freezer or on dry ice until specimen is picked up by carrier or transported to the laboratory. Label vial with patient's name, date, type of specimen (e.g., EDTA plasma, serum, urine).

Frozen

- Urine, respiratory exudates, and stool or feces (transport medium is not used) must all be refrigerated before transport. Specimen that must be kept at room temperature (ambient) before inoculation of media include spinal fluids and other body fluids, specimen for Neisseria gonorrhoeae isolation, and blood and wound cultures. Place specimen in the refrigerator for storage before pickup by courier.
- When packaging, place the specimen container in the zip-lock portion of bag and the required coolant in the outer pouch. If dry ice or a refrigerant is used, it must be placed outside the secondary container and the outer shipping container; the shock-absorbant material should be placed so that the secondary container does not become loose inside the outer shipping container as the dry ice evaporates.

Refrigerated (iced or cooled)

- Urine, respiratory exudates, and stool or feces (transport medium is not used) must all be refrigerated before transport. Specimen that must be kept at room temperature (ambient) before inoculation of media include spinal fluids and other body fluids, specimen for Neisseria gonorrhoeae isolation, and blood and wound cultures. Place specimen in the refrigerator for storage before pickup by courier.

Anaerobic

- Aspiration with a needle and syringe, rather than a swab, is the preferred method of collection of a specimen for recovery of anaerobic bacteria; once collected, the specimen must be protected from ambient oxygen and kept from drying until it can be processed in the laboratory. Transport container for anaerobic specimen includes:
  1. Syringe and needle for aspiration—valid only if specimen can be transported without delay. Remove needle cap before transporting. (Organism will survive in the aspirated material.)
  2. Port-a-cult (tube or vial)—tubes are used primarily for insertion of swab specimens; vials are used for inoculation of liquid specimen.
  3. Anaerobic culturette—plastic tube or jacket is fitted with a swab and contains either transport or pre-reduced medium. The culturette system also includes a vial or chamber separated by a membrane that contains chemicals that generate CO$_2$ catalysts and desiccants to get rid of any residual O$_2$ that may get into the system. The ampule must be crushed to activate this system.
  4. Bio-bag or plastic pouch system—transparent plastic bag that contains a CO$_2$-generating system, palladium catalyst cups, and an anaerobic indicator. Bag is sealed after inoculated plates have been inserted and the CO$_2$-generating system is activated. The advantage of this system is that the plates can be directly observed for early growth of colonies.

Clinical Alert

The Code of Federal Regulations governing the shipment of etiologic agents (S72.2 Transportation of Diagnostic Specimens, Biological Products, and Other Materials; Minimum Packaging Requirements) reads as follows:

No person may knowingly transport or cause to be transported in interstate traffic, directly or indirectly, any material, including, but not limited to, diagnostic specimens and biological products which such persons reasonable believe may contain an etiologic agent unless such material is packaged to withstand leakage of contents, shocks, pressure changes, and other conditions incident to ordinary handling in transportation.
Vitamin Testing

Both fat-soluble and water-soluble vitamins play a variety of physiologic roles in the body. Vitamin concentrations in blood, urine, and certain body tissues can be measured and reflect the nutritional status of the patient.

Vitamins have varying modes of action. For instance, vitamin E is an antioxidant, vitamin C is an enzyme cofactor, and vitamin A is an anti-infection agent.

Sources of fat-soluble vitamins include ingested (dietary) substances and biologic or intestinal microorganisms. Fat-soluble vitamins include vitamin A (known as retinol or carotene), vitamin D (calciferol), vitamin E (tocopherol), and vitamin K (consisting of phylloquinones or K₁, menaquinones or K₂, and menadiones or K₃).

The sources of water-soluble vitamins are dietary (ingested) substances and intestinal microorganisms. Water-soluble vitamins include ascorbic acid (vitamin C) and the B-complex vitamins, such as biotin, cobalamin (vitamin B₁₂), folate (folic acid), niacin (vitamin B₃), pyridoxine (vitamin B₆), riboflavin (vitamin B₂), thiamine (vitamin B₁), and pantothenic acid.

These tests are measurements of nutritional status. Low levels indicate recent inadequate oral intake, poor nutritional status, and/or malabsorption problems. They may not reflect tissue stores. High levels indicate excessive intake, intoxication, or absorption problems.

Reference Values

Dietary Reference Intakes (DRIs), the most recent approach adopted by the Food and Nutrition Board, Institute of Medicine, and National Academy of Sciences, provide estimates of vitamin intake. The DRIs look beyond deficiency disease and include the role of nutrients and food components in long-term health. The DRIs consist of four reference intakes: Recommended Daily Allowances (RDAs), Tolerable Upper Intake Levels (ULs), Estimated Average Requirements (EARs), and Adequate Intake (AI). When an RDA cannot be set, an AI is given as a normal value; both are to be used as goals for the patient. Levels are given for each individual vitamin. The RDAs are the amounts of ingested vitamins needed by a healthy person to meet daily metabolic needs, allow for biologic variation, maintain normal blood serum values, prevent depletion of body stores, and preserve normal body functions.

Procedure

1. Examine blood, urine, and hair or nail samples for vitamin levels. The types of specimens needed are listed in the table.
2. Test for vitamins by both direct and indirect methods.

Clinical Implications

Increased and decreased levels and critical ranges are shown in the table.

Interfering Factors

Factors that affect vitamin levels include age, season of the year, diarrhea or vomiting, certain drugs, various diseases, and long-term hyperalimentation.

Interventions

Pretest Patient Preparation

1. Assess overall nutritional status and address potential deficiencies. Oftentimes, one deficiency is accompanied by several nutrient deficiencies.
2. Evaluate signs and symptoms of disrupted vitamin-related metabolic reactions that indicate the need for testing.
3. Be aware that cost of testing (high) and time frames for obtaining test results (slow) are issues. Samples for vitamin tests are usually sent to specially laboratories, which increases cost and turnaround time dramatically.
4. Explain the purpose of the test before collecting blood, urine, hair, or nail specimens.
5. Inform the patient that vitamins are micronutrients that can be detected in the blood and urine as an indication of
...overt nutritional deficiency states, toxic levels, or subclinical hypovitaminosis. The potential for toxicity from excessive intake exists.

6. See Chapter 1 guidelines for safe, effective, informed pretest and intratest care.

7. Intratest care—Collect required specimens.

**Posttest Patient Aftercare**

1. Verify and report reference ranges (RR) and critical ranges (CR). Take appropriate action when values are too high or too low. Treat nutrient deficiencies and toxicities immediately.

2. In collaboration with other clinicians (eg, pharmacist, dietician), counsel the patient about abnormal results, follow-up tests, dietary changes, and treatment. Water-soluble vitamins are needed on a daily basis. Reference ranges vary and are method dependent. Check with your laboratory.

3. Follow Chapter 1 guidelines for safe, effective, informed posttest care.

### Substance Tested (Specimen Needed), Reference Range (RR), and Critical Toxic Range (CR) and RDIs When Available

#### FAT-SOLUBLE VITAMINS

**Vitamin A**

- **Retinol (serum)**
  - **RR:** 360–1200 µg/L or 0.70–1.75 µmol/L
  - **CR:** <10 µg/dL or <0.35 µmol/L indicates severe deficiency, >100–2000 µg/dL or >3.49–69.8 µmol/L indicates hypervitaminosis A
  - **Carotene (serum)**
    - **RR:** 50–300 µg/dL or 1.5–7.4 µmol/L
    - **CR:** >250 µg/dL or >4.65 µmol/L indicates carotenemia
  - **Retinyl esters <10 µg/L when selected**

- **Relative dose response (%)**
  - RR: >20; CR: >50 deficiency

- **Patient Preparation, Substance Function, and Indications for Test**
  - **Fasting. No alcohol 24 h before blood draw**
  - **Prevents night blindness and other eye problems (acne)**
  - **Enhances immunity, protects against pollution and cancer formation**
  - **Needed for maintenance repair of epithelial tissues**
  - **Aids fat storage**
  - **Acts as antioxidant (protects cells against cancer and other diseases)**
  - **Evaluate night blindness, mal-absorption disorders, chronic naphritis, acute protein deficiency, Bitot's spots, intestinal parasites, acute infections, chronic intake of >10 mg retinol equivalent (RE)**

- **Clinical Significance of Values**
  - **Increased**
    - Activation of phagocytes and/or cytotoxic T-cells
    - Alopecia
    - Amenorrhea
    - Arthralgia (gout)
    - Bitot's spots
    - Celiac disease
    - Cirrhosis of the liver
    - Congenital obstruction of the jejunum
    - Cystic fibrosis
    - Duodenal bypass
    - Fat malabsorption syndrome
    - Giardiasis
    - Immunity compromised (cell-mediated response, antibody response)
    - Insufficient dietary intake
    - Keratinization of lung, gastrointestinal tract, and urinary epithelia
    - Pregnancy
    - Premature epiphysial closure
    - Pseudotumor cerebri
    - Spontaneous abortions
    - Nystagloplasia (night blindness)
    - Oral contraceptives (carotene)
    - Pancreatic surgery
    - Protein-energy malnutrition (marasmus or kwashiorkor)
    - Perifollicular hyperkeratosis (Darier's disease)
    - Sprue
    - Xerophthalmia
    - Xerosis of the conjunctiva and cornea

- **Decreased**
  - Acute infections
  - Arthralgia (gout)
  - Bile duct obstruction
  - Bilio's spots
  - Celiac disease
  - Cirrhosis of the liver
  - Congenital obstruction of the jejunum
  - Cystic fibrosis
  - Duodenal bypass
  - Fat malabsorption syndrome
  - Giardiasis
  - Immunity compromised (cell-mediated response, antibody response)
  - Insufficient dietary intake
  - Keratinization of lung, gastrointestinal tract, and urinary epithelia
  - Pregnancy
  - Premature epiphysial closure
  - Pseudotumor cerebri
  - Spontaneous abortions
  - Nystagloplasia (night blindness)
  - Oral contraceptives (carotene)
  - Pancreatic surgery
  - Protein-energy malnutrition (marasmus or kwashiorkor)
  - Perifollicular hyperkeratosis (Darier's disease)
  - Sprue
  - Xerophthalmia
  - Xerosis of the conjunctiva and cornea

**Vitamin D**
1, 25-dihydroxycholecalciferol, calciferol (serum)
RR: 60 ng/mL or 150 nmol/L
Toxic: >150 ng/mL or >375 nmol/L
Deficient: <10 ng/mL or <25 nmol/L
CR: Serum calcium levels of 12–16 mg/dL or 3.0–4.0 mmol/L (vitamin D toxicity)
RDI: Adults: Cholecalciferol 10 µg/d or 400 U of vitamin D

Fasting
Synthesized by skin exposure to the sunshine
Required for absorption of calcium and phosphorous by the intestinal tract
Necessary for normal development of bones in children
Protects against muscle weakness, involved in regulation of heartbeat
Important in treatment of osteoporosis and hypocalcemia
Evaluate rickets, osteomalacia, fat malabsorption; disorders of parathyroid, liver or kidney; prolonged supplement intake of 2,000 IU/d
Gastrointestinal symptoms (anorexia, nausea, vomiting, constipation)
Infants: “elfin facies,” hypercalcemia with failure to thrive, mental retardation, stenosis of the aorta
Metastatic extraosseous calcification
Renal colic
Supplements
Williams’ syndrome
Anticonvulsants
Familial hypophosphatemic rickets (diabetes mellitus, Fanconi’s syndrome, hypoparathyroidism, renal osteodystrophy, renal tubular growth acidosis)
High phosphate or phytate intake
Inadequate diet
Inadequate exposure to sunlight (especially in the elderly)
Liver disease
Malabsorption syndromes
Osteomalacia (adults)
Rachitic tetany

Vitamin E
Serum alpha tocopherol, TE (most active)
RR (serum): Adults: 0.5–1.8 mg/dL or 12–42 µmol/L
NOTE: Concentration of vitamin E in newborns is less than half that of adults
RDI: Men: 10 mg Alpha tocopherol equivalent (a-TE)
Women: 8 mg/a-TE
Fasting. No alcohol 24 hours before draw
Antioxidant
Important in prevention of cancer and cardiovascular diseases
Promotes normal blood clotting, healing
Improves circulation necessary for tissue repair; maintains healthy nerves and muscles while strengthening capillary walls
Prevents cell damage by inhibiting the oxidation of lipids and formation of free radicals (antioxidants)
Aids utilization of vitamin A
Retards aging and may prevent age spots
Evaluate premature-birth-weight infants, abetalipoproteinemia, malabsorption
Low-birth-weight infants (sepsis, necrotizing enterocolitis)
Vitamin E supplementation increased bleeding tendency
Impaired leukocyte formation
Reduced cataract formation (with high ß-carotene and ascorbic acid levels)
Menstrual problems
Infertility (men and women)
Biliary atresia
Carotid deposits in muscle
Cholestasis
Dermatitis (flaky)
Edema
Malabsorption syndromes with steatorrhea
Neurologic syndromes affecting the spinal posterior columns and the retina
(abetal- or hyperlipoproteinemia), blinding loop syndrome, chronic pancreatitis, cystic fibrosis, inborn errors of metabolism, obstructive liver disease, short bowel syndrome)
Premature infants (broncho-pulmonary dysplasia, intraventricular hemorrhage, platelet dysfunction, low retinopathy)
Protein-energy malnourished children
Reperfusion injury
Platelet hyperaggregation, decreased erythrocyte survival and increased susceptibility to hemolysis

Vitamin K
Phylloquinone (K₁) plants; menaquinone (K₂ series) bacterial; menadione (K₂) synthetic
RR: 1.3–1.9 ng/mL, PIVKA 11 test (proteins induced in vitamin K absence). This test is superior. Plasma prothrombin concentration 10.5–12.5 seconds
RDI: Men: 80 µg/d Women: 60–65 µg/d

Fasting

Necessary for the production of prothrombin (blood clotting)
Essential for bone formation and repair
Necessary for synthesis of osteocalcin (the protein in bone tissue on which calcium crystalizes)
Therefore, prevents osteoporosis
Plays role in converting glucose into glycogen for storage in liver
Antibiotics interfere with absorption of vitamin K
Evaluate renal insufficiency and chronic antibiotic treatment

Glucose-6-phosphate dehydrogenase deficiency
Increased dietary intake or administered vitamin K preparation
Low-birth-weight infants (increased menadione)
Anemia with Heinz bodies
Hyperbilirubinemia
Kernicterus (bilirubin encephalopathy)
Loss of sucking reflex
Postkernicterus syndrome
Breast-fed infants (no Vitamin K received)
Conditions limiting for absorption or synthesis of vitamin K
Coumadin (warfarin)
Excessive oral mineral oil
Hypoprothrombinemia
Dietary lack
Lack of bile salts (external biliary fistulas, obstructive jaundice)
Liver disease
Nonabsorbable sulfonamides
Salicylate therapy
Megadoses of fat-soluble vitamins A or E are known to antagonize vitamin K
Long-term total parenteral nutrition
Chronic fat malabsorption, pancreatic disease, gastro-intestinal disease

WATER-SOLUBLE VITAMINS

Ascorbic Acid (Vitamin C)
RR: 0–114 µmol/L plasma; 0–2.0 mg/dL plasma, 114–301 nmol/10³ cells (mixed leukocytes), 20–53 µg/10³ cells (mixed leukocytes)
CR: <11 µmol/L plasma ascorbate, <0.2 mg/dl plasma ascorbate, <57 nmol/10³ cells (mixed leukocytes), <10 mg/10³ cells (mixed leukocytes)

Women consistently show higher vitamin C levels in tissues and fluids than men.
Plasma values are the best indicator of recent dietary intake.
Leukocyte vitamin C levels are indicative of cellular stores and body pool.

NOTE: Salivary vitamin C levels are not consistent; urinary vitamin C levels are not useful.
RDI: Adults: 60 mg/d

Antioxidant needed for tissue growth and repair, adrenal gland function, and healthy gums
Aids in production of antistress hormones and interferon; needed for metabolism of folic acid, tyrosine, and phenylalanine
Increases absorption of iron; reduces cholesterol levels and high blood pressure
Essential in neurotransmitter synthesis and metabolism
Essential in the formation of collagen; promotes wound healing; protects against infection
Enhances immunity
Evaluate scurvy, poor diet, and nephrolithiasis.

Decreased anticoagulant effect of heparin and warfarin (Coumadin)
Diarrhea
Overabсорption of iron Supplementation (alteration of tests for diabetes and occult blood)
Nausea
Some patients with history of kidney stones are at an increased risk of oxalate stones with too much vitamin C intake.

Adult scurvy (acne, listlessness, deep muscle hemorrhages, swan-neck deformity, gingivitis, perifollicular hemorrhages, hyperkeratosis, and hypochondriasis)
Alcoholism and drug abuse
Anemia (microcytic hypochromic)
Burns
Cold or heat stress
Edema, lower extremities
Gastric ulcers
Impaired iron absorption
Inadequate diet (especially elderly men)
Infantile scurvy (Barlow's disease, "pithed frog" position)
Inflammatory diseases, oxidative damage (proteins, DNA, human sperm DNA)
Lactation
Peteichae and total echymoses
Pregnancy
Risk of cancer (esophagus, oral cavity, uterine, cervix)
Smokers (decreased ascorbic acid half-life)
Thyrotoxicosis
Toxicity from chemical carcinogens (anthracene, benzpyrene, organochloride pesticides, heavy metals, nitrosamines)
Poor wound healing
Bleeding gums, dyspnea, edema, and weakness

Biotin
### Biotin

- **RR:** 0.82–2.87 nmol/L
- **CR:** <1.02 nmol/L deficiency (whole blood or serum)
- **RR:** 200–500 pg/mL or 0.82–2.05 nmol/L

Prenatal diagnosis of multiple carboxylase deficiency (MCD) by direct analysis of amniotic fluid for methylcitric acid or 3-hydroxyisovaleric acid.

**RDI:** 30 µg/d

- Biotin is produced by the gut flora.
- Aids in cell growth, fatty acid production, metabolism of fats, carbohydrates, and proteins, and utilization of other complex vitamins
- Promotes healthy sweat glands, nerve tissue, and bone marrow
- Needed for healthy hair and skin
- Assess for ingestion of raw eggs, inflammatory bowel disease, alcoholism, sulfonamide therapy, depression.

Biotin is also produced by the gut flora. It aids in cell growth, fatty acid production, metabolism of fats, carbohydrates, and proteins, and utilization of other complex vitamins. They promote healthy sweat glands, nerve tissue, and bone marrow. They are also needed for healthy hair and skin. They assess for ingestion of raw eggs, inflammatory bowel disease, alcoholism, sulfonamide therapy, and depression.

### Cobalamin (Vitamin B₁₂)

- **RR:** >200–835 pg/mL or 148–616 pmol/L
- **CR:** <100 pg/mL or <74 pmol/L deficiency

**RDI:** Adults 2.4 µg/d

- Overnight fast. Avoid heparin, ascorbic acid, fluoride, and alcohol before testing.
- Aids folic acid in formation of iron; prevents anemia
- Required for proper digestion, absorption of food, synthesis of protein and metabolism of fats and carbohydrates.
- Prevents nerve damage, maintains fertility, production of acetylcholine (neurotransmitter that assists memory and learning)
- Found mostly in animal sources, so strict vegetarians may need supplements
- Regional enteritis
- Evaluate strict vegetarian diet spanning 20–30 y, alcoholism, after gastrectomy, and parasitic infections.

Cobalamin is also known as Vitamin B₁₂. It is present in large amounts of raw egg white and ingestion of raw avidin. It can also be found in antibiotics, biotin responsive MCD syndromes, changes in mental status (depression), glossitis (magenta hue), high fetal resorption rate, hyperesthesia (algesia), immunodeficiency, increased serum, cholesterol and bile pigments, ingestion of large amounts of (6/d) of raw egg white, ingestion of raw avidin, localized paresthesia, maculosquamous dermatitis of the extremities, myalgia, pallor, long-term total parenteral nutrition after gut resection, if not supplemented, high blood sugar.

### Folate (Folic Acid)

- **RR:** 0.82–2.87 nmol/L
- **CR:** <1.02 nmol/L deficiency (whole blood or serum)
- **RR:** 200–500 pg/mL or 0.82–2.05 nmol/L

Prenatal diagnosis of multiple carboxylase deficiency (MCD) by direct analysis of amniotic fluid for methylcitric acid or 3-hydroxyisovaleric acid.

**RDI:** 30 µg/d

- Biotin is produced by the gut flora.
- Aids in cell growth, fatty acid production, metabolism of fats, carbohydrates, and proteins, and utilization of other complex vitamins
- Promotes healthy sweat glands, nerve tissue, and bone marrow
- Needed for healthy hair and skin
- Assess for ingestion of raw eggs, inflammatory bowel disease, alcoholism, sulfonamide therapy, depression.

Folate, also known as Folic Acid, is not only produced by the gut flora. It also aids in cell growth, fatty acid production, metabolism of fats, carbohydrates, and proteins, and utilization of other complex vitamins. They also promote healthy sweat glands, nerve tissue, and bone marrow. They are also needed for healthy hair and skin. They assess for ingestion of raw eggs, inflammatory bowel disease, alcoholism, sulfonamide therapy, and depression.

Deficiency caused by malabsorption—common in elderly and those with digestive disorders
- Alcoholism
- Addisonian pernicious anemia
- Thalassemia
- Diet lacking microorganisms and animal foods (sole B₁₂ sources)
- Distal sensory neuropathy ("glove and stockings")
- Sensory loss
- Gastrectomy
- Gastric atrophy (superficial gastritis, hereditary—degenerative congenital)
- Liver disease
- Pigmentation of skin creases and nailbeds (brownish)
- Polyendocrinopathy
- Pregnancy
- Renal disease
- Small intestine disorders (cancer, gluten-induced enteropathy—celiac disease, granulomatous lesions, intestinal resections, "stagnant bowel" syndrome, tropical sprue)
- Subacute combined degeneration of the cord
- Tapeworms
- Tinnitus and noise-induced hearing loss
- Tongue—red, smooth, shining, painful
- Vegans (and their breast-fed infants)
- Visual loss from optic atrophy
- Zollinger-Ellison syndrome
(pteroylglutamate, pteroylglutamic acid, 5-methyltetrahydrofolate)
Red blood cell folate (best indicator of status)
RR: 150–800 ng/mL or 340–18,120 nmol/L whole blood, corrected to packed cell volume of 45%
Tissue folate depletion (serum dietary fluctuations): <160 ng/mL or <360 nmol/L
RR: 3–21 ng/mL or 6.7–47.5 nmol/L
CR: <1.5 ng/mL or <3.4 nmol/L deficiency
Negative folate balance: <3 ng/mL or <7 nmol/L,
RDI: Adults: 400 µg/d
Other methods (infrequently used):
Deoxyuridine suppression test (DU or dUST), a functional indicator of folate status; in vitro laboratory test that defines presence of megaloblastosis and identifies which nutrient deficiency is responsible (folate or vitamin B₁₂)
Formiminoglutamic acid (FIGLU)—after histidine loading
Urine—24 hours after initial dose
RR: <35 mg/d or <201 µmol/d
Folate deficiency: <35 mg/d or <201 µmol/d

Fasting
Needed for energy production and formation of red blood cells
Strengthens immunity by aiding white blood cell functioning
Important for healthy cell division and replication (DNA and RNA synthesis)
Protein metabolism
Prevention of folic acid anemia
In pregnancy, regulates embryonic and fetal nerve cell formation, prevents premature birth
Works best when combined with vitamins B₁₂ and C
Cooking destroys folic acid
Evaluate megaloblastic anemia, cancer, inflammatory bowel disease, alcoholism, drug treatment with phenytoin, cholestyramine, sulfasalazine, oral contraceptives
Detect folate deficiency

Riboflavin (Vitamin B₂)

Folacin is dominant form in serum and RBC
Loss of seizure control
Acute renal failure
Active liver disease
Red blood cell hemolysis
Supplemental folate (400 µg/4 mg/d—side effects)

Alcohol, alcoholics
Liver disease
Elderly
Breast-fed infants of mothers taking estrogen-progesterone contraceptives
Cervical dysplasia
Cigarette smoking
Drug therapy (phenytoin, primidone, barbiturates)
methotrexate, metformin, cholestyramine, cycloserine, azathioprine, oral contra-ceptives, antacids)
Increased requirements
Hematopoiesis (thalassemia major)
Increased metabolism
Infancy
Lactating women
Malignancy
(lymphoproliferative)
Pregnancy
HPV-16 infection
Inadequate dietary intake
Malabsorption syndromes (celiac disease, sprue, blind loop syndrome)
Megaloblastosis
Neural tube defects (spina bifida, anencephaly)
Pancytopenia
Protection from malaria
Psoriasis
Renal dialysis
Scurvy
Tongue papillae atrophy (shiny, smooth)
Vitamin B₁₂ deficiency
Increased mean corpuscular volume
Depression
Methotrexate-treated patients
Hyperhomocysteinemia
Long-term unsupplemented total parenteral nutrition
Rheumatoid arthritis
(serum or plasma) RR: 4–24 µg/dL or 106–638 nmol/L
urine—much more sensitive to nutritional status RR: >80 µg/d or >213 nmol/dL erythrocyte: 10–50 µg/dL or 266–1330 nmol/L
Creatinine indicates deficiency <27 µg/g creatinine (urine) or <8 µmol/mol creatinine
Erythrocyte glutathione reductase assay, expressed in activity coefficients (AC). Test cannot be used in persons with glucose-6-phosphate deficiency RR: 10–50 µg/dL or 266–1330 nmol/L
RDI: Men: 1.3 mg/d Women: 1.1 mg/d

Fasting Necessary for red blood cell formation, antibody production, cell respiration, and growth
Alleviates eye fatigue and important in treatment and prevention of cataracts
Aids metabolism of fat, carbohydrates, and protein
With Vitamin A, maintains and improves mucous membranes in digestive tract
Helps absorption of iron and B6
Pure, uncomplicated riboflavin deficiency is rare—if seen, it is usually accompanied by multiple nutrient deficiencies.
Needed for metabolism of amino acid tryptophan, which is converted to niacin in the body
Easily destroyed by light, anti-biotics, and alcohol
Increased need for B2 with use of oral contraceptives or strenuous exercise
Assess poor dietary intake, as in congenital heart disease and some cancers.

Niacin (Vitamin B3)
Nicotinic acid, niacinamide (urinary N’-methyl nicotinamide, NMN) 24-h urine RR: 2.4–6.4 mg/d or 17.5–46.7 µmol/d CR: <0.8 mg/d or <5.8 µmol/d deficiency RDI: Men: 16 mg/d Women: 14 mg/d

Pyridoxine (Vitamin B6)

Abnormal liver function
Hypcholesterolemia
Use as hypolipidemic drug
Atrial fibrillation
Cystoid maculopathy
Epigastric discomfort
Glucose intolerance
Gout
Hyperglycemia
Hypotension
Pruritus
Smooth, swollen tongue
Upper body flushing

Alcoholics
Carcinoid syndrome
Casal’s necklace
Cirrhosis of the liver
Diarrheal disease
Diet lacking in niacin and tryptophan
Dyssebacia
Isoniazid therapy
Hartnup’s disease
Isoniazid therapy
Pellagra dermatosa; glossitis (scarlet, raw beef)
Gastrointestinal dysfunction
Central nervous system dysfunction
Organic psychosis
Encephalopathic syndrome
RR (direct)
Plasma vitamin B₆: 5–24 ng/mL or 20–97 nmol/L
Plasma pyridoxal 5 phosphate >7 ng/mL or >30 nmol/L
Plasma total vitamin B₆ > 10 ng/mL or >40 nmol/L
Urinary 4-pyridoxic acid (4rPA) <3.0 µmol/d (useful short-term index)
Urinary total vitamin B₆ antagonists B₆ >0.5 µmol/day (isoniazid, penicillamine, cycloserine)
RR (indirect):
Erythrocyte alanine transaminase index (EALT/EGPT) >1.25 (EALT is a better indicator than EAST; standardized approach needed to compare tests).

Fasting or urine collection Needed for production of hydrochloric acid and absorption of fats and protein, sodium and potassium balance, and red blood cell formation Required by nervous system for normal brain function

Tryptophan metabolism Niacin formation Gluconeogenesis Synthesis of nucleic acids, RNA and DNA: activates many enzymes and aids in absorption of vitamin B₁₂ Cancer immunity, prevents arteriosclerosis Mild diuretic—reduces premenstrual syndrome Diuretics and cortisone drugs block absorption of B₆ Antidepressants, estrogen therapy, and oral contraceptives increase need for B₆ Evaluate groups at risk, including newborn infants with low B₆, some cancers, excess alcohol.

Infants: neurologic symptoms and abdominal distress Peripheral neuropathy; progressive sensory ataxia; lower limb impairment Photosensitivity Neurotoxicity

Alcoholism
Anemias
Asthma
Breast cancer
Cheliosis
Coronary heart disease
Depression and confusion
Diabetes
Drugs (isoniazid, cycloserine, penicillamine, ethinyl estradiol, mestranol)
Glossitis
Hodgkin’s disease
Impaired interleukin-2 production
Increased metabolic activity
Infants (abnormal electro-encephalogram pattern, confusions)
Irritability
Lymphopenia
Peripheral neuropathy
Prenestral syndrome
Seborrheic dermatosis
Sickle cell anemia
Smokers
Stomatitis

RR (stimulation):

Thiamine (Vitamin B₁)

RR: 0.2–0.4 µg/dL or 5.9–118 nmol/L (serum or plasma)
2.5–7.5 µg/dL or 74–222 nmol/L (whole blood)
Late changes: <50 µg/d or <148 nmol/d urine with elevated blood pyruvate
Red blood cell transketolase measurement (most reliable method)
Enzyme assays—using thiamine pyrophosphate (TPP): 79–178 nmol/L
RR (stimulation): 0%–25%; deficiency, >20%
RDI: Men: 1.3 mg/d
Women: 1.1 mg/d

Parenteral dosages
High-carbohydrate diet increases need for B₁
Thiamin is poorly absorbed in adults with folate or protein deficiency.

Parenteral dosages

Antibiotics, sufla drugs, oral contraceptives
Alcoholism
Beriberi—dry beriberi (peripheral neurologic changes; ie, symmetric foot drop); infantile beriberi; wet beriberi
Cardiovascular (high-output congestive heart failure, low-output Shoshin disease)
Wernicke-Korsakoff syndrome (acute hemorrhagic polio-encephalitis)
Cerebral beriberi
Dependency states (thiamine-responsive megaloblastic anemia, lactic acidosis, ketoaciduria, subacute necrotizing encephalopathy, Leigh’s disease)
Dextrose infusions (frequent, long-continued or highly concentrated)
Folate deficiency
High-carbohydrate diet (mainly from milled [polished] rice)
Hyperthyroidism
Impaired absorption (ie, long-term diarrheas)
Impaired utilization (ie, severe liver disease)
Inadequate calorie or protein intake
Increased requirements (fever, lactation, pregnancy, strenuous physical exertion)
Poor memory
Renal dialysis
Long-term total parenteral nutrition

Fasting Enhances circulation and blood formation, carbohydrate metabolism, and production of hydrochloric acid Optimizes cognitive activity and brain function Has a positive effect on energy, growth, normal appetite, and learning capacity Needed for muscle tone of intestines, stomach, and heart Acts as antioxidant, protecting body from degenerative effects of aging, alcohol consumption, and smoking Evaluate alcoholism, impaired absorption, excess intravenous glucose infusion, in diets primarily of refined, unenhanced grain products

Antibiotics, sufla drugs, oral contraceptives
Alcoholism
Beriberi—dry beriberi (peripheral neurologic changes; ie, symmetric foot drop); infantile beriberi; wet beriberi
Cardiovascular (high-output congestive heart failure, low-output Shoshin disease)
Wernicke-Korsakoff syndrome (acute hemorrhagic polio-encephalitis)
Cerebral beriberi
Dependence states (thiamine-responsive megaloblastic anemia, lactic acidosis, ketoaciduria, subacute necrotizing encephalopathy, Leigh’s disease)
Dextrose infusions (frequent, long-continued or highly concentrated)
Folate deficiency
High-carbohydrate diet (mainly from milled [polished] rice)
Hyperthyroidism
Impaired absorption (ie, long-term diarrheas)
Impaired utilization (ie, severe liver disease)
Inadequate calorie or protein intake
Increased requirements (fever, lactation, pregnancy, strenuous physical exertion)
Poor memory
Renal dialysis
Long-term total parenteral nutrition
strenuous physical exertion
Poor memory
Renal dialysis
Long-term total parenteral nutrition
Unsupplemented
Mineral Testing

Minerals are micronutrients needed in relatively small amounts. Unlike vitamins, the source for minerals comes from nonliving, naturally occurring elements, such as mineral salts in the soil that become a part of the chemical constituents of food or minerals that are dissolved in ocean water and ingested in seafood. Mineral concentrations in blood, urine, and certain body tissues can be measured and reflect the nutritional status of the patient.

Minerals are either macronutrients (major) or micronutrients (trace or ultratrace). If the body requires a significant amount of the mineral (> 100 mg/d) and an RDA has been established, it is a macronutrient; if the body requires less (a few milligrams per day) and an RDA or Estimated Safe and Adequate Daily Dietary Intake (ESADDI) has been established, it is a micronutrient trace mineral; if the body requires <1 mg/d and no RDA or ESADDI has been established, it is a micronutrient ultratrace mineral.

Macronutrients (major minerals) include calcium, chloride, magnesium, phosphorous, potassium, sodium, and sulfur. Macronutrients are not listed in this table; they are explained in Chapter 6.

Trace minerals include the micronutrients chromium, cobalt, copper, fluorine, iodine, iron, manganese, molybdenum, selenium, and zinc.

Ultratrace minerals include the micronutrients arsenic, boron, bromine, cadmium, lead, lithium, nickel, silicon, tin, and vanadium.

Minerals found in the body without an assigned metabolic role include aluminum, antimony, beryllium, bismuth, cyanide (an anion that forms a salt with minerals), gold, mercury, silver, lead, thallium, and many others.

These measurements of minerals are used to assess environmental or occupational exposure and toxicity, monitor effectiveness of treatment, and evaluate mineral status along with other laboratory levels to verify deficiencies.

References Values

Dietary Reference Intakes (DRIs), the most recent approach adopted by the Food and Nutrition Board, Institute of Medicine, and National Academy of Sciences, provide estimates of mineral intake. The DRIs look beyond deficiency disease and include the role of nutrients and food components in long-term health and prevention of chronic disease (ie, calcium balance and calcium retention). The DRIs consist of four 4 reference intakes: Recommended Daily Allowances (RDAs), Tolerable Upper Intake Levels (ULs), Estimated Average Requirements (EARs), and Adequate Intake (AI). The RDAs and EAR levels have been established for some minerals, including those with and without an assigned role in the human body.

Procedure

1. Examine blood, urine, hair, or nail samples for mineral levels by indirect and direct methods.
2. Types of specimens required are listed in the table.

Clinical Implications

Increased and decreased levels and critical toxic ranges are found in the table.

Interfering Factors

Factors that affect mineral levels include: Genetic makeup

1. Age or stage of life cycle
2. Environmental factors
3. Drugs
4. Intestinal malabsorption
5. Stress
6. Strenuous physical activity
7. Smoking
8. Alcohol consumption
9. Dietary intake

**Interventions**

**Pretest Patient Preparation**

1. Evaluate overall nutritional status, dietary intake, and supplement usage to determine overconsumption.
2. Evaluate signs and symptoms of occupational and environmental toxicity and mineral deficiencies that indicate the need for testing.
3. Explain the purpose of the test before collecting blood, urine, hair, or nail specimens.
4. Inform the patient that minerals are nutrients that can be detected in the blood and urine as an indication of toxicity or exposure and nutritional status. The amounts needed are determined by what is necessary for optimal function and health and to prevent disease.
5. See Chapter 1 guidelines for safe, effective, informed pretest and intratest care.

**Posttest Patient Aftercare**

1. Verify and report reference ranges (RR) and critical toxic ranges (CR). Take appropriate action when values are too high or too low.
2. Counsel the patient appropriately about abnormal results, follow-up testing, occupational and lifestyle changes, treatment, and diet. Reference ranges vary and are method dependent. Check with your laboratory. Notify employer, workplace, and physician about exposure results.
3. Follow Chapter 1 guidelines for safe, effective, informed postest care.

<table>
<thead>
<tr>
<th>Substance Tested (Specimen Needed), Reference Range (RR), Critical Toxic Range (CR), and RDIs if Available</th>
<th>Patient Preparation, Substance Function, and Indications for Test</th>
<th>Clinical Implications</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Aluminum (Al)</strong> (serum) RR: 20–550 µg/L or 0.74–20.4 µmol/L dialysis patients (urine) RR: 5–30 µg/L or 0.19–1.11 µmol/L</td>
<td>Collect urine in acid-washed polypropylene container. Assess for occupational exposure, toxicity from antacids; monitor dialysis patients.</td>
<td>Increased</td>
</tr>
<tr>
<td></td>
<td>No metabolic role Metal used in other forms as an astringent (Burrow’s solution) and as an antacid</td>
<td>Aluminum absorption with citrate-containing drugs (effervescent or analgesics) Use of aluminum-containing astringents, hydroxide gels, aluminum-containing phosphate binders Excessive occupational exposure</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Toxicity:</strong> Aluminumosis (lung disease) Aluminum-induced encephalopathy Hypophosphatemia Dialysis dementia Iron-resistant microcytic anemia Aluminum-related osteomalacia In renal failure, when aluminum containing antacids are used; long-term intermittent dialysis</td>
</tr>
<tr>
<td><strong>Antimony (Sb)</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NOTE: Aluminum is a neurotoxin. The primary symptom is motor dysfunction leading to dysarthria, myoclonus, or epilepsy. Aluminum toxicity is not related to Alzheimer's disease. Aluminum can be found in laboratory solutions used with tissue samples and in laboratory dust. New testing methods are being adopted to rule out contamination.
<table>
<thead>
<tr>
<th>(24-h urine)</th>
<th>RR: &lt;10 µg/L or CR: &gt;1 mg/L or CR: &gt;82.1 nmol/L (plasma)</th>
<th>RR: 0.03–0.07 µg/dL or 2.5–5.7 nmol/L</th>
</tr>
</thead>
<tbody>
<tr>
<td>No metabolic role</td>
<td>Compounds used in alloys, medicines, poisons</td>
<td>Assess for occupational exposure and toxicity.</td>
</tr>
<tr>
<td>Excessive occupational exposure (ore from mining, bronze ceramics)</td>
<td>Ingested compounds (drugs used in parasitic infections)</td>
<td>Toxicity: Acid metallic taste, burning gastrointestinal pain (as in arsenic poisoning), throat constriction, dysphagia, pulmonary edema, liver and renal failure</td>
</tr>
<tr>
<td><em>Lethal dose</em>: 5–50 mg/kg body weight</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Arsenic (As)**

- **(hair or nails)**
  - RR: <1.0 µg/g of hair or nails (serum)
  - RR: >5 µg/mL or >0.07 µmol/L
- **Normal concentration**
  - RR: 0.03–0.07 µg/dL or 2.5–5.7 nmol/L
- **Toxic concentration**
  - RR: >20 µg/L or 2.22 µmol/L

**Ultratrace mineral; no function**

**Found in pesticides and paints**

**Used as a homicidal poison**

**Assess for occupational exposure and toxicity.**

Dermatoses (hyperpigmentation, hyperkeratosis, desquamation, and hair loss), hematopoietic for hair and nails

Depression

Liver damage characterized by jaundice

Peripheral neuropathy

Acute beryllium disease (a chemical pneumonitis)

- **Acute poisoning**
  - RR: >20 µg/L or >2.22 µmol/L
  - RR: 2–23 µg/L or 0.03–0.31 µmol/L

**Historical**—beryllium mining, electronics, chemical plants, manufacture of fluorescent lights (inhalation, introduction into or under skin and/or conjunctiva): berylliosis or granulomatosis

NOTE: Almost impossible to distinguish from sarcoidosis

<table>
<thead>
<tr>
<th>(24-h urine)</th>
<th>RR: negative, none detected</th>
<th>CR: &gt;20 µg/L or &gt;2.22 µmol/L</th>
</tr>
</thead>
<tbody>
<tr>
<td>No metabolic role; a metallic element.</td>
<td>Assess for occupational exposure and toxicity.</td>
<td>Acute beryllium disease (a chemical pneumonitis)</td>
</tr>
<tr>
<td>Excessive occupational exposure (metal extraction, refinery, rocket base, nuclear plant, extensive coal burning); secondary polycythemia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Historical—beryllium mining, electronics, chemical plants, manufacture of fluorescent lights (inhalation, introduction into or under skin and/or conjunctiva): berylliosis or granulomatosis</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NOTE: Almost impossible to distinguish from sarcoidosis
**Bismuth (Bi)**

- **(24-h urine)**
  - RR: 0.3–4.6 µg/L or 1.4–22.0 nmol/L
- **(plasma)**
  - 0.1–3.5 µg/L or 0.5–16.7 nmol/L

Collect urine in metal free container.

No metabolic role. Workers exposed in cosmetics, disinfectants, pigments, and solder industries Used in some drugs; poisoning as a consequence of therapy for syphilis Assess for occupational exposure, toxicity, and medication levels.

Bismuth used as treatment for syphilis in a growing child when mother has been treated during pregnancy

Treatment of peptic ulcer with bismuth-containing drug (zolimidine, colloidal bismuth subcitrate)

Bismuth subcarbonate, subgallate, and subnitrate compounds (used as antiseptics, astringents, sedatives, and to treat diarrhea and inflamed skin)

**Toxicity:** Ulcerative stomatitis, anorexia, headache, rash, renal tubular damage, bluish line at gum margin, albuminuria; resembles lead poisoning, without the blood changes and paralysis; rheumatic-like pain

**Boron (Bo)**

- **(blood, 4-mL serum)**
  - total RR: <2 mg/L or 33 µmol/L
  - CR: >20 mg/L or >330 µmol/L

Ultratrace mineral; a nonmetallic element, found as a compound such as boric acid or borax

Assess for exposure and toxicity, ingestion of boric acid, and un-expected absorption of boric acid from diapers or infant pacifier dipped in borax preparation and honey.

Increase in total plasma calcium concentrations and urinary excretions of calcium and magnesium

Decreased serum concentrations of 17B-estradiol, testosterone, and iodized calcium

Depressed mental alertness

**Bromine (Br), Bromide**

- **(serum)**
  - RR: 20–120 mg/dL or 2.5–15.0 mmol/L
  - RR: 1000–2000 mg/L or 12.5–25.0 mmol/L

Ultratrace mineral; a central nervous system depressant. Bromine is a liquid, nonmetallic element obtained from natural brines from wells and sea water; compounds used in medicine and photography

Assess for occupational exposure to bromide in medicine or photography.

Bromide acne

Neurologic disturbances

Increased spinal fluid pressure

**Toxicity:** Bromism or brominism

**Lethal dose:** 500–5000 mg/kg body weight

**Cadmium (Cd)**

- **(blood)**
  - RR: 0–5 ng/mL or 0–44 nmol/L
- **(urine—preferred)**
  - 0–5.0 µg/24h or 0–44 nmol/d
- **Toxic:** 100–3000 µg/L or 0.9–26.7 µmol/L

Ultratrace mineral, a metallic element in zinc ores Used in electroplating and in atomic reactors Its salts are poisonous.

Assess for occupational exposure, environmental poisoning.

In tissue, in prostatic and renal cancer

In urine, in hypertension, industrial exposure (electroplating—atomic reactors, zinc ores, cadmium solder)

**Toxicity:** Severe gastroenteritis, mild liver damage, acute renal failure; pulmonary edema; cough; duck-like gait; brown urine

**Lethal dose:** Several hundred mg/kg body weight

**Chromium (Cr)**
<table>
<thead>
<tr>
<th>Substance</th>
<th>Description</th>
<th>Reference Ranges</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Whole Blood</strong></td>
<td>Required for normal glucose metabolism; affects cholesterol synthesis</td>
<td>RR: 0.7–28.0 µg/L or 13.4–538 nmol/L</td>
<td>Excessive industrial exposure (carcinogenic)</td>
</tr>
<tr>
<td><strong>24-h Urine</strong></td>
<td></td>
<td>RR: 0.1–2.0 µg/d or 1.9–38.4 nmol/d</td>
<td>Increased risk for congestive heart disease</td>
</tr>
<tr>
<td></td>
<td>Assess for occupational exposure, poor diet, elderly at risk. Severe trauma and stress increase need.</td>
<td></td>
<td>Hypercholesterolemia</td>
</tr>
<tr>
<td><strong>Cobalt (Co)</strong></td>
<td>Essential element in vitamin B₁₂ — stimulates production of red blood cells</td>
<td>RR: 0.1–2.0 µg/d or 1.9–38.4 nmol/d</td>
<td>Cardiomyopathy after industrial exposure, during maintenance dialysis, and after drinking beer contaminated with cobalt during processing</td>
</tr>
<tr>
<td><strong>Copper (Cu)</strong></td>
<td>Required for hemoglobin synthesis, essential component of several enzyme systems; present in the liver and excreted by the kidneys and in bile</td>
<td>RR: 85–150 µg/dL or 13.3–23.6 µmol/L</td>
<td>T-cell proliferation, Hepatic glutathione</td>
</tr>
<tr>
<td><strong>Plasma</strong></td>
<td></td>
<td>RR: 90–150 µg/dL or 14.1–23.6 µmol/L</td>
<td>Wilson's disease (hepatolenticular degeneration)</td>
</tr>
<tr>
<td></td>
<td>Assess for excessive antacid intake, nephronic malabsorptive disorder, hemodialysis, and consumption of water high in copper by infants.</td>
<td></td>
<td>Ingestion of solutions of copper salts</td>
</tr>
<tr>
<td></td>
<td>Ceruloplasmin is an indirect test for copper</td>
<td></td>
<td>Contaminated water or dialysis fluids</td>
</tr>
<tr>
<td></td>
<td>RR: 21–53 mg/dL, 210–530 mg/L</td>
<td>Native-American childhood cirrhosis; Female rheumatoid arthritis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(neonate: 5–18 mg/dL, 50–180 mg/L)</td>
<td>Oral contraceptive use, Inflammatory conditions</td>
<td></td>
</tr>
<tr>
<td><strong>Cyanide (Cn Radical)</strong></td>
<td>No metabolic role</td>
<td>Industrial exposure (pesticides, metallurgy)</td>
<td>Cardiomyopathy after industrial exposure, during maintenance dialysis, and after drinking beer contaminated with cobalt during processing</td>
</tr>
<tr>
<td></td>
<td>The most common and most deadly poison — stops cellular respiration by inhibiting the actions of cytochrome oxidase, carbonic anhydrase, and other enzyme systems.</td>
<td></td>
<td>Ingestion of hydrocyanic acid and fumes from burning nitrogen-containing products</td>
</tr>
<tr>
<td></td>
<td>Toxicity comes from inhalation or ingestion—a hazard to firefighters.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Assess for industrial exposure, inhalation, or accidental poisoning from ingestion.</td>
<td>Toxicity: Lethal dose is &lt;5 mg/kg body weight (small child), fatal dose = 5–25 seeds. Death within 5 min of ingestion/inhalation. Adverse effects are dizziness, weakness, mental and motor impairment, and sudden death.</td>
<td></td>
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<tr>
<td><strong>Fluorine (F)</strong></td>
<td>Gaseous chemical found in soil in combination with calcium.</td>
<td>Fluorosis (excess fluorine use; &gt;4 million ppm in water; treatment of osteoporosis, multiple myeloma, or Paget's disease)</td>
<td>Marginal to deficient dietary intake from deficiencies in geochemical environments</td>
</tr>
<tr>
<td><strong>Plasma</strong></td>
<td>Used as a compound (fluoride) in toothpaste</td>
<td>Osteosclerosis</td>
<td>Dental caries</td>
</tr>
<tr>
<td></td>
<td>RR: 0.01–0.2 µg/mL or 0.5–10.5 µmol/L</td>
<td>Exostoses of spine and genuvalgum</td>
<td>Skeletal changes, especially in long bones</td>
</tr>
<tr>
<td></td>
<td>(urine)</td>
<td>Excess ingestion from swallowing fluoridated toothpaste</td>
<td></td>
</tr>
<tr>
<td></td>
<td>RR: 0.02–3.2 µg/mL or 1.05–168.3 µmol/L</td>
<td>Toxicity: Peculiar taste with salivation and thirst (salty-soapy), hemorrhagic gastroenteritis; hypoglycemia; central nervous system depression; renal failure</td>
<td></td>
</tr>
<tr>
<td></td>
<td>RDI: Men 4 mg/d Women 3 mg/d</td>
<td>Lethal dose: 50–500 mg/kg body weight</td>
<td></td>
</tr>
<tr>
<td><strong>Gold (Au)</strong></td>
<td>Required for normal glucose metabolism; affects cholesterol synthesis</td>
<td>Excessive industrial exposure (carcinogenic)</td>
<td>Insulin resistance (hyperinsulinemia)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Renal damage</td>
<td>Impaired glucose</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Increased risk for congestive heart disease</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Hypercholesterolemia</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Decreased fertility</td>
</tr>
</tbody>
</table>

**Cobalamin (vitamin B₁₂) deficiency**

**Menkes' steely hair disease:** lack of pigmentation of skin and hair

**Collagen abnormalities, osteoporosis**

**Ataxia**

**Hypochromic anemia**

**Unresponsive to iron therapy**

**Hypercholesterolemia**

**Impaired cardiovascular system**

**Rheumatoid arthritis**

**Oversupplementation of zinc**

**Altered interleukin-2 production**

**Neutropenia, leukopenia**

**Liver disease, kidney disease**

**Fluorosis (excess fluorine use; >4 million ppm in water; treatment of osteoporosis, multiple myeloma, or Paget's disease)**

**Osteosclerosis**

**Exostoses of spine and genuvalgum**

**Excess ingestion from swallowing fluoridated toothpaste**

**Toxicity:** Peculiar taste with salivation and thirst (salty-soapy), hemorrhagic gastroenteritis; hypoglycemia; central nervous system depression; renal failure

**Lethal dose:** 50–500 mg/kg body weight

**5–10 g sodium fluoride**
(colloidal gold in cerebrospinal fluid) RR: minute amount (serum) RR: <10 µg/dL or <0.5 µmol/L (therapeutic range) 100–200 µg/dL or 5.1–10.2 µmol/L Collect in metal-free container. No metabolic role; a metallic element. Salts used in early rheumatoid arthritis and in nondisseminated lupus erythematosus. Detectable in serum 10 mo after cessation of treatment. Assess for toxicity in treatment of rheumatoid arthritis.

**Iodine (I)**

(plasma) RR: 2–4 µg/dL, 60 ng/mL
Deficiency: IDD (iodine deficiency disorders) (daily urine) Mild IDD, RR: 50–100 µg/d (median urine, 3.5 µg/dL)
Moderate IDD, RR: 25–49 µg/d (median urine, 2–3.4 µg/dL)
Severe IDD, RR: <25 µg/d (median urine, 0–1.9 µg/dL)
RDI: adults 150 µg/d
Nonmetallic element belonging to the halogen group. Aids in the development and function of the thyroid gland, formation of thyroxine, and prevention of goiter. Assess for goiter.

Prolonged excessive intake of iodine leading to iodide-goiter and myxedema (common with pre-existing Hashimoto's thyroiditis) Excessive consumption of seaweed, kelp supplements; caffeine High dietary intake of known goitrogens (rutabagas, turnips, cabbages) Hypothyroidism in autoimmune thyroid diseases, inhibition of thioamide drugs Dysgeusia Acne-like skin lesions

Toxicity: Mucous membranes stained brown; burning pain in mouth and esophagus, laryngeal edema, shock, nephritis, circulatory collapse

**Iron (Fe)**

Rheumatoid arthritis if gold sodium thiomalate or gold thioglucose (aurothioglucose) is given parenterally; oral gold compound

Toxicity: At least 35% of patients undergoing chrysotherapy develop some degree of toxicity. Pruritus, dermatitis, stomatitis, albuminuria with or without nephrotic syndrome, agranulo-cytosis, thrombocytopenic purpura, and aplastic anemia

Adverse reactions: Enterocolitis, intrahepatic cholestasis, skin hyperpigmentation, peripheral neuropathy, and pulmonary infiltrates

Simple, endemic, colloid, or euthyroid goiter
Endemic cretinism (neurologic and/or myxedematous) Fetus: abortions, stillbirths, congenital anomalies
Child/teen: impaired mental function, retarded physical development
Adult: hypo- or hyperthyroidism, impaired mental function

Lethal dose: 5–50 mg/kg body weight
(serum, 5 mL, diurnal; morning specimen shows higher values)
RR: 35–140 µg/dL toxic: >300 µg/dL
Iron RR values: Males: 65–175 µg/dL or 11.6–31.3 µmol/L Females: 50–170 µg/dL or 9.0–30.4 µmol/L Newborn: 100–250 µg/dL or 17.9–44.8 µmol/L
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17.9–44.8 µmol/L
Total iron binding capacity (TIBC) RR: 250–100 µg/dL
Transferin RR values: Adult: 250–425 mg/dL or 44.8–76.1 µmol/L Newborn (0–4 days): 130–275 mg/dL or 1.30–2.75 g/L Child: 203–360 mg/dL or 2.03–3.60 g/L RDI: Adults: 10–15 mg/d
Essential to hemoglobin formation, transportation of oxygen, and cellular respiration Plays a role in the nutrition of epithelial tissues and the development of red blood cells Assess for ingestion of iron pills or vitamin and mineral pills (toxicity). Populations at risk for deficiency are infants and children 0.5–4.0 y, early adolescents, and women who are pregnant.
Diets high in heme iron or high in promoters of nonheme iron absorption Excessive iron absorption hereditary hemochromatosis (African or “Bantu” siderosis); prolonged therapeutic administration of iron to subjects not iron deficient; chronic alcoholism or liver disease, pancreatic insufficiency potential; “shunt hemochromatosis”; severe anemia with ineffective erythropoiesis and increased hemolysis; diabetes in 80% of patients Transfusional hemosiderosis β-thalassemia major, some chronic sideroblastic anemias, hypoplastic or other refractory anemias Other: cancers (primary hepatic carcinoma, acute leukemia, early breast cancer); demyelinating disease; Alzheimer's disease; increased risk of congestive heart disease, listeriosis Also see Chapter 6
Iron deficiency anemia: inadequate diet (grossly iron deficient, high in cereals, low in animal protein and vitamin C) Koilonychia (spoon-shaped nails) Excessive menstrual loss Pregnancy, lactation Blood donors Premature infants Intestinal helminthiasis (especially hookworm disease) Malabsorption syndromes, chronic diarrhea, gastrectomy, patients with atrophic gastritis and achlorhydria, occult gastrointestinal bleeding Hereditary hemorrhagic telangiectasia Turner's syndrome Angiodyplasia (vascular ectasis or arteriovenous anomaly) Blue rubber bleb nevi (hereditary cutaneous hemangiomas) Ménitére’s disease Zollinger-Ellison syndrome, pseudo-Zollinger-Ellison syndrome (hypersecretion of gastric HCl) Drugs (aspirin and ethanol), adrenocorticosteroids or nonsteroidal anti-inflammatory agents Sports anemia Patterson-Kelly (Plummer-Vinson) syndrome Facitia iron deficiency anemia (Lasthenie de Ferjol syndrome—self-induced bloodletting) Poor dietary intake Transferrin: severe protein-energy malnutrition Iron sequestration (idiopathic pulmonary hemosiderosis, paroxysmal nocturnal hemoglobinuria, chronic disease with inability to metabolize iron from reticulo-endothelial cell deposits, congenital atransferrinemia [rare]) Vitamin A deficiency—lack in developmental periods causes deficits in neural functioning and behavior

Lead (Pb)
Molybdenum (Mo)

(blood, preferred specimen)

2 mL, collect with oxalate-fluoride mixture

RR: <25 µg/dL or <0.21 mmol/L in children
and in most adults without occupational exposure

CR: = 100 µg/dL or 4.8 µmol/L in adults

(24-h urine)

RR: <80 µg/dL or <0.39 µmol/L (hair)

RR: <5 µg/g

CR: >2 µg/g

Collection:

Collect specimen in lead-free container and avoid airborne contaminants. For blood, use specifically manufactured tubes for blood lead collection.

Description:

Molybdenum is a metallic element—its compounds are poisonous, and any level of lead in blood is abnormal. Lead oxides are used in paint pigment; lead additives in gasoline provide air pollutants.

Interpretation:

Toxic: 3.0–294.4 mmol/L or RR: 0.6–59.0 µg/L

Children and adults, hypochromic microcytic anemia

Lead sources: Ingested or inhaled leaded paint (renovation dust), contaminated soil; contaminated water (lead pipes, lead solder on copper pipes, softened water); retention of a lead object in the stomach or joint (shot, curtain weight, fishing weight, bauble), contaminated acidic foods and beverages (storage in lead-glazed ceramics, leaded crystal, galvanized or nonstainless steel pots); inhalation (burning lead-painted wood or battery casings in home fireplaces or stoves); leaded gas fumes; occupational exposure

Lethal dose: 30 g/kg body weight

Use acid-washed, leakproof container, keep specimen on ice.

No metabolic role. Mercury is the only metal that is liquid at ordinary temperatures. Primarily absorbed by inhalation, but can also be absorbed through the skin and gastrointestinal tract. It is then distributed to the central nervous system and kidneys and excreted in the urine. Evaluated for mercury toxicity, neurologic findings related to inorganic or organic mercurials, inhalation of mercury vapors. Assess for occupational exposure, toxicity, and poisoning from contaminated fish.

Manganese (Mn)

Essential for lipid and carbohydrate metabolism, bone and tissue formation, and reproductive processes

Assess for occupational exposure and evaluate certain diseases.

Therapy for bipolar disorder

Diabetes insipidus

Renal failure, weight gain

Diminished taste perception

High “hard water” levels

Lithium (Li)

Ultratrace mineral; a metallic element

Lithium carbonate is used as drug to treat manic phase of manic-depressive illness. Decreased dietary sodium intake lowers the excretion rate of lithium. Assess psychotherapeutic drug monitoring.

Chronic inhalation of airborne manganese (mines, steel mills, chemical industries)

“Manganic madness,” permanent crippling neurologic disorder of the extrapyramidal system (similar to lesions in Parkinson's disease)

Increased urine levels in acute hepatitis, myocardial infarction and rheumatoid arthritis

Low tissue values in children with maple syrup disease and phenylketonuria

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RR: <5 µg/g

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Description:

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Lethal dose: 30 g/kg body weight

Use acid-washed, leakproof container, keep specimen on ice.

No metabolic role. Mercury is the only metal that is liquid at ordinary temperatures. Primarily absorbed by inhalation, but can also be absorbed through the skin and gastrointestinal tract. It is then distributed to the central nervous system and kidneys and excreted in the urine. Evaluated for mercury toxicity, neurologic findings related to inorganic or organic mercurials, inhalation of mercury vapors. Assess for occupational exposure, toxicity, and poisoning from contaminated fish.
<table>
<thead>
<tr>
<th>Element</th>
<th>Serum/Plasma</th>
<th>RR (µg/L or nmol/L)</th>
<th>Clinical Considerations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Molybdenum (Mo)</td>
<td>Serum: 0.1–3.0 or 1.0–31.3 nmol/L</td>
<td></td>
<td>A trace element, associated with the inborn error of molybdenum metabolism. Assess for genetic and dietary molybdenum deficiency.</td>
</tr>
<tr>
<td></td>
<td>Urine: 0.1–10 µg/d or 2–170 nmol/L</td>
<td></td>
<td>Massive ingestion of tungsten (W)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Occupational and high dietary intake (elevated uric acid blood concentration, gout)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sulfur amino acid toxicity</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Growth depression and anemia similar to copper deficiency</td>
</tr>
<tr>
<td>Nickel (Ni)</td>
<td>Serum or Plasma: 0.14–1.0 or 2.4–17.0 nmol/L</td>
<td></td>
<td>Ultratrace mineral; metallic element</td>
</tr>
<tr>
<td></td>
<td>Urine: 0.1–10 µg/d or 2–170 nmol/L</td>
<td></td>
<td>Nickel carbonyl is an industrial chemical used in plating metals—toxic when inhaled, causes pulmonary edema.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Assess for occupational exposure.</td>
</tr>
<tr>
<td>Selenium (Se)</td>
<td>Serum: 46–143 or 0.58–1.62 µmol/d</td>
<td></td>
<td>A chemical element resembling sulfur, found in soil. Has a role in the metabolism of enzymes. As a sulfide, used in treating dandruff and tinea versicolor (ie, Selsun Blue).</td>
</tr>
<tr>
<td></td>
<td>Urine: 6.0–15.0 mg/24 h or 214–534 µmol/d</td>
<td></td>
<td>Determine cause of loss of pigmentation of hair and skin.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Elevated dietary intake owing to high soil concentrations (North Dakota, USA; Venezuela).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>As a sulfide, used in treating dandruff and tinea versicolor (ie, Selsun Blue).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Parenteral nutrition</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Parenteral nutrition</td>
</tr>
<tr>
<td>Silicon (Si)</td>
<td>Plasma: 0.13–0.15 mg/L or 4.63–5.43 µmol/L</td>
<td></td>
<td>Ultratrace mineral; nonmetallic element in soil. Occurs in traces in skeletal structures (bones and teeth).</td>
</tr>
<tr>
<td></td>
<td>Urine: 6.0–15.0 mg/24 h or 214–534 µmol/d</td>
<td></td>
<td>Necessary for the formation of collagen, bones, and connective tissue; healthy nails, skin and hair; and calcium absorption in early stages of bone formation. Needed to maintain flexible arteries and major role in cardiovascular disease. Important in prevention of Alzheimer’s disease and osteoporosis; inhibits aging process in tissues.</td>
</tr>
<tr>
<td>Silver (Ag)</td>
<td>Serum: 0.21 ± 0.15 ng/dL, 19.47 ± 13.90 nmol/L</td>
<td></td>
<td>Collect in metal-free container. No metabolic role. Salts used as antiseptic and bacteriostatic agents. In normal individuals, silver slowly accumulates in body tissue with age but causes no apparent harm.</td>
</tr>
<tr>
<td></td>
<td>Urine: &lt;1 µg/d or &lt;9.3 nmol/d</td>
<td></td>
<td>Assess for occupational exposure or toxicity from medicinal uses of silver.</td>
</tr>
<tr>
<td>Thallium (Tl)</td>
<td></td>
<td></td>
<td>Lethal dose: 3.5–35 g total dose</td>
</tr>
<tr>
<td>Element</td>
<td>Range (blood)</td>
<td>Range (urine)</td>
<td>Collect</td>
</tr>
<tr>
<td>---------</td>
<td>---------------</td>
<td>--------------</td>
<td>---------</td>
</tr>
<tr>
<td>RR: &lt;0.5 µg/dL or &lt;24.5 nmol/L</td>
<td>CR: 10–800 µg/dL or 0.5–39.1 µmol/L</td>
<td>RR: &lt;2 µg/L or &lt;9.8 nmol/L or 4.9–97.8 µmol/L</td>
<td>Collect in metal-free container.</td>
</tr>
<tr>
<td>Tin (Sn)</td>
<td>RR: 24–50 µg/L or 202–421 nmol/L</td>
<td></td>
<td>Collect in metal-free container.</td>
</tr>
<tr>
<td>Vanadium (V)</td>
<td>Serum: RR: 0.01–0.23 µg/L or 0.20–4.51 nmol/L or Hair: RR: 0.10–0.16 µg/g dry wt. or 1.96–3.14 nmol/g dry wt.</td>
<td>Urine: RR: &lt;0.24 µg/L or &lt;4.7 nmol/L</td>
<td>Collect in metal-free container.</td>
</tr>
<tr>
<td>Zinc (Zn)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Zinc**

- **Fasting morning specimen**
  - Plays a role in protein synthesis; critical for growth and sexual maturation
  - Important in wound healing and sensory perception (particularly taste and smell)
  - Important in activating certain serum enzymes and in insulin and porphyrin metabolism
  - Assess population that may have increased needs for intake—alcoholism, chronic illness, stress, trauma, surgery, mal-absorption, lactovegetarians, children consuming vegetarian diets, decubitus ulcers, anorexics.

- **24-h urine**
  - RR: 70–120 µg/dL or 10.7–18.4 µmol/L
  - RR: 150–1200 µg/d or 2.3–18.4 µmol/d
  - Zinc therapy for Wilson's disease
  - Ingestion of food or beverage contaminated by storage in a galvanized container
  - Long-term ingestion of excessive zinc supplements >150 mg/d (secondary copper deficiency)
  - Low serum high-density lipoprotein
  - Gastric erosion
  - Depressed immune system
  - Lethargy in dialysis patients
  - Hyperzincuria increasing with the severity of diabetes
  - Inhalation of zinc oxide fumes causing neurologic damage (metal fume fever, brass-founders' ague, zinc shakes), metallic taste, bloody diarrhea
  - Decreased intake (chronic alcoholics, vegetarians, young women with anorexia nervosa), diarrhea
  - Decreased circulatory and splenic T lymphocytes
  - Prolonged bedrest
  - Decrease in absorption of tetracycline
  - Rheumatic diseases
  - Infection
  - Growth retardation
  - Male hypogonadism and hypospermism
  - Nyctalopia (night blindness)
  - Hypogeusia (blunting of sense of taste)
  - Impaired wound healing, long-term total parenteral nutrition without zinc supplement
  - Chronic liver disease
  - Acrodermatitis enteropathica (dermatitis of the extremities seen in children between 3 weeks–18 months)
  - Dwarfism
  - Parasitism (Egypt)
  - Compromised immune function
  - Low facteur thyrique sérique (FTS) (thyrmulin, hormone secreted by the thymus)
  - Impaired embryogenesis
  - Behavioral disturbances (impaired hedonic tone)
  - Skeletal abnormalities, defective collagen synthesis, alopecia, impaired protein synthesis
  - Some cancers
Appendix H Examples of Forms

The health care profession assists patients to properly use and complete required forms as part of their pretest, intratest, and posttest care. These forms are part of the documentation and recording process and, in some cases, may become part of the permanent health care record. Accurately documenting health care diagnostic procedures may be necessary for reimbursement, legal, or compliance issues. For example, an informed consent form signifies a freely given agreement on the part of the patient to undergo certain diagnostic, surgical, or treatment procedures. Before signing any form, the patient, guardian, or witness should carefully read the entire form and ask questions if not thoroughly understood. If the patient cannot read and sign a given form, documentation of a qualified individual (e.g., parent or guardian) who is signing the form should be recorded. Anything that occurs out of the ordinary when preparing, administering, explaining or signing a form should also be documented.

Forms included in this appendix are used for HIV testing, drug screening, sleep logs and questionnaires, evidentiary specimen collection, sexual assault examination, interviewing/ videotaping a patient, and molecular genetic testing. These are not all-inclusive but provide examples of commonly used forms.

**FIGURE H.1** State of Maryland Health Department form for HIV testing.

**FIGURE H.2** Consent form for witnessed urine drug testing in community, occupational, or clinic setting.

**FIGURE H.3** Sleep log.

FIGURE H.5 Sleep disorders questionnaire.


FIGURE H.7 Male evidence collection. Reference: Olshaker JS, Jackson MC, Smock WS: Forensic Emergency Medicine,


**FIGURE H.10** Consent form for interviewing, videotaping, and/or photographing a patient. Reference: Olshaker JS, Jackson MC, Smock WS: Forensic Emergency Medicine, Philadelphia, Lippincott Williams & Wilkins, 2001 (modified).
**FIGURE H.11** Example of a consent form used by neighborhood nurses in public health offices prior to testing clients for STDs.

**FIGURE H.12** Example of a consent form used for sickle cell anemia testing program.

**FIGURE H.13** Non-consenting or deceased patient authorization form for HIV testing, State of Wisconsin, USA.

**FIGURE H.14** Consent for molecular genetic testing. Source: CompGene, Comprehensive Genetics Services, S.C., Milwaukee, WI, USA.
Appendix I Panic or Critical Values

It is recognized that critical laboratory values (eg, low or high critical value) indicate the need for prompt clinical intervention. The health care professional reviews records for any sudden change in values that may also signal alarm (eg, new diagnoses of leukemia, sickle cell anemia, aplastic crisis). Critical (panic) laboratory values represent serious medical conditions that may be life threatening unless immediate actions are taken. Notification and collaboration with the clinician and other members of the health care team must take place when critical values are identified so that prompt treatment can begin (eg, a blood glucose level <70 or >300 mg/dL [<4 or >17 mmol/L], an increased bleeding time >15 minutes], a hemoglobin <7.0 or >20.0 g/dL [<70 or >200 g/L], a brown to black-gray urine upon standing with an increasing pH, or an elevated digoxin =2.0 mg/mL [>2.6 mmol/L]). (See Table I.1).

<table>
<thead>
<tr>
<th>Test</th>
<th>Low Critical Value</th>
<th>SI Units</th>
<th>High Critical Value</th>
<th>SI Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lead/capillary blood</td>
<td>0–&lt;9 µg/dL (RR)</td>
<td>0–&lt;0.43 µmol/L</td>
<td>&gt;45 µg/dL</td>
<td>&gt;2.17 µmol/L</td>
</tr>
<tr>
<td>Antibody identification</td>
<td>Positive</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cord blood workup</td>
<td>Positive direct Coombs</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Absolute neutrophil count</td>
<td>=500 × 10³/mm³</td>
<td>=500,000/mm³</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hematocrit</td>
<td>&lt;20%</td>
<td>&lt;0.20</td>
<td>&gt;60%</td>
<td>&gt;0.60</td>
</tr>
<tr>
<td>Hematocrit (neonate)</td>
<td>&lt;14.7 mmol/L</td>
<td></td>
<td>&gt;70 vol %</td>
<td>&gt;31.2 mmol/L</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>&lt;9.5 g/dL</td>
<td>&lt;95 g/dL</td>
<td>&gt;22 g/dL</td>
<td>&gt;220 g/L</td>
</tr>
<tr>
<td>Hemoglobin (neonate)</td>
<td>&lt;2 × 10⁹/mm³</td>
<td>&lt;2000/mm³</td>
<td>&gt;30 × 10³/mm³</td>
<td>&gt;30,000/mm³</td>
</tr>
<tr>
<td>Platelets, adult</td>
<td>&lt;40 × 10³/mm³</td>
<td>&lt;40,000/mm³</td>
<td>&gt;10,000 × 10³/mm³</td>
<td>&lt;1,000,000/mm³</td>
</tr>
<tr>
<td>Activated partial thromboplastin time (APTT)</td>
<td>None</td>
<td></td>
<td>&gt;78 seconds</td>
<td></td>
</tr>
<tr>
<td>Fibrinogen</td>
<td>&lt;100 mg/dL</td>
<td>&lt;2.9 µmol/L</td>
<td>&gt;700 mg/dL</td>
<td>&gt;20.6 µmol/L</td>
</tr>
<tr>
<td>Prothrombin time (PT)</td>
<td>None</td>
<td></td>
<td>INR: &gt;3.6 or 730 sec or 3 × control</td>
<td></td>
</tr>
<tr>
<td>Bleeding Time</td>
<td>None</td>
<td></td>
<td>&gt;15 minutes</td>
<td></td>
</tr>
<tr>
<td>NOTE exceptions:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hematocrit high critical value</td>
<td></td>
<td></td>
<td>&gt;60 does not apply to newborns.</td>
<td></td>
</tr>
<tr>
<td>Hemoglobin high critical value</td>
<td></td>
<td></td>
<td>&gt;20 does not apply to newborns.</td>
<td></td>
</tr>
</tbody>
</table>

The presence of blast cells and sickle cells is to be reported at once.

CHEMISTRY

<table>
<thead>
<tr>
<th>Test</th>
<th>Low Critical Value</th>
<th>SI Units</th>
<th>High Critical Value</th>
<th>SI Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bilirubin, adult</td>
<td>None</td>
<td>None</td>
<td>&gt;15 mg/dL</td>
<td>&gt;257 µmol/L</td>
</tr>
<tr>
<td>Bilirubin, neonate</td>
<td>None</td>
<td>None</td>
<td>&gt;13 mg/dL</td>
<td>&gt;222 µmol/L</td>
</tr>
<tr>
<td>BUN</td>
<td>2 mg/dL</td>
<td>0.71 mmol/L</td>
<td>&gt;80 mg/dL</td>
<td>&gt;28.6 mmol/L</td>
</tr>
<tr>
<td>Calcium</td>
<td>&lt;6 mg/dL</td>
<td>&lt;1.5 mmol/L</td>
<td>&gt;13 mg/dL</td>
<td>&gt;3.2 mmol/L</td>
</tr>
<tr>
<td>Carbon dioxide (CO₂)</td>
<td>&lt;10 mEq/L</td>
<td>&lt;10 mmol/L</td>
<td>&gt;40 mEq/L</td>
<td>&gt;40 mmol/L</td>
</tr>
<tr>
<td>Creatinine</td>
<td>0.4 mg/dL</td>
<td>35 µmol/L</td>
<td>2.8 mg/dL</td>
<td>247 µmol/L</td>
</tr>
<tr>
<td>Glucose, adult</td>
<td>&lt;70 mg/dL</td>
<td>&lt;3.9 mmol/L</td>
<td>&gt;300 mg/dL</td>
<td>&gt;16.6 mmol/L</td>
</tr>
<tr>
<td>Glucose, neonate</td>
<td>&lt;30 mg/dL</td>
<td>&lt;1.7 mmol/L</td>
<td>&gt;325 mg/dL</td>
<td>&gt;18 mmol/L</td>
</tr>
<tr>
<td>Magnesium</td>
<td>&lt;1.0 mg/dL</td>
<td>&lt;0.4 mmol/L</td>
<td>None</td>
<td>&gt;1.0 mmol/L</td>
</tr>
<tr>
<td>Osmolality</td>
<td>250 mOsm/kg</td>
<td>250 mmol/kg</td>
<td>&gt;325 mOsm/kg</td>
<td>325 mmol/kg</td>
</tr>
<tr>
<td>Phosphorus</td>
<td>1.2 mg/dL</td>
<td>0.4 mmol/L</td>
<td>9 mg/dL</td>
<td>2.87 mmol/L</td>
</tr>
<tr>
<td>Potassium, adult</td>
<td>&lt;2.8 mEq/L</td>
<td>&lt;2.8 mmol/L</td>
<td>&gt;6.7 mEq/L</td>
<td>&gt;6.7 mmol/L</td>
</tr>
<tr>
<td>Potassium, neonate</td>
<td>&lt;2.8 mEq/L</td>
<td>&lt;2.8 mmol/L</td>
<td>&gt;7.0 mEq/L</td>
<td>&gt;7.0 mmol/L</td>
</tr>
<tr>
<td>Sodium</td>
<td>&lt;120 mEq/L</td>
<td>&lt;120 mmol/L</td>
<td>&gt;160 mEq/L</td>
<td>&gt;160 mmol/L</td>
</tr>
<tr>
<td><strong>HEMODIALYSIS</strong></td>
<td><strong>Calcium</strong></td>
<td>&lt;8 mg/dL</td>
<td>&lt;2 mmol/L</td>
<td>&gt;11 mg/dL</td>
</tr>
<tr>
<td><strong>Glucose</strong></td>
<td>&lt;70 mg/dL</td>
<td>&lt;3.9 mmol/L</td>
<td>&gt;300 mg/dL</td>
<td>&gt;16.6 mmol/L</td>
</tr>
<tr>
<td><strong>Hematocrit</strong></td>
<td>&lt;20%</td>
<td>0.20</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td><strong>Hemoglobin</strong></td>
<td>&lt;6.0 g/dL</td>
<td>&lt;60 g/L</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td><strong>Phosphorus</strong></td>
<td>&lt;2.0 mg/dL</td>
<td>0.71 mmol/L</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td><strong>Potassium</strong></td>
<td>&lt;2.8 mEq/L</td>
<td>&lt;2.8 mmol/L</td>
<td>&gt;6.5 mEq/L</td>
<td>&gt;6.5 mmol/L</td>
</tr>
</tbody>
</table>

**BLOOD GASES**

### Arterial
- **CO (carbon monoxide)**
  - None
  - None
  - >10%
  - >0.10
- **HCO₃ bicarbonate**
  - <10 mEq/L
  - <10 mmol/L
  - >40 mmol/L or mEq/L
  - >40 mmol/L
- **Methemoglobin**
  - None
  - None
  - >10%
  - >0.10
- **O₂ content**
  - >9.0 vol %
  - <4.0 mmol/L
  - None
  - None
- **O₂ saturation**
  - <75%
  - <0.75
  - None
  - None
- **pCO₂**
  - <20 mm Hg
  - <2.7 kPa
  - >77 mm Hg
  - >10.2 kPa
- **pH**
  - <7.20
  - <7.20
  - >7.60
  - >7.60
- **pO₂**
  - <40 mm Hg
  - <5.3 kPa
  - None
  - None
- **Potassium**
  - <2.5 mEq/L
  - <2.5 mmol/L
  - >6.5 mEq/L
  - >6.5 mmol/L
- **Sodium**
  - <120 mEq/L
  - <120 mmol/L
  - >160 mEq/L
  - >160 mmol/L

### Venous
- **HCO₃**
  - <10 mEq/L
  - <10 mmol/L
  - >40 mEq/L
  - >40 mmol/L
- **Saturation**
  - <40%
  - <0.40
  - >85%
  - >0.85
- **pCO₂**
  - <20 mm Hg
  - <2.7 kPa
  - >60 mm Hg
  - >8.0 kPa
- **pH**
  - <7.20
  - <7.20
  - >7.60
  - >7.60

### Capillary
- **pCO₂**
  - <20 mm Hg
  - <2.7 kPa
  - >60 mm Hg
  - >8.0 kPa
- **pH**
  - <7.20
  - <7.20
  - >7.60
  - >7.60
- **pO₂**
  - <25 mm Hg
  - <3.3 kPa
  - None
  - None
- **Total T₃**
  - <50 ng/dL
  - <0.8 nmol/L
  - >300 ng/dL
  - >4.6 nmol/L
- **Total T₄**
  - <2.0 µg/dL
  - <25.7 nmol/L
  - >15.0 µg/dL
  - >193 nmol/L

### MICROBIOLOGY (QUALITATIVE RESULTS) AND SEROLOGY

| Blood culture/smear growth | None | Positive—heavy of pathogen |
| CSF culture/smear Gram's stain | None | Positive |
| Bacterial antigens | None | Antigen detected |
| Blood cross match | None | Incompatible |
| Blood parasites—malaria | None | Present |
| Stool culture/smear | None | Enteric pathogen present. |
| Clostridium difficile toxin (Inpatients and residents of long-term care facilities) | None | Positive |
| Cryptosporidium antigen | None | Positive |
| Culture, Group B Strep, genital (delivery and nursery) | None | Positive |
| Culture/Smear, acid-fast bacilli | None | Positive |
| Culture/Smear, Legionella E. coli K1 antigen | None | Positive |
| Enteric pathogens | None | Positive |
| Giardia antigen | None | Positive |
| Haemophilus antigen | None | Positive |
| Legionella antigen | None | Positive |
| Meningococcal antigen | None | Positive |
| Methicillin-resistant | None | Positive |
| Staphylococcus aureus | None | Positive |
| Ova and parasites | None | Positive |
| Pneumococcal antigen | None | Positive |
| Pneumocystis carinii DFA | None | Positive |
| Strep Group B antigen (delivery and nursery) | None | Positive |
| Strep Group B antigen (spinal fluid) | None | Positive |
| Vancomycin-resistant Enterococcus | None | Positive |

### SPECIFIC ORGANISMS—BLOOD
- **Tuberculosis stain**
  - None
  - Positive acid-fast or culture
<table>
<thead>
<tr>
<th>Test</th>
<th>Normal Range</th>
<th>Abnormality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hepatitis</td>
<td>None</td>
<td>Positive</td>
</tr>
<tr>
<td>Syphilis</td>
<td>None</td>
<td>Positive</td>
</tr>
<tr>
<td>AIDS</td>
<td>None</td>
<td>Positive</td>
</tr>
<tr>
<td>Malaria smear</td>
<td>None</td>
<td>Positive</td>
</tr>
<tr>
<td><strong>SPINAL FLUID (CSF)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CSF—total protein</td>
<td>Low–none</td>
<td>High: &gt;45 mg/dL</td>
</tr>
<tr>
<td>CSF—glucose</td>
<td>Low &lt;80% of blood level</td>
<td></td>
</tr>
<tr>
<td>CSF—WBC</td>
<td>Low–none</td>
<td>High: increased over 20 segmented neutrophils</td>
</tr>
<tr>
<td>CSF—blasts or malignant cells</td>
<td>Low–none</td>
<td>High: &gt;10/mm$^3$</td>
</tr>
<tr>
<td><strong>URINALYSIS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urine, micro</td>
<td>None</td>
<td>Pathologic crystals (urate, cysteine, leucine, tryptine)—present</td>
</tr>
<tr>
<td>Urine RBC casts, casts micro</td>
<td>None</td>
<td>Presence of RBC</td>
</tr>
<tr>
<td>Urine glucose, macro</td>
<td>None</td>
<td>Strongly positive</td>
</tr>
<tr>
<td>Urine ketone, macro</td>
<td>None</td>
<td>Strongly positive</td>
</tr>
</tbody>
</table>

**NOTE:** There are variance in unpublished lists—this is not an official list.

**NOTE EXCEPTIONS:**
- Urine glucose in preop patient
- Urine glucose in patient <2 years
- Urine ketones in patient <2 years
- Urine glucose in patient <2 years
- >250 mg/dL
- >14 mmol/L
Appendix J Effects of the Most Commonly Used Drugs on Frequently Ordered Laboratory Tests (Blood, “Whole” Plasma, Serum, Stool, and Urine)

**DRUGS AND LABORATORY TEST OUTCOMES**

Many prescription medications, over-the-counter medications, vitamins, minerals, and herbal preparations can influence the results of laboratory tests. Mechanisms of drug effects are either pharmacologic (eg, furosemide usually increases the excretion of potassium, resulting in a low serum potassium level) or analytic (eg, when a drug in a patient's body fluid or tissue interferes with a chemical step in a laboratory test, resulting in an erroneous test result). The drug classes that cause the majority of analytical interferences include antibiotics, antihypertensives, anticonvulsants, hormones, and antidepressants.

Accurate and complete medication histories, including prescription medications, over-the-counter medications, vitamins, minerals, and herbal preparations, are essential to interpret laboratory test results that fall out of the normal range.

The following table (Table J.1) is by no means exhaustive, and whenever laboratory results are suspected of being spurious, further research is necessary. There are many references available, including drug monographs in *American Hospital Formulary Service* (AFHS), *Drug Information* (published by the American Society of Health System Pharmacists), drug package inserts, and the *Physician's Desk Reference* (PDR), which contains official product information. Services that maintain this information include the Iowa Drug Information Service (IDIS) and DRUGDEX.

### Discoloration of Feces Caused by Drugs

1. **Black**—acetazolamide, aluminum hydroxide, aminophylline, 5-aminosalicylic acid, amphetamine, amphotericin B, antacids, anticoagulants, aspirin, betamethasone, bismuth, charcoal, chloramphenicol, chlorpropamide, clindamycin, corticosteroids, cortisone, cyclo-phosphamide, cytarabine, digitalis, ethacrynic acid, ferrous salts, furosemide, fluoroquinolones, halothane, heparin, hydralazine, hydrocortisone, ibuprofen, indomethacin, iodine, levarterenol, levodopa, manganese, melphalan, methotrexate, methylene blue, paraldehyde, phenacetin, phenolphthalein, potassium, potassium salts, prednisolone, procarbazine, senna, sulfonamides, tetracycline, theophylline, thiotepa, trimethaphan, warfarin
2. **Gray**—colchicine
3. **Green**—indomethacin, iron, mebendazole, mesalamine, naphthalene
4. **Dark brown**—dexamethasone
5. **Blue**—chloramphenicol, methylene blue
6. **Pink**—anticoagulants, aspirin, salicylates
7. **Red**—anticoagulants, aspirin, phenolphthalein, salicylates, tetracycline
8. **Orange**—phenazopyridine, rifampin
9. **Tarry**—warfarin, ergot preparations, ibuprofen, salicylates
10. **White/speckling**—aluminum hydroxide, indocyanine green
11. **Yellow**—senna

### Discoloration of Urine Caused by Drugs

1. **Black**—cascara, co-trimoxazole, ferrous salts, levodopa, methocarbamol, methylxypyrrolidone, naphthalene, quinine, sulfonamides
2. **Dark**—cascara, levodopa, metronidazole, primaquine, quinine, senna
3. **Brown**—cascara, chloroquine, levodopa, methocarbamol, methyldopa, metronidazole, nitrofurantoin, primaquine, quinine, rifampin, senna, sulfonamides
4. **Blue**—indigo blue, methylene blue, mitoxantrone, nitrofurantoin, triamterene
5. **Blue/green**—amitriptyline, Doan's pills, indigo blue, indomethacin, magnesium salicylate, methylene blue, propofol
6. **Orange**—chloroxazone, dihydroergotamine, heparin, phenazopyridine, rifampin, sulfasalazine, warfarin
7. **Pink**—anthraquinone dyes, aspirin, cascara, deferoxamine, methyldopa, phenytin, salicylates, senna
8. **Purple**—phenolphthalein
9. **Red**—cascara, chlorpromazine, daunorubicin, deferoxamine, dihydroergotamine, dimethylsulfoxide, doxorubicin, heparin, ibuprofen, methyldopa, phenazopyridine, phenolphthalein, phenothiazines, phenytin, rifampin, senna
10. **Red-purple**—chloroxazone, ibuprofen, senna

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**BIBLIOGRAPHY**

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### Discoloration of Feces Caused by Drugs

1. **Black**—acetazolamide, aluminum hydroxide, aminophylline, 5-aminosalicylic acid, amphetamine, amphotericin B, antacids, anticoagulants, aspirin, betamethasone, bismuth, charcoal, chloramphenicol, chlorpropamide, clindamycin, corticosteroids, cortisone, cyclo-phosphamide, cytarabine, digitalis, ethacrynic acid, ferrous salts, furosemide, fluoroquinolones, halothane, heparin, hydralazine, hydrocortisone, ibuprofen, indomethacin, iodine, levarterenol, levodopa, manganese, melphalan, methotrexate, methylene blue, paraldehyde, phenacetin, phenolphthalein, potassium, potassium salts, prednisolone, procarbazine, senna, sulfonamides, tetracycline, theophylline, thiotepa, trimethaphan, warfarin
2. **Gray**—colchicine
3. **Green**—indomethacin, iron, mebendazole, mesalamine, naphthalene
4. **Dark brown**—dexamethasone
5. **Blue**—chloramphenicol, methylene blue
6. **Pink**—anticoagulants, aspirin, salicylates
7. **Red**—anticoagulants, aspirin, phenolphthalein, salicylates, tetracycline
8. **Orange**—phenazopyridine, rifampin
9. **Tarry**—warfarin, ergot preparations, ibuprofen, salicylates
10. **White/speckling**—aluminum hydroxide, indocyanine green
11. **Yellow**—senna

### Discoloration of Urine Caused by Drugs

1. **Black**—cascara, co-trimoxazole, ferrous salts, levodopa, methocarbamol, methylxypyrrolidone, naphthalene, quinine, sulfonamides
2. **Dark**—cascara, levodopa, metronidazole, primaquine, quinine, senna
3. **Brown**—cascara, chloroquine, levodopa, methocarbamol, methyldopa, metronidazole, nitrofurantoin, primaquine, quinine, rifampin, senna, sulfonamides
4. **Blue**—indigo blue, methylene blue, mitoxantrone, nitrofurantoin, triamterene
5. **Blue/green**—amitriptyline, Doan's pills, indigo blue, indomethacin, magnesium salicylate, methylene blue, propofol
6. **Orange**—chloroxazone, dihydroergotamine, heparin, phenazopyridine, rifampin, sulfasalazine, warfarin
7. **Pink**—anthraquinone dyes, aspirin, cascara, deferoxamine, methyldopa, phenytin, salicylates, senna
8. **Purple**—phenolphthalein
9. **Red**—cascara, chlorpromazine, daunorubicin, deferoxamine, dihydroergotamine, dimethylsulfoxide, doxorubicin, heparin, ibuprofen, methyldopa, phenazopyridine, phenolphthalein, phenothiazines, phenytin, rifampin, senna
10. **Red-purple**—chloroxazone, ibuprofen, senna

---

**Table J.1 Effects of the Most Commonly Used Drugs on Frequently Ordered Laboratory Tests**

<table>
<thead>
<tr>
<th>Test</th>
<th>Increased Can Lead to a False-Positive Value</th>
<th>Decreased Can Lead to a False-Negative Value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Acid Phosphatase (Serum)  
Increased by: p-aminosalicylic acid

Decreased by: clofibrate, mirtazapine, pemoline

Alanine Aminotransferase  
Increased by: acetobutol, acetaminophen, acetohexamide, acyclovir, albendazole, aldesleukin, allorpinol, alprazolam, aminoglutethimide, aminosalicylic acid, amiodarone, amitriptyline, amoxapine, amphotericin B, ampicillin, amrinone, anabolic steroids, anastrozole, anticonvulsants, antifungal agents, ardeparin, arsenic trioxide, asparaginase, aspirin, atorvastatin, atovaquone, auranofin, aurothioglucose, azathioprine, aztreonam, barbiturates, barium, BCG vaccine, benazepril, bepridil, betaxolol, bicatulamidine, bismuth subsalicylate, bisoprolol, bitolterol, bromocriptine, buspiron, busulfan, calcitriol, candesartan, capecitabine, carbenicillin, carmustine, cephalosporin antibiotics, cervatentin, cetirizine, chenodiol, chloral hydrate, chlorambucil, chloramphenicol, chloridiazepoxide, chlorothiazide, chlorpheniramine, chlorpromazine, chloropropamide, chlorotetracycline, chlortaldione, chlorozoxazine, cholestyramine, choline magnesium trisalicylate, cidofovir, cimetidine, cinocicin, ciprofloxacin, cisplatin, cladribine, clarihyalin, clindamycin, clofazimine, clofibrate, clophene, clopimargine, clonidine, clopidogrel, clobazapate, cloxacinil, clozapine, copepilin, conjugated estrogens, cortisone, cyclobenzaprine, cyclophosphamide, cyproheptadine, cytarabine, dactinomycin, dalte-parin, danazol, dantrolene, dapsone, demeclocycline, desipramine, dizapam, diclofenac, didanosine, dienestrol, diethylstilbestrol, diffusional, diltiazem, disopyramide, disulfiram, doxetacel, doxorubicin, doxycycline, dronabinol, enalapril, enoxaparin, erythromycin, estersified estrogens, estropipate, ethacrynic acid, ethambutol, etchilorynol, ether, etodolac, etoposide, etretinates, famotidine, farnamate, fenofibrate, fenoprofen, fencaineide, flucanoan, flucytoseine, fluorouracil, fluoxymesterone, fluphenazine, flurazepam, flutamide, fluvastatin, fluvaxetine, foscarnet, fospencytoin, furazolidone, furosemide, ganciclovir, gemfibrozil, gentamicin, glemipime, glycopyrrolate, gold, gosferelin, granisetron, griseofulvin, guanethidine, haloperidol, heptatis A vaccine, heparins, lomefloxacin, loracarbef, loratadine, lovastatin, loxapine, MAO inhibitors, mepatoline, mephitritamine, mefloquine, melphalan, meserase, mepramabate, mercaptopurine, mepenoxan, mesalamine, methotrexate, methoxsalen, methyldopa, methylphenidate, methyltestosterone, metoclopramide, metotolozane, metoprolol, mexiletine, minocycline, mirtazapine, mitomycin, mitoxantrone, moexiplol, molindone, montelukast, moricizine, morphine, moxalactam, muromonob-CD3, mycophenolate, nabumetone, nafarelin, nacillin, nalidixic acid, naltrexone, nadrolone, naproxen, nefazodone, nefilina, netilmicin, nepitavrin, niasicin, niasicinamide, nicardipine, nifedipine, nilutamide, nisoldipine, nitrofurantoin, nifedipine, northeptrolone, norfaxacin, norotyplite, octreotide, ofloxacine, oleandomycin, olsalazine, omeprazole, ondansetron, oral contraceptive, oxacinil, oxaprin, oxazepam, oxazepemone, palivizumab, papaverin, paraglyne, paroxetine, pegasparagase, pemoline, penicillamine, pentoxyllylifene, perphenazine, phenazopyridine, phenezilin, phenobarbital, phenothiazines, phenytoin, phosphoryl, phosphorus, pindolol, piroglitazone, pipercalin, pirroricam, polithyazine, pralidoxine, pravastatin, prazosin, probenecid, procainamide, prochlorperazine, propafenone, propoxyphene, propranolol, propritopril, protropilin, pyrazinamide, pyrtrimithine, quazepam, quinapril, quinethazone, quinidine, ramipril, ranitidine, rifampin, rufine, rupidone, ritonavir, rosiglitazone, saquinivir, sargramostimustin, sibutramine, sildenafil, simvastatin, sparflxacin, spectinomycin, stanozolol, stavudine, streptokinase, streptomycin, streptozocin, sulfadiazine, sulfamethoxazole, sulfanilamide, sulfasalazine, sulfonamoxo, sulfonyleurea, sulindac, sumatriptan, tetracilum, terbutaline, terbutalil, tercyclazine, thiabendazole, thiazides, theilhyperazine, theicyanate, thiguanine, thiopenal, thiouracil, thiocyanate, thiopental, thioridazine, thiophixene, thioracil, ticarcillin, ticlopidine, timolol, tinzaparin, tobramycin, tocinidine, tolazamide, tolazoline, tobutamide, tocolpine, tometin, trandolapril, traloylpyrphenone, trastuzumab, treino, trichlorothiazide, trifluoperazine, trimethoprim, trimetrexate, trimipramine, trioxsalen, triproterol, triglazitone, troleando-myacin, trovafloxacin, uracil mustard, ursodiol, valproic acid, valpoxetine, verapamil, vidarabine, warfarin, zalcitabine, zidovudine, zileuton, zolmitriptan, zolpidem

Decreased by: aspirin, carvedilol, cycloxygenase, interferon, ketoprofen, phenothiazines, simvastatin, toremifene, ursodiol

Alanine Aminotransferase (Serum)  
Increased by: amikacin, basiliximab, calcitriol, carbiamaezone, carbediol, cispilatin, diazoxide, doxorubicin, gentamicin, lansoprazole, lithium, mesalamine, mycophenolate, nabumetone, naproxen, nifedipine, norfloxacin, ofloxacine, oral contraceptives, radioactive agents, sevoflurane, triazolam, venlafaxine, verapamil, zalcitabine

Decreased by: atenolol, captopril, cilostazol, dipyrismonde, enalapril, fosinopril, furosemide, ibuprofen, indapamide, metoprolol, quinapril, ranipril

Aldolase (Serum)  
Increased by: corticotropin, diclofenac, itraconazole, lovastatin, nacian, quinidine (IM injection), simvastatin, thiobendazole, vasopressin

Decreased by: propranolol

Aldosterone (Plasma)  
Increased by: amiloride, ammonium chloride, chlorathaldione, corticotropin, dobutamine, fenoldopam, fosinopril, furosemide, hydatazine, hydrochlorothiazide, indomethacin, laxesatives, metoclopramide, nifedipine, opiate, pravastatin, spironolactone, triamterene, verapamil

Decreased by: probucol
**Alkaline Phosphatase (Serum)**

**Increased by:**
- Ammonia (Plasma)
- Alkaline Phosphatase
- MAO inhibitors, neomycin, tetracycline, tromethamine

**Decreased by:**
- tetracycline, thiazides, valproic acid
- felbamate, furosemide, hydroflumethiazide, ion exchange resins, isoniazid, pegasparagase,
- parathyroid extract, phenobarbital, phenylephrine, phenylpropanolamine, primidone, ProSobee,
- levodopa, mafenide, methamphetamine, methyldopa, neomycin, nystatin, penicillamine,
- erythromycin, gentamicin, hydrocortisone, ifosfamide, isomil, insulin, kanamycin, levarterenol,
- cephradine, cisplatin, cloxacillin, colistin, corticotropin, cytarabine, doxorubicin, ephedrine,
- bismuth subsalicylate, brompheniramine, carbenicillin, carbenicilllin, cefaclor, cefadroxil, cephalexin,
- increased by:
- prednisolone, prednisone, tamoxifen, trifluoperazine, ursodiol, vitamin D
- danazol, estrogens, etidronate, norethindrone, oral contraceptives, pamidronate, penicillamine,
- azathioprine, calcitonin, calcitriol, carvedilol, chemotherapy, clofibrate, colchicine, cyclosporine,
- Decreased by:
- valproic acid, venlafaxine, verapamil, vidarabine, vitamin D, warfarin, zalcitabine, zidovudine,
- trimipramine, trioxsalen, troglitazone, troleandomycin, trovafloxacin, uracil mustard, ursodiol,
- tocainide, tolazamide, tolbutamide, tolcapone, tolmetin, toremifene, tramadol,
- thioguanine, thiopental, thioridazine, thiothixene, thiouracil, ticarcillin, ticlopidine, timolol,
- sulindac, tacrolimus, terbinafine, tetracycline, thiabendazole, thiazides, thiethylperazine,
- sulfadiazine, sulfamethoxazole, sulfanilamide, sulfasalazine, sulfisoxazole, sulfonyleureas,
- rifampin, nitric oxide, nicardipine, nifedipine, nitroglycerin, nitrofurantoin, nifedipine,
- fluphenazine, flurazepam, flutamide, fuflavastatin, foscarnet, fosphenytoin, ganciclovir,
- gem-citabine, gemfibrozil, gentamicin, glupyrrole, gold, granulocyte colony-stimulating factor, griseofulvin, haloperidol, hepatitis A vaccine, hepatitis B vaccine, human growth hormone, hydralazine, hydroflumethiazide, ibuprofen, idarubicin, ilutaside, indomethacin, interferon, interferon, interleukin, intravenous, isoniazid, isradipine, lacosamide, lamotrigine, levodopa, levotyroxine, lincomycin, lisinopril, lithium, lomefloxacin, loracarbef, loratadine, lovastatin, MAO inhibitors, mercloethromycin, meclofenamate, medroxyprogesterone, melphalan, meprobamate, mercaptopurine, meropenem, mesalamine, metaxalone, methimazole, methotrexate, methoxsalen, methylxylon, methyltestosterone, metoclopramide, metolazone, metoprolol, minocycline, mirtazapine, misoprostol, mitoxantrone, moexipril, molindone, morphine, moxalactam, mycophenolate, nabumetone, nafarelin, naladixic acid, nadrothine, naproxen, neflinavir, netilmicin, niacin, niacinamide, nicardipine, nifedipine, nitrofurantoin, nizatidine, norfloxacin, nortriptyline, octreotide, olofoxacin, oleandomycin, osalazine, omeprazole, oral contraceptives, oral hypoglycemics, oxacillin, oxaprinon, oxazapam, oxymetholone, papaverine, pargylene, paroxetine, pegaspargase, penicillamine, perphenazine, phenazopyridine, phenobarbital, phenothiazines, phenytoin, phosphorus, pindolol, piperacillin, piroxicam, plicamycin, polythiazide, procainamide, prochlorperazine, progesterone, promazine, promethazine, propafenone, propoxyphene, propylthiouracil, protriptyline, pyrazinamide, pyrimethamine, quazepam, quinapril, quinethazone, quinidine, ramipril, ranitidine, rifampin, riluzole, risperidone, sargramostim, sildenafil, sibutramine, stanozolol, streptokinase, sulfadiazine, sulfamethoxazole, sulfinamide, sulfoasalazine, sulfisoxazole, sulfonyleureas, sulindac, tacrolimus, terbinafine, tetracycline, thienamidine, thiazides, thiotepa, thiothixene, thiouacil, ticarcillin, ticlopidine, timolol, tocolol, tolazamid, tolazamide, tolbutamide, tolcapone, toltrazurin, toremifene, trilostane, trastuzumab, treintoin, triazolam, trichloromethiazide, trifluoperazine, trimethoprim, trimethoxat, trimipramine, trioxsalen, troglitazone, troenadromycin, trovoxatiazine, ursodiol, vitamin D, warfarin, zalcitabine, zidovudine, zolmitriptan, zolpidem

**Amino acids (total/fractions) (Urine)**

**Increased by:**
- acetaminophen, amikacin, amino-caproic acid, amphoterein, ampicillin, aspirin, phenelzine, subsalicylatic acid, brompheniramine, carbenicillin, cefaclor, cefadroxil, cephalaxin, cephradine, ceftriaxone, colistin, corticotropin, cytarabine, doxorubicin, ephedrine, erythromycin, gentamicin, hydrocortisone, ifosfamide, isosum, insulin, kanamycin, levamisole, levodopa, mafenide, methamphetamine, methyldopa, neomycin, nystatin, penicillamine, paroxtaloid extract, phenobarbitol, phenylephrine, phenylpropanolamion, pramide, ProSobee, pseudoephedrine, streptozocin, tetracycline, trimetazidine, trimethoprim, unidazole, vancomycin, vitamin D

**Decreased by:**
- insulin

**Ammonia (Plasma)**

**Increased by:**
- acetazolamide, asparaginase, chlorothiazide, chlorothalidone, ethacrynic acid, felbamate, furosemide, hydroflumethiazide, ion exchange resins, isoniazid, pegaspargase, promazine, tricyclic, thyroid extract, vitamin D, warfarin, zalcitabine, zidovudine

**Decreased by:**
- cefotaxime, diphenhydramine, lamotrigine, lactobacillus acidophilus, levodopa, MAO inhibitors, neomycin, tetracycline, thrombomycin
Amylase (Serum)  
Increased by:  
- aceterminophen, aminosalicylic acid, amoxapine, amphotericin B, asparaginase, asprin, atovaquone, azathioprine, azithromycin, benzthiazide, betablockers, calcium, calcitriol, cerivastatin, chlorothiazide, chlorpromazine, clonidine, guanfacine, glutethimide, hydrocortisone, indomethacin, prednisone, retinoids, sodium bicarbonate, sodium fluoride, sodium valproate, sulindac, thiamine, ticlopidine, vitamin E, xanthines

Angiotensin Converting Enzyme (Serum)  
Increased by:  
- acebutolol, amitriptyline, anticoagulants, bisoprolol, captopril, chlorpromazine, chlorthalidone, cyclosporin A, diltiazem, furosemide, hydroflumethiazide, ibuprofen, ketoprofen, naproxen, nifedipine, quinapril, ramipril, trandolapril

Antidiuretic Hormone (Plasma)  
Increased by:  
- chlorthalidone, cisplatin, ether, furosemide, hydrochlorothiazide, lithium methyclothiazide, polythiazide

Decreased by:  
- chlorpromazine, clonidine, guanfacine

Antinuclear Antibody (ANA) (Serum)  
Increased by:  
- acebutolol, amitriptyline, anticoagulants, bisoprolol, captopril, chlorpromazine, chlorthalidone, cyclosporin A, diltiazem, furosemide, hydroflumethiazide, ibuprofen, ketoprofen, naproxen, nifedipine, quinapril, ramipril, trandolapril

Aspartate  
Increased by:  
- aceterminophen, aminosalicylic acid, amoxapine, amphotericin B, asparaginase, asprin, atovaquone, azathioprine, azithromycin, benzthiazide, betablockers, calcium, calcitriol, chlorothiazide, chlorpromazine, clonidine, guanfacine, glutethimide, hydrocortisone, indomethacin, prednisone, retinoids, sodium bicarbonate, sodium fluoride, sodium valproate, sulindac, thiamine, ticlopidine, vitamin E, xanthines

Aminothransferase (AST) (Serum)  
Increased by:  
- acebutolol, amitriptyline, anticoagulants, bisoprolol, captopril, chlorpromazine, chlorthalidone, cyclosporin A, diltiazem, furosemide, hydroflumethiazide, ibuprofen, ketoprofen, naproxen, nifedipine, quinapril, ramipril, trandolapril
<table>
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<tr>
<th>Drug Class</th>
<th>Effect</th>
<th>Examples</th>
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<tr>
<td>Apolipoprotein A</td>
<td>Increased by</td>
<td>carbachamazepine, furosemide, gemfibrozil, nisoldipine, oral contraceptives</td>
</tr>
<tr>
<td>(Serum)</td>
<td>Decreased by</td>
<td>lovastatin</td>
</tr>
<tr>
<td>Apolipoprotein B</td>
<td>Increased by</td>
<td>acenocumarol, atenolol, clorhidalione, conjugated estrogens, cyclophosphine,</td>
</tr>
<tr>
<td>(Serum)</td>
<td>Decreased by</td>
<td>estron/progestin therapy, etretinale, furosemide, gemfibrozil, isoretinoin,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>levonorgestrel, methycoid-thiazide, metoporal, oral contraceptives, phenobarbital,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>radioactive iodine, simvastatin, stanozolol</td>
</tr>
<tr>
<td>Atrial Natriuretic</td>
<td>Increased by</td>
<td>atenolol, captopril, carotenol, cyclophosphine, dipyrimal, doxorubicin, morphine,</td>
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<td>Peptide (Plasma)</td>
<td></td>
<td>furosemide, gemfibrozil, inosine, microthelin, prednisolone, procainamide,</td>
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<td>Decreased by</td>
<td>ritonavir, saquinavir, sargramostim, sorbitan, stanozolol, stavudine, streptokinase, streptomycin, sulfadiazine,</td>
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<tr>
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<td>sulfasalazine, sulfisomizide, sufloxylurea, sulindac, sumatriptan, tacrocin,</td>
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<td></td>
<td></td>
<td>tocapleine, tometopine, toremifene, tramadol, tracyflcyromyne, trastu-zumab,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>trazolam, trichloroethazide, trifluoperazine, trimethoprim, trimetrexate, trimipran,</td>
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<td></td>
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<td>trioxsalen, triprolidine, troglitazone, trolessamid, uracil musterd, urapidil,</td>
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<tr>
<td></td>
<td></td>
<td>valsartan, venlafaxine, verapamil, vidarbine, vinorelbine, warfarin, zalcitabine,</td>
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<tr>
<td></td>
<td></td>
<td>zidovudine, zolmitriptan, zolpidem</td>
</tr>
<tr>
<td>Basophils</td>
<td>Increased by</td>
<td>desipramine, paroxetine, tretinoin, triazolam, venlafaxine</td>
</tr>
<tr>
<td></td>
<td>Decreased by</td>
<td>procainamide, thiopeptial</td>
</tr>
<tr>
<td>Bilirubin (Serum)</td>
<td>Increased by</td>
<td>acetybutilol, acetaminophen, ace-azolamide, acetohexamide, acobephazensane,</td>
</tr>
<tr>
<td></td>
<td>Decreased by</td>
<td>atenolol, captopril, chlortalkidone, conjugated estrogens, cyclophosphine,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>estron/progestin therapy, etretinale, furosemide, gemfibrozil, isoretinoin,</td>
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<tr>
<td></td>
<td></td>
<td>levonorgestrel, methycoid-thiazide, metoporal, oral contraceptives, phenobarbital,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>radioactive iodine, simvastatin, stanozolol</td>
</tr>
</tbody>
</table>

**Decreased by:**
- alluporinol, ascorbic acid, clomipramine, cyclosporine, ibuprofen, ketoprofen, metronidazol, naltrexone, penicillin, pindolol, pindosiol, progesterone, rifampin, simvastatin, toremifene, trifluoperazine, urapidil, valsartan, venlafaxine, verapamil, vidarbine, vinorelbine, warfarin, zalcitabine, zidovudine, zolmitriptan, zolpidem
Increased by:

- Phenytoin, phosphp, piper-acillin, piroxicam, polythiazide, prazosin, primaquine, probenecid, procainamide, procarbazine, prochlorperazine, progesterone, promazine, promethazine, propafenone, propoxphene, protyphioauracil, pyrazinamide, quazepam, quinapril, quinethazone, quinidine, quinupristin, radiographic agents, ramipril, ranitidine, rifampin, salicylate, saqui-navir, sargramostim, sorbitan, stanozolol, stavudine, streptomycin, sulfaacetamide, sulfadiazine, sulfadiazine, sulfadoxine, sulfadimethoxine, sulfadiazine, sulfafurazone, sulfamethazine, sulfinpyrazone, sulfisoxazole, sulfonyleurase, sulindac, tacrolimus, tamoxifen, terbinafine, tetracycline, thia bendazole, thiazides, thymiperazine, thioguanine, thioridazine, thiotoxine, thioracil, ticarcillin, ticlidpine, timolol, tobramycin, tocaidine, tolazamide, tolazoline, tolbutamide, tolcapone, toletin, topotecan, toremifene, tramadol, trandolapril, tranlycromine, trastuzumab, treintoxin, triazolam, trimethobenzamide, trimethoprim, trimetrexate, trimipramine, tryptophan, trole-andromycin, uracil mustard, valproic acid, venlafaxine, verapamil, vidarabine, vinorelbine, vitamin K, zalcitabine, zidovudine, zolmitriptan, zolpidem

Decreased by:

- Amikacin, anticonvulsants, aspirin, barbiturates, carbamazepine, cyclosporine, hydroxyurea, isoretinoin, penicilllin, pindolol, prednisone, sulfisoxazole, theophylline, thioridazine, Ursodiol, valproic acid

**Bilirubin (Urine)**

- Increased by: acetohexamide, acetophenazine, aminosalicyclic acid, chlorpromazine, dapsone, etodolac, flufenazine, imipramine, isoniazid, methyldopa, nabumetone, norethandrolone, perphenazine, phenothiazines, tolmetin

Decreased by: ascorbic acid, chlorothalidone

**Bleeding time (Blood)**

- Increased by: acetylsalicyclic acid, aminocaproic acid, ampicillin, aspirin, carbo oil, carbencillin, clodipogrel, dexam, dithran, diflunisal, flutiazam, fluoxetine, gabapentin, hetastarch, ketorolac, moxalactam, nafcillin, naproxen, nifedipine, nonsteroidal anti-inflammatory drugs, oxaprozin, penicillcin, piroxicam, procycline, propranolol, streptokinase, sulindac, ticarcillin, tidione, tolmetin, valproic acid, warfarin

Decreased by: conjugated estrogens, desmopressin, epropitin alfa, erythropoietin

**Blood Cells and Red Casts (Urine)**

- Increased by: abicimab, aldeisleukin, alprostadil, alpetase, ardeparin, aurothiglo, basilikimab, BCG vaccine, bicalutamide, candesartan, carvedilol, cefuroxime, chlorothiazide, cidofovir, clofibrate, clofibrate, clobidogrel, danazol, diclofenac, disopyramide, doxazosin, doxorubicin, doxorubicin, ethosuximide, etretinate, fenoprofen, gabapentin, ganciclovir, gemcitabine, gold, goserelin, ibuprofen, indinavir, indomethacin, isoretinoin, itraconazole, ketoprofen, ketorolac, lansoprazole, leuprolide, lomefloxacin, mephenamic acid, mesalamine, metformin, metoprolol, micardis, minocycline, misoprostol, mycophenolate, naproxen, nifedipine, olsalazine, omeprozole, oxaprozin, oxycodeone, paroxetine, pegasparagase, penicillamine, piroxicam, probenecid, pyrimethamine, rifampin, risperidone, somatotropin, sotalol, sulfinpyrazone, sulfisoxazole, sulindac, ticlidpine, tolcapone, tolmetin, valproic acid, warfarin

Decreased by: ascorbic acid, captoprill, finasteride, lamotrigine

**Blood Urea Nitrogen (BUN)**

- Increased by: ACE inhibitors, acetaminophen, acetylaminol, acyclovir, albendazole, aldeisleukin, alkaline antacids, allopurinol, altretamite, amantadine, amikacin, amiloride, amino acids, aminocaproic acid, amiodarone, amphotericin B, amyI nitrite, anabolic steroids, arsenicals, asparaginase, aspirin, atenolol, azathioprine, azithromycin, bacitracin, benazepril, benzthiazide, betaxolol, bismuth subsalicylate, bisoprolol, busulfan, calcitriol, candesartan, cannabin, capreomycin, captopril, carbamazepine, carvedilol, castor oil, cefaclor, cefamandole, cefazolin, cefalexin, cefonicid, cefoperazone, cefotaxime, cefotetan, cefoxitin, cefoproxime, ceftriaxone, clindamycin, clopidogrel, codeine, colchicine, conjugated estrogens, desmopressin, epoetin alfa, ethacrynic acid, ethambutol, ethosuximide, etilornor, etretinate, fenoprofen, gabapentin, ganciclovir, gemcitabine, gold, goserelin, ibuprofen, indinavir, indomethacin, isoretinoin, itraconazole, ketoprofen, ketorolac, lansoprazole, leuprolide, lomefloxacin, mephenamic acid, mesalamine, metformin, metoprolol, micardis, minocycline, misoprostol, mycophenolate, naproxen, nifedipine, olsalazine, omeprozole, oxaprozin, oxycodeone, paroxetine, pegasparagase, penicillamine, piroxicam, probenecid, pyrimethamine, rifampin, risperidone, somatotropin, sotalol, sulfinpyrazone, sulfisoxazole, sulindac, ticlidpine, tolcapone, tolmetin, valproic acid, warfarin

Decreased by: ascorbic acid, captoprill, finasteride, lamotrigine

**C3 Complement (Serum)**

- Increased by: cimetidine, cyclophosphamide, oral contraceptives

**C4 Complement (Serum)**

- Decreased by: danazol, hydralazine, methyldopa, phencytoin

- Increased by: cyclophosphamide, danazol, oral contraceptives

- Decreased by: dextran, methyldopa, penicillamine
**C-Peptide (Plasma)**

*Increased by:* betamethasone, deferoxamine, glimepiride, glyburide, indapamide, isoproterenol, oral contraceptives, prednisone, terbutaline

*Decreased by:* atenolol, calcitonin, miglitol

**Calcitonin (Plasma)**

*Increased by:* estrogen/progesterin therapy, pentagastrin

*Decreased by:* estrogens, octreotide, phenytoin

**Calcium (Serum)**

*Increased by:* aldesleukin, alkaline antacids, aluminum, anabolic steroids, antacids, basiliximab, calcitriol, calcium gluconate, captopril, cefotaxime, chlorothiazide, chloropropamide, chlorothalidone, dienestrol, diethylstilbestrol, dihydrotestosterone, doxorubicin, estramustine, estropipate, etretinate, fluoxymesterone, hydralazine, hydrochlorothiazide, iron, leuprolide, lithium, levodopa, megestrol, methandienone, methandrostenolone, methadone, mifepristone, nandrolone, norethisterone, oral contraceptives, oxymetholone, parathyroid hormone, paroxetine, pentostatin, phenobarbital, polystyrene sulfonate, polynucleotides, propranolol, rifampicin, sirolimus, spironolactone, tamoxifen, theophylline, thiazides, toremifene, trastuzumab, tretinoin, trichlormethazine, vitamin D, zalcitabine

*Decreased by:* acetazolamide, aldesleukin, alendronate, amifostine, amlodipine, amphotericin B, anticonvulsants, arsenic trioxide, asparaginase, aspirin, basiliximab, bisphosphonates, calcitonin, carprofen, chloroquine, chlorothiazide, cidofovir, cisplatin, corticosteroids, corthione, diuretics, doxorubicin, erythropoietin, estrogen/progesterin therapy, estropipate, etidronate, etretinate, felbamate, fosarnet, furosemide, gallium, gentamicin, glucocorticoids, hydrochlorothiazide, insulin, interferon, iron dextran, isoniazid, ketoconazole, laxatives, magnesium salts, methicillin, mycophenolate, oral contraceptives, pamidronate, paracetamol, pentamidine, phenobarbital, phenytoin, plicamycin, polystyrene sulfonate, prednisone, riluzole, sirolimus, spironolactone, tamoxifen, theophylline, thiazides, toremifene, trastuzumab, tretinoin, trichlormethazine, vitamin D, zalcitabine

**Calcium (Urine)**

*Increased by:* acetazolamide, aldesleukin, alendronate, amifostine, amlodipine, amphotericin B, anticonvulsants, arsenic trioxide, asparaginase, aspirin, basiliximab, bisphosphonates, calcitonin, carprofen, chloroquine, chlorothiazide, cidofovir, cisplatin, corticosteroids, corthione, diuretics, doxorubicin, erythropoietin, estrogen/progesterin therapy, estropipate, etidronate, etretinate, felbamate, fosarnet, furosemide, gallium, gentamicin, glucocorticoids, hydrochlorothiazide, insulin, interferon, iron dextran, isoniazid, ketoconazole, laxatives, magnesium salts, methicillin, mycophenolate, oral contraceptives, pamidronate, paracetamol, pentamidine, phenobarbital, phenytoin, plicamycin, polystyrene sulfonate, prednisone, riluzole, sirolimus, spironolactone, tamoxifen, theophylline, thiazides, toremifene, trastuzumab, tretinoin, trichlormethazine, vitamin D, zalcitabine

*Decreased by:* alendronate, bicarbonate, bisphosphonates, calcitonin, chloroquine, chlorothalidone, diuretics, etidronate, furosemide, hydrochlorothiazide, insulin, interferon, iron, lapatinib, metformin, metoclopramide, minoxidil, nandrolone, norethisterone, oral contraceptives, pamidronate, parathyroid extract, phenytoin, polystyrene sulfonate, prednisone, raloxifene, sirolimus, spironolactone, tamoxifen, theophylline, thiazides, trichlormethazine, vitamin D, zalcitabine

**Chloride (Serum)**

*Increased by:* acetazolamide, acetylcysteine, ammonium chloride, aspirin, cannabis, carbamazepine, carprofen, cefotaxime, chloride salts, chlorothiazide, cholestyramine, corticosteroids, diazoxide, etretinate, guanethidine, hydrochlorothiazide, hydrocortisone, ibuprofen, insulin, lactulose, lithium, methicillin, methyltestosterone, metoprolol, metoclopramide, minoxidil, nandrolone, n-butylscopolamine, nitrates, oral contraceptives, plicamycin, polystyrene sulfonate, prednisone, raloxifene, sirolimus, spironolactone, tamoxifen, theophylline, thiazides, trichloroethylene, vitamin D, vitamin K

*Decreased by:* allopurinol, amiloride, aspirin, bicarbonate, bumetanide, cefotaxime, cholestyramine, chlorpropamide, chlorothalidone, corticosteroids, corticotropin, cortisone, diuretics, etretinate, furosemide, hydrochlorothiazide, hydrocortisone, hydrofluoromethane, hydroxyurea, insulin, ketoprofen, lithium, magnesium, methicillin, metoclopramide, minoxidil, nandrolone, norethisterone, oral contraceptives, pamidronate, parathyroid extract, phenytoin, polystyrene sulfonate, prednisone, raloxifene, sirolimus, spironolactone, tamoxifen, theophylline, thiazides, trichlormethazine, vitamin D, vitamin K

**Cholesterol (Serum)**

*Increased by:* acetohexamide, acetophenazine, aminoglutethimide, amiodarone, amphotericin B, anabolic steroids, ampicillin, aspirin, basiliximab, bisphosphonates, calcitonin, carprofen, chloroquine, chlorothiazide, ciprofloxacin, clonidine, clofibrate, clonidine, clomipramine, conjugated estrogens, corticosteroids, cortisone, cyclophosphamide, cyclosporine, danazol, dantrolene, dapsone, diclofenac, disulfiram, etanercept, etretinate, fluoxetine, furosemide, hydrochlorothiazide, hydrocortisone, hydrofluoromethane, hydroxyurea, insulin, ketoprofen, lithium, metformin, minoxidil, nandrolone, norethisterone, oral contraceptives, oxymetholone, paroxetine, plicamycin, polystyrene sulfonate, prednisone, raloxifene, sirolimus, spironolactone, tamoxifen, theophylline, thiaoacetylene, thiazides, thioracil, ticlopidine, tolcapon, tretinoin, troglitazone, venlafaxine, vitamin D, zolpidem
Decreased by: acarbose, acetobutol, albuterol, aldesleukin, allopurinol, aluminum hydroxide, amikacin, amiloride, aminosalicylic acid, amidodarone, amiodipine, ampicillin, ascorbic acid, asparaginase, aspirin, atenolol, atorvastatin, azathioprine, bisoprolol, captopril, carvedilol, chlorambucil, chlorofom, chloropropamide, chlordihalodone, chlorymestane, clazaprin, clofibrate, clomiphene, clonidone, coenzyme Q10, colchicine, colistimethate, conjugated estrogens, diottazine, doxorubicin, enalapril, erthyromycin, es supplied estrogens, estrogen therapy, fenofibrate, fluoroquinolone, fluvastatin, fosinopril, glyburide, glycyrrhizinic factor, guanabenz, haloperidol, HMG CoA-rectductase inhibitors, idomethacin, insulin, isoniazid, isornisid, ispriflokrin, ketocanzone, lansoprazole, levonorgestrel, levothyrroxine, lincomycin, lisinopril, LMW heparin, losartan, MAO inhibitors, medroxyprogesterone, metformin, methyldopa, metoprolol, metronidazole, nandrolone, neomycin, niacin, nicotinic acid, nifedipine, Norplant, oral contraceptives, oxandrolone, oxymetholone, perindopril, phenytoin, pindolol, pravastatin, prazosin, prednisolone, probucol, progestosterone, pyllium, raloxifene, ramipril, simvastatin, spironolactone, statins, streptokinase, tacrolimus, tamoxifen, terazosin, tetracycline, thiazides, thyroid, tolbutamide, trozadone, ursodiol, valproic acid, verapamil

**Creatinine (Serum)**

**Increased by:** anticoagulants, carbicillin, dicumarol, pegpyraparagase, plicamycin, tetracycline, ticarcillin

**Decreased by:** amphenyline, oral contraceptives

**Positive result by:** aminosalicylic acid, aztreonam, cefadroxil, cefazidime, cefcloxime, cefoxime, chloromazine, chloropropamide, ethosuximide, hexadralazine, ibuprofen, imipenem cilastatin, isoniazid, levodopa, mefenamic acid, melphalan, methyldopa, moxalactam, penicillin, phenytoin, procainamide, quinidine, quinine, streptomycin, sulfonyleurases, tetracycline

**Cortisol (Plasma)**

**Increased by:** anticonvulsants, aspirin, atropine, benzodiazipines, citalopram, clomipramine, corticotropin, corticocortin-releasing hormone, cortisone, dcoxidide, diclofenac, estrogens, ether, ethyl, estradiol, fenoprofen, furosemide, gemfibrozil, glyburide, hydrocortisone, insulin, interferon, interleukin, lithium, methadone, methoclopramide, naloxone, octreotide, opiates, oral contraceptives, prednisolone, prednison, ranitidine, spironolactone, tumor necrosis factor, vasopressin

**Decreased by:** aminoglutethimide, barbiturates, beclomethasone, budesonide, clonidone, corticosteroids, danazol, dexamethasone, dextroamphetamine, diazoxide, epinphen, etomidate, fluocinolone, indomethacin, ketoconazole, labelotol, levodopa, lithium, magnesium sulfate, medroxyprogesterone, megestrol, mesalamine, midazolam, morphine, nifedipine, nitrous oxide, norethindrone, oxazepam, phenobarbital, phenytoin, pravastatin, prednisolone, ranitidine, rifampin, sumatRIPTAN, trimcinolone, trimipramine

**Creatinine (Serum)**

**Increased by:** ACE inhibitors, acetobutol, acetyhemide, acemetaphen, acetyclovir, albendazole, aldesleukin, ankle antacids, alprazolam, alprostadil, altretamine, amikacin, amiloride, amidodarone, amphotericin B, ascorbic acid, asparaginase, aspirin, azathioprine, azithromycin, aztreonam, barbiturates, benzazepin, betazolol, bicalutamide, bisoprolol, danesartan, capreomycyn, captropil, carbamazepine, carvedilol, cefamandole, cefazolin, cefixime, cefoperazone, cefotaxime, cefotolin, cefpodoxime, cefuroxime, ceftriazone, ceftriaxone, cephapilin, cephradine, cetrizine, chlorothiazide, clorpropamide, chlordihalodone, cidofovir, cimetidine, cinacixin, ciprofloxacin, cispalatin, clarithromycin, clofibrate, clonidine, clorazepate, co-trimoxazole, colistimethate, colistin, cyclosporine, danazol, demecycloline, dextran, diclofenac, disopyramide, doxycycline, enalapril, enflurane, epoetin alfa, esprotan, ethambutol, etidronate, etretinate, fenoprofen, furocystone, fludarabine, foscarin, furosemide, gabapentin, gemcitabine, gemfibrozil, gentamicin, griseofulvin, imipenem, indomethacin, interleukin alfa-2, irbesartan, isomethacin, kanamycin, ketoprofen, ketorolac, labelotol, lamotrigine, lansoprazole, leuprolide, levodopa, lidocaine, lisoprinol, lithium, loracarbef, losartan, lovastatin, manitol, meclofenamate, mephenemine, meropenem, mesalamine, methicillin, methotrexate, methylhydroxy, methylprednisolone, metoprolol, micardis, mitomycin, mitoxantrone, moexipril, moxalacamt, mycophenolate, niadilaxid acid, naproxen, neomycin, nelamicin, nifedipine, niletuline, nisoldpine, nonsteroidal anti-inflammatory drugs, norflaxacin, ofloxacacin, olsalazine, oxacillin, oxaprozin, paminodurate, paromycin, pegasparagase, penicillamine, pencillin, pentamide, pentostatin, phenazopyridine, phosphorus, piperacillin, piroxicam, plicamycin, prodnisone, propafenone, quazepam, quinapril, radioactive agents, ramipril, ranitidine, risperidone, salisalate, sargramostim, sevoflurane, streptokinase, streptomycin, stropozocin, sulfamethoxazole, sulfasalazine, sulfisoxozone, sulindac, suprofen, tacrolimus, tetracycline, thiazides, ticarcillin, ticlopidine, timolol, tobramycin, tramadol, trandolapril, trinol, trimaterene, trimazolam, trimethoprime, trimetrexate, trovafloxacin, ursodiol, valsartan, vancomycin, vasopresin, venlafaxine, vitamin D

**Decreased by:** amikacin, ascorbic acid, cannabis, captopril, dobutamine, ibuprofen, interferon alfa-2a, lisinopril, methylhydroxy, nicardipine, prodnisone, terazosin, triazolam, tromethamine, valproic acid, zidovudine
Creatinine Kinase (CPK) (Serum) Increased by: acetylsalicylic acid, amikacin, ascorbic acid, dantrolene, phenothiazines, pindolol, prednisone, sulfamethoxazole

Creatinine (Urine) Increased by: acetylsalicylic acid, amikacin, ascorbic acid, dantrolene, phenothiazines, pindolol, prednisone, sulfamethoxazole

Eosinophils (Blood) Decreased by: al-glucerase, antithyroid therapy, basiliximab, corticotropin, danazol, dexfenfluramine, erythropoietin, etretinate, glucocorticoids, hydrochlorothiazide, mycophenolate, pilocarpine

Erythrocytes (Blood) Increased by: al-glucerase, antithyroid therapy, basiliximab, corticotropin, danazol, dexfenfluramine, erythropoietin, etretinate, glucocorticoids, hydrochlorothiazide, mycophenolate, pilocarpine

Erythrocyte Sedimentation Rate (Blood) Increased by: anticonvulsants, aspirin, carbamazepine, cephalothin, cephradin, clozapine, cyclosporine A, dexamethasone, etretinate, fluvasatin, hydralazine, indomethacin, isoretinoin, lomefloxacin, methotrexate, misoprostol, oxazepam, oral contraceptives, prednisone, piroxicam, quinapril, sulfamethoxazole, zolpidem
Erythropoietin (Serum)
Increased by: anabolic steroids, daunorubicin, erythropoietin, fluoroxymesterone, hydroxyurea, theophylline, zidovudine
Decreased by: acetazolamide, amphotericin B, cisplatin, enalapril, furosemide, theophylline

Ethanol (Serum)
Increased by: ascorbic acid, atropine, phenobarbital, propranolol
Decreased by: acobarbital, acebutolol, amino acids, asparaginase, aspirin, atenolol, clofibrate, glyburide, insulin, isoproterenol, levotyroxine, metformin, metoprolol, neomycin, niacin, nicotinic acid, nifedipine, prazosin, propranolol, propylthiouracil, simvastatin, sotalol, streptozocin

Euglobulin Clot Lysis time (Blood)
Increased by: cyclosporine A
Decreased by: asparaginase, clofibrate, dextran, gemfibrozil, streptokinase

Fatty Acids (Free) (Plasma)
Increased by: amiodarone, ampicillin, aspirin, antacids, anticonvulsants, aspirin, barbiturates, chloramphenicol, cholestyramine, cycloserine, diethylstilbestrol, erythromycin, espnadin, insulin, ivermectin, metformin, methotrexate, nitrofurantoin, oral contraceptives, penicillin, pentamidine, phenobarbital, phenylethanolamine, rifampin, salsalate, sulfasalazine, sulfisoxazole, tetracycline, triamterene, trimethoprim
Decreased by: acarbuzolamide, aminosalicylic acid, ampicillin, ascorbic acid, aspirin, atenolol, clonidine, dexamethasone, cholestyramine, cycloserine, diethylstilbestrol, erythromycin, espnadin, insulin, ivermectin, metformin, methotrexate, nitrofurantoin, oral contraceptives, penicillin, pentamidine, phenobarbital, phenylethanolamine, rifampin, salsalate, sulfasalazine, sulfisoxazole, tetracycline, triamterene, trimethoprim

Fibrinogen (Plasma)
Increased by: ascorbic acid, atropine, phenobarbital, propranolol
Decreased by: acobarbital, acebutolol, amino acids, asparaginase, aspirin, atenolol, clofibrate, glyburide, insulin, isoproterenol, levotyroxine, metformin, metoprolol, neomycin, niacin, nicotinic acid, nifedipine, prazosin, propranolol, propylthiouracil, simvastatin, sotalol, streptozocin

Ferritin (Blood)
Increased by: ferrous sulfate, iron preps, oral contraceptives, theophylline
Decreased by: antithyroid therapy, ascorbic acid, deferoxamine, methimazole

Ethanol (Serum)
Increased by: anabolic steroids, daunorubicin, erythropoietin, fluoroxymesterone, hydroxyurea, theophylline, zidovudine
Decreased by: acetazolamide, amphotericin B, cisplatin, enalapril, furosemide, theophylline

Euglobulin Clot Lysis time (Blood)
Increased by: cyclosporine A
Decreased by: asparaginase, clofibrate, dextran, gemfibrozil, streptokinase

Fatty Acids (Free) (Plasma)
Increased by: amiodarone, ampicillin, aspirin, antacids, anticonvulsants, aspirin, barbiturates, chloramphenicol, cholestyramine, cycloserine, diethylstilbestrol, erythromycin, espnadin, insulin, ivermectin, metformin, methotrexate, nitrofurantoin, oral contraceptives, penicillin, pentamidine, phenobarbital, phenylethanolamine, rifampin, salsalate, sulfasalazine, sulfisoxazole, tetracycline, triamterene, trimethoprim
Decreased by: acarbuzolamide, aminosalicylic acid, ampicillin, ascorbic acid, aspirin, atenolol, clonidine, dexamethasone, cholestyramine, cycloserine, diethylstilbestrol, erythromycin, espnadin, insulin, ivermectin, metformin, methotrexate, nitrofurantoin, oral contraceptives, penicillin, pentamidine, phenobarbital, phenylethanolamine, rifampin, salsalate, sulfasalazine, sulfisoxazole, tetracycline, triamterene, trimethoprim

Folic acid (Serum)
Increased by: asparaginase, clofibrate, dextran, gemfibrozil, streptokinase
Decreased by: asparaginase, clofibrate, dextran, gemfibrozil, streptokinase

Folic acid Stimulating Hormone (FSH) (Plasma)
Increased by: bicalutamide, bromocriptine, cimetidine, clofungine, danazol, erythropoietin, finasteride, follicle-stimulating hormone, gonadotropin-releasing hormone, growth hormone-releasing hormone, hydrocortisone, ketoconazole, leuprolide, levodopa, metformin, methotrexate, nitrofurantoin, oral contraceptives, penicillin, pentamidine, phenobarbital, phenyltoin, primidone, pyrimethamine, rifampin, salsalate, sulfasalazine, sulfisoxazole, tetracycline, triamterene, trimethoprim
Decreased by: acarbose, acebutolol, amino acids, asparaginase, aspirin, atenolol, clofibrate, glyburide, insulin, isoproterenol, levotyroxine, metformin, metoprolol, neomycin, niacin, nicotinic acid, nifedipine, prazosin, propranolol, propylthiouracil, simvastatin, sotalol, streptozocin

Ferritin (Blood)
Increased by: ferrous sulfate, iron preps, oral contraceptives, theophylline
Decreased by: antithyroid therapy, ascorbic acid, deferoxamine, methimazole

Fibrinogen (Plasma)
Increased by: ascorbic acid, atropine, phenobarbital, propranolol
Decreased by: acobarbital, acebutolol, amino acids, asparaginase, aspirin, atenolol, clofibrate, glyburide, insulin, isoproterenol, levotyroxine, metformin, metoprolol, neomycin, niacin, nicotinic acid, nifedipine, prazosin, propranolol, propylthiouracil, simvastatin, sotalol, streptozocin

Folic acid (Serum)
Increased by: niacin, nitrofurantoin, oral contraceptives, theophylline, zidovudine
Decreased by: theophylline, zidovudine

Free Thyroxine Index (Serum)
Increased by: amiodarone, carbamazepine, furosemide, levothyroxine, oral contraceptives, orphenadrine, phenobarbital, propranolol
Decreased by: amiodarone, aspirin, clofungine, co-trimoxazole, corticosteroids, ferrous sulfate, iodide, isoretinoin, lovastatin, methimazole, phenobarbital, phenyltoin, primidone, radioactive iodine, salsalate

Glucagon (Plasma)
Increased by: asparaginase, danazol, glucocorticoids, guanabenz, hydrochloorthizide, insulin, interferon alpha-2a, nifedipine, prednisolone, propranolol
Decreased by: atenolol, insulin, metoprolol, octreotide, pindolol, propranolol, troglitazone, verapamil

Glucose (Cerebrospinal Fluid)
Increased by: cefotaxime, dexamethasone
Decreased by: cefotaxime

Glucose (Urine)
Increased by: acarbose, ampicillin, ascorbic acid aspirin, bisacodyl, chloral hydrate, cholestyramine, diazepam, digoxin, ferrous sulfate, flurazepam, furosemide, hydroquinone, insulin, levodopa, oxytetacycline, phenazopyridine, phenobarbital, radiographic agents, secoaromatic, tetracycline, vitamin preparations
Decreased by: acarbose, ampicillin, ascorbic acid aspirin, bisacodyl, chloral hydrate, cholestyramine, diazepam, digoxin, ferrous sulfate, flurazepam, furosemide, hydroquinone, insulin, levodopa, oxytetacycline, phenazopyridine, phenobarbital, radiographic agents, secoaromatic, tetracycline, vitamin preparations
Increased by:
- acarbose, atenolol, clofibrate, fluoroxyemterone, glyburide, guanethidine, lisinopril, MAO inhibitors, metformin, metoprolol, nandrolone, norethindrone, octreotide, pargylne, phentoin, prazosin, terazosin, troglitazone

Decreased by:
- aceturbutol, atenolol, beclomethasone, betamethasone, calcitonin, cannabis, chlorothiazide, chloropromazine, clorhohalidone, clofibrate, conjugated estrogens, cortisone, danozol, dexamethasone, diazoxide, diethylstilbostrol, estropipate, ethacrynic acid, helodipine, fluoroxyemterone, foscarin, furosemide, hormone high-density, hydrochlorothiazide, hydroxychloroquine, imipramine, interferon alfa-2a, iron, levonorgestrel, lithium, medroxypogesterone, mefenamic acid, mestranol, methadrenostenolone, methylprednisolone, metoprolol, naproxen, niacin, niacinamide, nicotinic acid, nifedipine, nitrofurantoin, norethindrone, octreotide, oral contraceptives, penphenezone, phentoin, pindolol, polythiazide, prednisolone, prednisone, promazine, quinethazone, spironolactone, streptozocin, thiazides, trimcinolone, triamterene, verapamil

Glycosylate Hemoglobin (Hb A1c) (Blood)
Increased by:
- aspirin, atenolol, beta-blockers, gemfribozol, gleipiride, hydrochlorothiazide, indapamide, lovastatin, niacin, nicardipine, nicotinic acid, proranolol

Decreased by:
- acarbace, deferoxamine, diltiazem, enalapril, gilpizide, glyburide, insulin, lisinopril metformin, nissoline, pravastatin, ramipril, terazosin, verapamil

?-Glutamyltransferase (GTT) (Serum)
Increased by:
- acetaminophen, allopurinol, alprazolam, amiodarone, aphotonerin B, anastrozole, anticonvulsants, aurothiogluocese, azithromycin, barbiturates, captorpl, carbamazepine, cefdinir, cefclidnic, cefpodoxime, ceftazidime, ceftrizine, chloramphenicol, chlorpromazine, chloropromamide, cimetidine, cisplatin, clomipramine, clozapine, cyclopersin A, dactinomycin, dantrole, diciofenac, disopryamide, disulfiram, doxorubucin, doxycycline, enalapril, esterified estrogens, etopropate, etoposide, etretinat, famoldine, fluconazole, flucytosine, fluoroxyemterone, flutamide, flavutan, fosphtomycin, gold, griseofulvin, haloperidol, hepatitis A vaccine, hydrochlorothiazide, ibuprofen, interferon alfa-3, isoniazid, isoretinoin, israpi-dine, ketamine, ketoprofen, labetalol, lansoprazole, levthoxyzone, linsitocin, lomefloxacin, loratadine, lovastatin, medroxypogesterone, meropenem, melsalmine, metothrasete, methylodpa, metoprolol, moexipril, mycoppenolate, nabumetone, naproxen, nelfinavir, nevirinp, niacin, nicardipine, nisoldipine, nitrofurantoin, nortryptiline, octreotide, ofloxacin, olsalazina, omeprazole, oral contraceptives, papaverine, pegaspargase, phenazyoridinede, phenobarbital, phenothiazines, phenytoin, piroxicam, prazosin, probenecid, prochlorperazine, propafenone, propoxyphene, quinapril, quindine, rifampin, riluzole, ritonavir, silbutramine, stanzolol, streptokinase, sulfamethoxazole, sulfasalazine, sulfisoxazol, sulindac, terbinafine, thiaendazole, thiethyl-perazine, thiopental, thioridazine, tocainide, tolmetin, trifluoperazine, triolitazone, trole-andomycin, valproic acid, warfarin, zalcitabile, zidovudine

Decreased by:
- azahliphore, clofibrate, conjugated estrogens, metothrasete, ursodiol

Growth Hormone (Plasma)
Increased by:
- amino acids, anabolic steroids, citalopram, clomipramine, clonidine, desipramine, dexamethasone, diazepam, ethyl estradiol, growth hormone-releasing hormone, indomethacin, insulin, interferon, interleukin, methambemthamine, metoclopramide, midazolam, niacin, oral contraceptives, phenytin, proranolol, pyridostigmine, tumor necrosis factor, varospesrin

Decreased by:
- bromocriptine, chlorpromazine, corticosteroids, hydrocortisone, medroxypogesterone, methylodpa, octreotide, prednisone, propantheline, valproic acid

5-Hydroxyindoleacetic Acid (Urine)
Increased by:
- acetaminophen, chlordiazepoxide, cisplatin, diazepam, ephedrine, flurazepam, fluorouracil, guaufenesin, melphalan, methocarbolip, naproxen, phenobarbital, phenoltamine, raufwofla, sulfasalazine, resperine

Decreased by:
- asphin, chlorpromazine, corticoprin, imipramine, interferon alfa-2a, isoniazid, levodopa, MAO inhibitors, methylodpa, octreotide, phenothiazines, promazine, streptozocin

Complement CH 50 (Serum)
Increased by:
- chlorpromazine, cyclophosphamide

Decreased by:
- hydralazine

High-Density Liprotein (HDL) Cholesterol (Serum)
Increased by:
- acarbose, ACE inhibitors, albuterol, atorvastatin, beclomethasone, captopripl, carbonazepine, carvedilol, cerivastatin, chenodiol, cholestyramine, cimetidine, clofibrate, coenzyme Q10, colestipol, corn oil, diltiazem, doxazocin, estrogen therapy, estrogen/ progesterin therapy, faveinrez, fenofibrate, fluvastatin, furosemide, gemfribozol, glyburide, goserellin, hydroxychloroquine, indapamide, insulin, isradipine, ketcnonazole, lovastatin, medroxypogesterone, meformin, methimazole, minoxidil, nafarelin, niacin, nicardipine, nicotinic acid, nifedipine, nissoline, norplant, oral contraceptives, perindopril, phenobarbital, phenytin, pindolol, pravastatin, prazosin, prednisone, ramipril, rosiglitazone, simvastatin, tiaprofen, trifluoperazine, troglitazone, trole-andomycin, valproic acid, warfarin, zidovudine

Decreased by:
- acebutolol, ascobic acid, atenolol, azathioprine, bisoprolol, beta-blockers, carvedilol, chenodiol, chlorpromamide, clofibrate, cyclopersin A, danazol, estrogen/progesterin therapy, etretinate, gemfribozol, hydrochlorothiazide, indapamide, indomethacin, interferon alfa-2a, isoretinoin, levthoxyzone, linseed oil, lisinopril, medroxypogesterone, methimazole, methylodpa, metoprolol, nadolol, nandrolone, neomycin, norplant, oral contraceptives, prednisolone, probucol, propranolol, psyllium, rallocenaza, sotalol, spironolactone, stanzolol, tamoxifen, thiazides, timolol, trichloroethazide, ursodiol, verapamil

Hydroxyproline (Urine)
Increased by:
- anticonvulsants, corticosteroids, cyclosporine, danazol, hormone replacement, interferon, levthoxyzone, nafarelin, parathyroid hormone, phenobarbital, phenytin, somatotropin, thyroid, tobutamide, vitamin D
**Increased by:**
- antineoplastic agents, ascorbic acid, aspirin, bisphosphonates, budesonide, calcitonin, conjugated estrogens, corticosteroids, estrogen/progesterin therapy, etidronate, gallium nitrate, glucocorticoids, medroxyprogesterone, pamidronate, plicamycin, prednisolone, prednisone, pranoproanol

**Immuno globulin E Antibody (Serum)**

**Increased by:**
- aztreonam, penicillin G

**Decreased by:**
- phenytoin

**Insulin (Plasma)**

**Increased by:**
- acetoheptadine, adenosine, albuterol, amino acids, aspirin, beclomethasone, calcium gluconate, cannabis, captopril, chlorpropamide, chlordihalidone, deferroxamine, glipizide, glyburide, insulin, alfa-2a, alfa-2b, alproserenol, levodopa, lisinopril, medroxyprogesterone, megestrol, metyldrenisone, metoprolol, niacin, nicotinic acid, norenthalone, oral contraceptives, perindopril, prazosin, prednisolone, prednisone, quinidine, rifampin, ritodrine, secretin, spironolactone, streptozocin, terbutaline, tolbutamide, tramadol, triiodothyronine, verapamil

**Decreased by:**
- acarbose, acetoheptadine, calcitonin, chlorpropamide, clofibrate, conjugated estrogen, diazoxide, diltiazem, doxazosin, enalapril, enprostil, ethacrynic acid, either, furosemide, hydrochlorothiazide, metformin, midazolam, morphine, nafacid, nefilipide, octreotide, phenytoin, prazosin, pranoproanol, pyrilium, tolazamide, toltudam, troglitazone

**Iron (Blood)**

**Increased by:**
- acetylsalicylic acid, cefotaxime, chemotherapeutic agents, chloramphenicol, cisplatin, ferrous sulfate, iron, iron dextran, methicillin, methazol, methotrexate, multivitamin, oral contraceptives, pyrazinamide

**Decreased by:**
- allopurinol, aspirin, cholestyramine, corticoterop, cortisone, deferoxamine, metformin, oxymetholone, pergolide, pyrazinamide, risperidone

**Ketones (Urine)**

**Increased by:**
- acetylcoenzyme, aminosalicylic acid, aspirin, captopril, cefdinir, cefixime, dimeracaprol, ether, etodolac, ifosfamide, isoniazid, levodopa, mesna, metformin, methyldopa, nacin, mibitahide, penicillamine, phenazopyridine, phenolpehtaline, phenothiazines, streptozocin, valproic acid

**Decreased by:**
- aspirin, phenytoin pyridinylacide

**Lactate Dehydrogenase (LD) (LDH) (Serum)**

**Increased by:**
- acetobutol, almidarone, amphetomin B, anabolic steroids, aspirin, auranofin, azithromycin, betaxolol, captopril, carbenicillin, cefdinir, cefonicid, cefotaxime, cefetan, cefoxitin, cefpodoxime, ceftazidime, cefuroxime, chloramphenicol, chloropamide, chlorothaldehyde, cimetidine, ciprofloxacin, clindamycin, clofibrate, codein, capsope, diclofenac, diltiazem, donepezil, doxorubicin, estramustine, etretinate, fenoprofen, fleroxidine, fluphenazine, fluvoxime, foscarcin, ganciclovir, gentamicin, gold, granulocyte colony-stimulating factor, hydralazine, ibuprofen, imipramine, interferon alpha-2a, interferon-alpha-2b, isotretinoin, irarecan, levozocin, levodopa, meperidine, mesalamine, methylprednisolone, methotrexate, metyldopam, metoprolol, morphine, pyrophosphine, nefazodone, nelfinavir, nifedipine, nitrofurantoin, norflaxacin, oxacillin, oxaprin, paroxetine, pegfilgrastim, pemoline, penicillamine, pipidol, Piperacill, plicamycin, prophophyrex, pranoprololin, propylthiouracil, quinidine, quinupristin/dalfopristin, rifuzole, sibutramine, simvastatin, streptokinase, streptozocin, sulfamethoxazole, sulfasalazine, sulfosoxazole, sulindac, tacrolimus, tetracycline, thiopental, ticarcillin, tobramycin, tolmetin, valproic acid, vasopressin, verapamil

**Decreased by:**
- amnikcin, anticonvulsants, ascorbic acid, cefotaxime, cefobase, enalapril, hydroxurea, metronidazole, naltrexone

**Luteinizing Hormone (LH) (Plasma)**

**Increased by:**
- bicalutamide, bromocriptin, clophimine, finasteride, gonadotropin-releasing hormone, gosereolin, growth-releasing hormone, hydrocortisone, ketoconazole, leuroplide, nalozone, nilutamide, spironolactone, tamoxifen, valproic acid

**Decreased by:**
- acetabol, amidori, amphetomin B, anabolic steroids, aspirin, auranofin, azithromycin, betaxolol, captopril, carbenicillin, cefdinir, cefonicid, cefotaxime, cefetan, cefoxitin, cefpodoxime, ceftazidime, cefuroxime, chloramphenicol, chloropamide, chlorothaldehyde, cimetidine, ciprofloxacin, clindamycin, clofibrate, codein, capsope, diclofenac, diltiazem, donepezil, doxorubicin, estramustine, etretinate, fenoprofen, fleroxidine, fluphenazine, fluvoxime, foscarcin, ganciclovir, gentamicin, gold, granulocyte colony-stimulating factor, hydralazine, ibuprofen, imipramine, interferon alpha-2a, interferon-alpha-2b, isotretinoin, irarecan, levozocin, levodopa, meperidine, mesalamine, methylprednisolone, methotrexate, metyldopam, metoprolol, morphine, pyrophosphine, nefazodone, nelfinavir, nifedipine, nitrofurantoin, norflaxacin, oxacillin, oxaprin, paroxetine, pegfilgrastim, pemoline, penicillamine, pipidol, Piperacill, plicamycin, prophophyrex, pranoprololin, propylthiouracil, quinidine, quinupristin/dalfopristin, rifuzole, sibutramine, simvastatin, streptokinase, streptozocin, sulfamethoxazole, sulfasalazine, sulfosoxazole, sulindac, tacrolimus, tetracycline, thiopental, ticarcillin, tobramycin, tolmetin, valproic acid, vasopressin, verapamil

**Decreased by:**
- amnikcin, anticonvulsants, ascorbic acid, cefotaxime, cefobase, enalapril, hydroxurea, metronidazole, naltrexone

**Lipase (Serum)**

**Increased by:**
- acetaminophen, asparaginase, azathioprine, bethanechol, calcitro, cerivastatin, chlorothiaizide, cholinergic agents, cidofovir, cimetidine, cloaprine, codeine, conjugated estrogens, cyclosporine, demeclocycline, desipramine, diazoxide, didanosine, donepezil, doxurubicin, enalapril, estradiol, estrogens, estradiol, ethinyl estradiol, finasteride, goserelin, ketoterepine, ketoconazole, leuroplide, medroxyprogesterone, megestrol, metformin, octreotide, oral contraceptives, phenothiazines, phenoytoin, pimozide, pravastatin, prednisone, progesterone, stanozolol, tamoxifen, thioridazine, toremifen, valproic acid

**Decreased by:**
- calcium, hydroxyurea, mesalamine, protamine, simvastatin, tamoxifen, toremifene, valproic acid, venlafaxine

**Lymphocytes (Blood)**

**Increased by:**
- acetylsalicylic acid, cefaclor, cefdinir, cefazidime, chlorambucil, chlorpropamide, dexamethasone, gabapentin, granulocyte colony-stimulating factor, glycoflavin, haloperidol, levodopa, levodopa, narconics, niacinamide, oxazolac, paroxetine, pergolide, propylthiouracil, quazepam, spironolactone, triazolam, valproic acid, venlafaxine

**Decreased by:**
- aplazolam, asparaginase, benzodiazepines, bupropion, cefdinir, cefpodoxime, celtriaxin, chlorambucil, cyclosporine, dexamethasone, fluphenazine, folic acid, furosemide, glucocorticoids, hydrocortisone, ibuprofen, inotocin, levetracetam, levofloxacin, lithium, meclotheramine, miremazapine, miremazapine, nicaralbumin-CD3, nelfinavir, niacin, oxazolac, olsalazine, pamidronate, pentostatin, phenytoin, quazepam, rabeprazole, sirolimus, barbifamine, thiamine, trastuzumab, triazolam
Magnesium (Serum)  
Increased by: alkaline antacids, amiloride, aminoglycosides, aspirin, calcitriol, cefotaxime, felodipine, hydroflumethiazide, lithium, magnesium salts, medroxyprogesterone, progesterone, sodium bicarbonate, tacrolimus, triamterene  
Decreased by: albuterol, aldesleukin, amphotericin B, arsenic trioxide, azathioprine, basiliximab, calcitriol, calcium gluconate, cefotaxime, chlorothiazide, chlorthalidone, cisplatin, cyclosporine, digoxin, doxorubicin, ethacrynic acid, foscarnet, furosemide, gentamicin, haloperidol, hydrochlorothiazide, hydroflumethiazide, insulin, metolazone, neomycin, oral contraceptives, pamidronate, pentamidine, prednisolone, sirolimus, tacrolimus, theophylline, thiazides, tobramycin, trastuzumab, zalcitabine

Magnesium (Urine)  
Increased by: acetzolamide, ammonium chloride, amphotericin B, bumetanide, calcitonin, chlorothiazide, cisplatin, cyclosporin A, ethacrynic acid, furosemide, gentamicin, hydrochlorothiazide, lithium, magnesium hydroxide, methyclothiazide, thiazides, torsemide, triamterene  
Decreased by: acetzolamide, amiloride, calcium gluconate, interferon alfa-2a, oral contraceptives, parathyroid extract

Methemoglobin (Blood)  
Increased by: acetaminophen, aminosalicylic acid, amyl nitrite, analgesics, antimalarials, aspirin, benzocaine, bismuth nitrate, chloramphenicol, chlorpheniramine, co-trimoxazole, dapsone, dimercaprol, furazolidone, isoniazid, isosorbide, local anesthetics, methicillin, methylene blue, metoclopramide, nitrofurantoin, nitrofurazone, nitroglycerin, phenazopyridine, phenytoin, potassium chloride, primaquine, probenecid, quinidine, quinine, sulfactamamide, sulfamethizole, sulfinilamide, sulfisoxazole  
Decreased by: sulfamethoxazole, sulfasalazine, trimethoprim

Monocytes (Blood)  
Increased by: alprazolam, ampicillin, carbenicillin, chlorpromazine, granulocyte colony-stimulating factor, griseofulvin, haloperidol, lomefloxacin, methsuximide, paroxetine, penicillamine, piperacillin, prednisone, propylthiouracil, quazepam  
Decreased by: alprazolam, glucocorticoids (transient), granulocyte colony-stimulating factor, triazolam

Oxalate (Urine)  
Increased by: ascorbic acid, bumetanide, ethylene glycol  
Decreased by: ascorbic acid, calcium carbonate

Parathyroid Hormone Assay (Plasma)  
Increased by: chemotherapy, cyclosporine, estrogen/progestin therapy, foscarnet, human growth hormone, hydrocortisone, isoniazid, ketoconazole, methyltestosterone, prednisolone, tamoxifen, valdecoxib, verapamil  
Decreased by: aluminum hydroxide, calcitriol, cimetidine, diltiazem, foscarnet, hydrocortisone, human growth hormone, hydrochlorothiazide, mannitol, medroxyprogesterone, nafarelin, nifedipine, nitrofurantoin, oral contraceptives, parathyroid hormone, pindolol, prednisone, thiazides, vitamin D

Phosphate (Serum)  
Increased by: aldesleukin, aluminium hydroxide, anabolic steroids, aspirin, azithromycin, basiliximab, bisoprolol, cefdinir, cefotaxime, dipyridamole, epoetin alfa, erythropoietin, etidronate, etretinate, famotidine, foscarnet, furosemide, human growth hormone, hydrochlorothiazide, imipramine, minocycline, nitrofurantoin, nifedipine, nitrofurantoïn, oral contraceptives, paroxetine, prednisone, sirolimus, tacrolimus, thiazides, vitamin D  
Decreased by: acetazolamide, albuterol, aldesleukin, alendronate, alkali antacids, aluminium salts, amino acids, amiodarone, anesthetic agents, anticonvulsants, azathioprine, calcitriol, calcitonin, calcium, calcium gluconate, calcium, calcitriol, carbamazepine, cefdinir, cefotaxime, dexamethasone, doxercubin, ethacrynin, etretinate, famotidine, famotidine, fritide, hydrochlorothiazide, insulin, neomycin, oral contraceptives, pamidronate, pentamidine, prednisone, propanolol, sirolimus, theophylline, triazolam, vitamin D

Plasminogen (Blood)  
Increased by: anabolic steroids, danazol, fluoxymesterone, norethandrolone, oral contraceptives, oxandrolone, oxymetholone, stanozolol  
Decreased by: alteplase, anistreplase, asparaginase, chemotherapy, dextran, gemfibrozil, streptokinase

Platelet (Blood)  
Increased by: alglucerase, amoxapine, auranofin, cefazolin, cefdinir, cefotetan, cefpodoxime, ceftobuten, ceftriaxone, cefuroxime, clindamycin, clozapine, danazol, diltiazem, dipyridamole, donepezil, epoetin alfa, etrapenem, erythropoietin, estropipate, etretinate, fludarabine, gemfibrozil, glucoctocoids, imipenem/cilastin, immune globulin, indinavir, interferon alfa-2a, isotretilnoin, lansoprazole, lithium, lomefloxacin, megestrol, mepoten, mesalamine, methylprednisolone, metoprolol, metyrosine, miconazole, moxalactam, netilmicin, ofloxacin, oral contraceptives, paroxetine, pentoxifiline, piperazine, piroxicam, propanolol, sirolimus, ticlopidine, venlafaxine, zalcitabine, zidovudine
Increased by:
- abciximab, acetaminophen, acetylazolamide, albendazole, albuterol, aldesleukin, alemuzumab, allopurinol, altretamine, aminocaproic acid, aminoglutethimide, ami噪one, amitriptyline, amoxicillin, amphotericin B, ampicillin, amrinone, anagrelide, anticonvulsants, antineoplastic agents, ardeparin, arsenic trioxide, asparaginase, aspirin, auranofin, aurothioglucose, azathioprine, azithromycin, bacampicillin, barbiturates, basiliximab, BCG vaccine, benazepril, benzthiazide, betaxolol, bicatalumide, bismuth subsalicylate, b-lowmycin, bupropion, candesartan, capreptagin, caphromycycin, captopril, carbamazepine, carbenicillin, carmustine, carvedilol, cefadroxil, cefamandole, cefazolin, cefditoren, cefixime, cefonicid, cefotetan, cefotixin, cepodoxime, cefotibutin, cefotizoxime, cefotaxime, ceftriaxone, cefuroxime, cefizine, chlorambucil, chloramphenicol, chloridiazepoxide, chloroquine, chlorothiazide, chlorpheniramine, chlorproamone, chlorpropamide, chlortetracycline, chlorthaldione, cimetidine, cladrabrine, clermastine, clindamycin, clindoflox, clofibrate, clofibrate, clopimilum, clonazepam, clopidogrel, co-trimoxazole, codeine, colchicine, cyclobenzaprine, cyclophosphamide, cyproheptadine, ctarabazine, dacarbazine, dactinomycin, dalteparin, danalox, demeclocycline, desipramine, dextroamphetamine, diamethoxazine, diacerein, didanoxine, dihydropropion, diethylosulfat, digitals, digitoxin, dilataz, diphenhydramine, disopyramide, docetaxel, donepezil, doxepin, doxorubicin, doxycycline, eflornithine, enalapril, enoxaparin, epirubicin, eprosartan, erythromycin, eseopemprazone, estramustine, etanercept, ethacrynic acid, ethchlorvynol, ethosuximide, etodronate, etoposide, etretinate, factor VII a, farnolitad, fenoprofen, flecanide, fluconazole, fluoxetine, fludarabine, fluorouracil, fluphenazine, fluorovasatin, fluvoxamine, fomiwires, fondaparinux, fosphenytoin, fusidateine, gabapentin, galantamine, ganciclovir, gatifloxacin, gemcitabine, gentamicin, glibperide, glyburide, gold, granisetron, heparitin B vaccine, hydralazine, hydrochlorothiazide, hydroxychloroquine, hydroxyurea, ibuprofen, idarubicin, ifosfamide, imatinib, imipenem/clisiast, imipramine, immunoglobulin, indinov, indinavir, indinavir, interferon alpha-2a, interferon-2k, ironotocan, isoniazid, isosorbide dinitrate, isrotetinoin, irlaconazole, ketoprofen, lamivudine, lamotrigine, lanroxifene, lepirudin, levodopa, lincomycin, levamisole, levocin, leucin, lomace, loracarbef, lovatatrin,loxapine, maprotiline, measles virus vaccine, meclotheramine, mephenicap, melfoquine, meloxicam, melphalan, meproprame, mergercaptone, meropenem, mesalamine, mesoridazine, melformin, methazolamide, meftillrin, methimazole, methotrexate, methsuximide, methylothiazide, methylalp, methylphenidate, methysergide, metroporol, metronidazole, metronixin, mexiletine, mitirinone, minocycline, mirtazapine, misoprostol, mitomycin, mitoxantrone, moricicnine, morphine, moxalaclam, mumps virus vaccine, muromonab-CD3, mycophenolate, nabumetone, naldixic acid, naproxen, netilmicin, nevirapine, niacin, nicardipine, nitrofurantoain, nitroglycerin, nizatidate, norfloxacin, nortriptiyline, nystatin, ofloxacin, olsalazine, omeprazole, onphenadrine, oxacin, oxetoxycycline, pacitaxel, pamidronate, pegaspargase, pemoline, penicillamine, penicillin, pentamidine, pentostatin, pentoxifylline, pergolide, perphenazine, phenobarbitol, phenothiazines, pindolol, piroxicam, plicamycin, polo virus vaccine, polythiazone, potassium iodide, pravastatin, pravadoline, primadone, probenecid, procainamide, procarbazone, promazine, promethazine, propafenone, propranolol, propyl-thiouracil, propritriptyline, pyrazinamide, pyrimethamine, quazepam, quinidone, quinupristin/dalfopristin, raveprazole, rafoxine, ramipril, reserpine, rifampin, risperidone, rivastigmine, rubella virus vaccine, saquinavir, sargramostim, sirolimus, smallpox vaccine, spironolactone, staudvine, streptomycin, streptococin, sulfamethoxazole, sulfasalazine, sulfisoxazole, sulfonureas, sulindac, tacrolimus, tamoxifen, temozolomide, tetracycline, thiabendazole, thiadiazides, thioeguaine, thioridazine, thiotepa, thiothixene, ticarcillin, ticcapone, tidlopin, tinzaparin, tobramycin, tocinadine, tolazamide, tolazoline, tolbutamide, tolcapone, tolphotin, topotecan, toremifene, tranylcypromine, trastuxumab, trichlormethiazide, trifluoperazine, trimethoprime, trimetrexate, trimipramine, triflunoxacin, uracil mustard, valdecoxib, valganciclovir, valproic acid, vancomycin, vinblastine, vincristine, vinorelbine, zidovudine, zolmitriptan

**Porphobilinogen (Urine)**
- Increased by: aminosaliclyc acid, anticonvulsants, barbiturates, cascarra, chloridiazepoxide, chlorpropamide, griseofulvin, imipenem/clisiast, meprobamate, oral contraceptives, phenothiazines, pentazocine, phenytoin, probeaine, tolbutamide

**Porphyrins (Urine)**
- Increased by: antipyretics, barbiturates, chloral hydrate, chloridiazepoxide, chlorpropamide, ciprofloxacinc, diazepam, ergot preparations, gresofulvin, nalidixic acid, hydantoin derivatives, norfloxacin, meprobamate, methylap, hydroxychloroquine, ofloxacin, oxetoxycycline, pentazocine, phenazopyridine, progastim derivatives, sulfamethoxazole, tetracycline, vitamin K
- Decreased by: oral contraceptives

**Potassium (Serum)**
- Increased by: ACE inhibitors, aldesleukin, amiloride, aminocaproic acid, aminoglutethimide, ammonium chloride, amphetamine B, atenolol, azathioprine, azithromycin, basiliximab, benazepril, betaxolol, bispnlol, candesartan, cannabis, captopril, cefdin, cefoxime, cefuroxime, clofibrate, cyclosporine, danazol, dexamethasone, digoxin, doxorubicin, enalapril, epsilon alfa, erythropyriinet, etretinate, foleodipine, fosphetyoin, indomethacin, isonizid, ketocaonazole, ketoroloc, labetolad, lisinopril, lithium, LMW heparin, lovastatin, manitol, mehiolin, methyltestosterone, metoprolol, micardis, mycophenolate, naproxen, netilmicin, nifedipine, nonsteroidal anti-inflammatory drugs, norfloxacin, ofloxacin, paroxetine, penicillin, pentamidine, perindopril, pindolol, piroxicam, propranolol, quinapril, quinupristin/dalfopristin, ramipril, risedrone, sirolimus, somatropin, spironolactone, succinylcholine, sulindac, tacrolimus, timolol, trandolapril, triamterene, trimethoprism, valsartan, venlafaxine, zalcitabine
Increased by:
- acetzolamide, albuterol, aldosterone, amiodarone, ammonium chloride, amphotericin, arsenic trioxide, aspirin, azathioprine, basiliximab, betamethasone, betaxolol, bisacodyl, bumetanide, candesartan, capreomycin, captopril, carbamazepine, carbenicillin, carvedilol, cathartics, cephalaxin, chloroquine, chlorothiazide, chlorothalidone, cidofovir, cisplatin, corticosteroids, corticotropin, cortisone, dexamethasone, digoxin immune fab, diuretics, dobutamine, donepezil, doxazosin, doxorubicin, enalapril, epoprostenol, ethacrynic acid, etretinate, fluconazole, furosemide, fludrocortisone, foscarnet, fosinopril, fosphenytoin, furosemide, gentamicin, hydrochlorothiazide, imatinib, indapamide, insulin, itraconazole, ketoprofen, laxatives, levodopa, lithium, lomeflaxacin, methazolamide, methyclothiazide, methylprednisolone, metoclopramide, melatonin, moxalactam, mycophenolate, nabumetone, naproxen, neomycin, nifedipine, noscapine, ondansetron, pamidronate, paroxetine, penicillin, perphenazine, phenytoin, prednisolone, tolvaptan, tramadol, trifluoperazine, vancomycin, vidarabine, zalcitabine

Potassium (Urine)
Increased by:
- acetzolamide, ammonium chloride, antibiotics, aspirin, betamethasone, bumetanide, calcitonin, carbenicillin, cathartics, chlorothalidone, corticosteroids, corticotropin, cortisone, dexamethasone, diuretics, ethacryninc acid, fenoldopam, fludrocortisone, gentamicin, hydrochlorothiazide, hydrocortisone, indomethacin, isosorbide, levodopa, lithium, mafenide, methylothiazide, methylothiazide, nalidixic acid, nifedipine, potassium, propranolol, ranitidine, sodium bicarbonate, spironolactone, streptozocin, tacrolius, terbutaline, tetracycline, theophylline, thiazides, ticarcillin, tobramycin, triamterene, triflurazethidiazide, trimethoprim, venlafaxine

Pregnanediol (Urine)
Increased by:
- corticotropin, phenazopyridine, tamoxifen

Progestosterone (Plasma)
Increased by:
- corticotropin, ketoconazole, progesterone, tamoxifen

Prolactin (Serum)
Increased by:
- amitriptyline, amoxapine, butorphanol, carbipoda, chlorpromazine, cimetidine, citalopram, clozapine, danazol, desipramine, diethylstilbestrol, enalapril, fenoldopam, fluphenazine, flusofamid, ganadotropin-releasing hormone, growth hormone-releasing hormone, haloperidol, imipramine, insulin, interferon, interleukin, labetalol, loxapine, megestrol, mirtal, methylprednisolone, melatonin, methylprednisolone, morphine, nitrous oxide, oral contraceptives, parathyroid hormone, perphenazine, phenytoin, prochlorperazine, promazine, ranitidine, reserpine, risperidone, thioridazine, thiopental, thyrutropin-releasing hormone, trifluoperazine, trimipramine, tumor necrosis factor, verapamil

Decreased by:
- acetzolamide, amiloride, anesthetic agents, carbamazepine, cyclosporin A, diazoxide, felodipine, ketoconazole, levarterenol, niasin, ramipl, sulfamethoxazole, trimethoprim

Protein (Cerebrospinal fluid)
Increased by:
- ampicillin, aspirin, cefotaxime, chloramphenicol, ibuprofen, imipramine, methicillin, methotrexate, oxytetracycline, penicillin, perphenazine, phenothiazines, procaaine, radiographic agents, streptomyacin, sulfadiazine, sulfanilamide, sulfisoxazole, tetracycline, tolbutamide, trifluoperazine, vancomycin

Decreased by:
- aceltominophen, cefotabrine, cefotaxime, dexamethasone, ranitidine, rifampin, ropinrole, tamoxifen, toremifene, valproic acid

Protein (Urine)
Increased by:
- acetzolaminophen, acetzolamide, aldeleukin, aminophylline, aminosalicylic acid, amphotericin B, ampicillin, arsenicals, ascorbic acid, asparaginase, aspirin, auranofin, averothioglucose, bactracin, benazepril, betaxolol, bicarbonate, bismuth subsalicylate, capreomycin, carbamazepine, castor oil, cefaclor, cefamandole, cefdinir, cefalothin, chloral hydrate, chlorhexidine, chloromycetin, chlorpromazine, chlorpheniramine, chlorpropamide, chlorothalidone, cidofovir, cisplatin, clindamycin, colchicine, codeine, colistin, corticosteroids, corticotropin, cyclosporine, dantrolene, demeclocycline, diethylstilbestrol, enalapril, furosemide, gentamicin, hydrochlorothiazide, imatinib, indapamide, insulin, itraconazole, ketoprofen, laxatives, levodopa, lithium, lomeflaxacin, methazolamide, methyclothiazide, methylprednisolone, metoclopramide, melatonin, moxalactam, mycophenolate, nabumetone, naproxen, neomycin, nifedipine, noscapine, ondansetron, pamidronate, paroxetine, penicillin, perphenazine, phenytoin, prochlorperazine, promazine, ranitidine, reserpine, risperidone, thioridazine, thiopental, thyrutropin-releasing hormone, trifluoperazine, trimipramine, tumor necrosis factor, verapamil

Decreased by:
- acetzolamide, amiloride, anesthetic agents, carbamazepine, cyclosporine, dexamethasone, ranitidine, rifampin, ropinrole, tamoxifen, toremifene, valproic acid

Renin (Plasma)
Elevated by:
- captopril, furosemide

Decreased by:
- oral contraceptives
**Sodium (Serum)**

Increased by: aldesleukin, amiloride, amino acids, ampicillin, anabolic steroids, betamethasone, cannabinoids, carbamazepine, cefotaxime, chlorothalidone, cholestyramine, clonidine, corticosteroids, cortisone, diazoxide, doxorubicin, etretinate, fluocortisone, guanethidine, human growth hormone, hydrocortisone, isosorbide, mannitol, methyclothiazide, methylprednisolone, nifedipine, oxymetholone, penicillin, phenytoin, propranolol, stanozolol, sulfonylureas, tolbutamide

Decreased by: aldesleukin, amiloride, aminoglutethimide, ammonium chloride, amphotericin, atovaquone, benazepril, captopril, carbamazepine, carvedilol, cathercids, chlorothiazide, chloropropamide, chlorthalidone, cisplatin, clozapine, cyclophosphamide, dapsone, desmopressin, diuretics, doxepin, doxorubicin, esomeprazole, ethacrynic acid, etretinate, fluoxetine, fluvoxamine, foscarnet, furosemide, gentamicin, gliburide, glycerin, haloperidol, hydrochlorothiazide, hydroflumethiazide, indomethacin, isosorbide dinitrate, itraconazole, ketocanozole, ketoprofen, ketorolac, laxatives, lisinopril, lithium, mannitol, methyclothiazide, methylprednisolone, metolazone, miconazole, morphone, nicardipine, nifedipine, nisoldipine, nonsteroidal anti-inflammatory drugs, omeprazole, paroxetine, pentostatin, phenoxymazine, pimozide, polythiazide, propafenone, quinethazone, ramipril, risperidone, sertraline, sirolimus, sodium bicarbonate, somatostatin, spironolactone, sulfonylureas, tacrolimus, theophylline, thiazides, ticlopidine, tobramycin, trastuzumab, triamterene, trimethoprim, trimethrexate, trovafloxacin, valproic acid, vasopressin, venlafaxine, vidarabine, vincristine, zalcitabine

**Sodium (Urine)**

Increased by: ACE inhibitors, acetazolamide, amiloride, ammonium chloride, aspirin, atenolol, benzthiazide, bumetanide, calcitonin, captopril, carvedilol, chlorothiazide, chlorthalidone, cisplatin, clofibrate, cyclohexiazide, dexamethasone, digitalis, doxepin, enalapril, ethacrynic acid, felodipine, fentanyl, furosemide, hydrocortisone, hydroflumethiazide, ifosfamide, indomethacin, insulin, isosorbide, levodopa, lithium, losartan, mannitol, methyclothiazide, metoprolol, metolazone, niacin, niacinamide, oral contraceptives, parathamine, parathyroid extract, polythiazide, progestosterone, quinethazone, secrecin, spironolactone, tetraclycline, thiazides, tersemine, trimcinolone, triamterene, triclothemide, trimethoprim, verapamil, vincristine

Decreased by: anesthetic agents, carbamazepine, corticosteroids, cortisone, cyclosporine, diazoxide, etodolac, fluoxetine, ibuprofen, indomethacin, insulin, ketorolac, levartenenol, lithium, methylprednisolone, naproxen, nifedipine, octreotide, omeprazole, propranolol, ramipril

**Somatomedin C (Insulin-like Growth Factor I) (Plasma)**

Increased by: AINE inhibitors, acetazolamide, amiloride, ammonium chloride, aspirin, atenolol, benzthiazide, bumetanide, calcitonin, captopril, carvedilol, chlorothiazide, chlorthalidone, cisplatin, clofibrate, cyclohexiazide, dexamethasone, digitalis, doxepin, enalapril, ethacrynic acid, felodipine, fentanyl, furosemide, hydrocortisone, hydroflumethiazide, ifosfamide, indomethacin, insulin, isosorbide, levodopa, lithium, losartan, mannitol, methyclothiazide, metoprolol, metolazone, niacin, niacinamide, oral contraceptives, parathamine, parathyroid extract, polythiazide, progestosterone, quinethazone, secrecin, spironolactone, tetraclycline, thiazides, tersemine, trimcinolone, triamterene, triclothemide, trimethoprim, verapamil, vincristine

Decreased by: estrogens, ethinyl estradiol, methimazole, octreotide, oral contraceptives, tamoxifen

**Testosterone (Serum)**

Increased by: anabolic steroids, barbiturates, bromocriptine, Casodex, cimetidine, clomiphene, danazol, estrogen/progestin, finasteride, flutamide, goserelin, leuprolide, levonorgestrel, mifepristone, nafarelin, nalmefene, naloxyne, nitilumate, oral contraceptives, phentoyin, pravastatin, rifampin, tamoxifen, valproic acid

Decreased by: carbamazepine, cimetidine, conjugated estrogens, cyclophosphamide, danazol, dexamethasone, diazoxide, diethylstilbestrol, digoxin, estradiol valerate, fenoldopam, finasteride, follicle-stimulating hormone, gemfibrozil, goserelin, interleukin, ketocazone, leurozole, leuprolide, magnesium, medroxyprogesterone, metformin, methylprednisolone, nafarelin, octreotide, oral contraceptives, pravastatin, prednisone, spironolactone, stanozolol, tamoxifen, tetracycline, verapamil

**Thrombin Time (Blood)**

Increased by: anistreplase, asparaginase, streptokinase, urokinase

Decreased by: dextran, LMW heparin

**Thyroglobulin (Serum)**

Increased by: amiodarone

Decreased by: neomycin, thyroxine

**Thyroid-Stimulating Hormone (TSH) (Serum)**

Increased by: amiodarone, atenolol, calcitonin, carbamazepine, chlorpromazine, clomiphene, conjugated estrogens, ferrous sulfate, iodide, levothyroxine, lithium, lovastatin, methimazole, metoprolol, morphon, phenothiazine, potassium iodide, prazosin, prednisone, propranolol, radiographic agents, rifampin, sumatriyan, tamoxifen, thyrotropin-releasing hormone, valproic acid

Decreased by: amiodarone, anabolic steroids, aspirin, carbamazepine, clofibrate, corticosteroids, danazol, fenoldopam, growth-releasing hormone, hydrocortisone, interferon alfa-2, levothyroxine, nifedipine, octreotide, somatostatin, thyroxine, troleandomycin

**Thyroxine-Binding Globulin (TBG) (Serum)**

Increased by: carbamazepine, clofibrate, diethylstilbestrol, ethyrypoetin, estrogens, mestranol, oral contraceptives, perphenazine, phenothiazines, progestosterone, ralfoxiene, tamoxifen

Decreased by: anabolic steroids, asparaginase, clenostipol, corticosteroids, corticotropin, cortisone, danazol, fluoxymesterone, methyltestosterone, nandrolone, norethandrolone, norethindrone, oxymetholone, phenytoin, prednisone, propranolol, stanozolol

**Triiodothyronine Uptake (Serum)**

Increased by: aminogluthetimide, aminosalicylic acid, amiodarone, anabolic steroids, aspirin, barbiturates, chlorpropamide, coab, corticosteroids, Coumadin, fluoxymesterone, furosemide, methyltestosterone, metoprolol, miotane, nandrolone, Norplant, orphenadrine, oxymetholone, penicillin, phenytoin, propranolol, stanozolol, sulfonfonylures, tamoxifen, thyroxin, tolbutamide
Increased by:
- amiodarone, chloridiazepoxide, clofibrate, diazepam, diflunisal, estradiopate, fluoxymesterone, lithio, medroxyprogesterone, methimazole, oral contraceptives, perphenazine, phenothiazines, thiazides

Triiodothyronine (T₃) (Serum)

Increased by:
- amiodarone, amphetamine, clofibrate, erythropoitin, estradiopate, fluorouracil, insulin, levothyroxine, mestranol, methadone, opiates, oral contraceptives, phenothiazines, phenytoin, propylthiouracil, ranitidine, tamoxifen, terbutaline, thyrotropin-releasing hormone, valproic acid

Decreased by:
- amiodarone, anabolic steroids, asparaginase, aspirin, atenolol, carbamazepine, cholestyramine, cimetidine, clophemene, clopimramine, cotrimoxazole, corticosteroids, danazol, dexamethasone, diclofenac, furosemide, glucocorticoids, hydrocortisone, interferon α-2a, iodide, isoretinoin, lithium, methimazole, metoprolol, mitotane, naproxen, netilmicin, oral contraceptives, penicillamine, phenobarbital, phenytoin, potassium iodide, prednisone, propranolol, propylthiouracil, radiographic agents, salsalate, somatostatin, stanozolol, sulfonylureas, theophylline

Thyroxine (T₄) Free (Serum)

Increased by:
- amiodarone, aspirin, carbamazepine, danazol, enoxaparen, erythropoitin, furosemide, levothyroxine, phenytoin, propranolol, propylthiouracil, radiographic agents, tamoxifen, thyroxine, valproic acid

Decreased by:
- amiodarone, anabolic steroids, carbamazepine, clofibrate, corticosteroids, estrogen therapy, isoretinoin, levothyroxine, lithium, mestranol, methadone, methimazole, norethindrone, octreotide, oral contraceptives, phenobarbital, phenytoin, ranitidine

Triglycerides (Serum)

Increased by:
- acetylsalicilc acid, amiodarone, ardeparin, aspirin, atenolol, bisoprolol, beta-blockers, carbamazepine, carvedilol, Casodex, chloridane, chlorothiazide, chlortalidone, cholestyramine, colchicine, cyclosporin, danazol, didanosine, enalapril, estrogen/progestin therapy, etrogestens, estrogens, fluconazole, flutamide, furosemide, glucocorticoids, glycine, goserelin, hydrochlorothiazide, interferon α-2a, isoretinoin, linaconazole, labetalol, levotyroxine, methylclohiside, methylodopa, metoprolol, miconazole, mirtazapine, nafarelin, norfloxacain, oflloxacin, oral contraceptives, perindolol, pindolol, polythiuride, prazosin, prednisolone, prednison, propranolol, radioactive iodine, riperidone, ronavir, simvastatin, sotalol, spironolactone, tamoxifen, tenovir, thiazides, ticlopidine, timolol, tretinoin, trichormethiazide, warfarin, zalciabine

Decreased by:
- acarbose, ACE inhibitors, acetylsalicilc acid, amiodarone, amloidine, anabolic steroids, ascorbic acid, asparaginase, atorvastatin, azathioprine, bisoprolol, captopril, carvedilol, cervatistatin, chenolidol, chlortalidone, cholestyramine, clofibrate, colestopil, danazol, dexfenfuramine, diiltiazem, doxazosin, enalapril, estrogen therapy, fenofibrate, flaxseed oil, fuvastatin, gemfibrozil, glyburide, HMG CoA-reductase inhibitors, hydroxychlororouquine, hydroxyurea, indomethacin, insulin, ketoconazole, levodopa, levonorgestrel, levothyroxine, LMW heparins, lovastatin, medroxyprogesteron, metformin, metimazole, methadone, methylclobidz, micotin, nicardipine, nifdefine, nifdefine, nifdefine, nifdefine, oxandrolone, pentoxyfylino, pindolol, pravastatin, prazosin, prednisolone, probucol, psylgium, simvastatin, stanozol, terazosin, trifalotazine, unfractionated heparin, verapamil

Uric Acid (Serum)

Increased by:
- acetaminophen, acetazolamide, aldesleukin, amiloride, ampicillin, anabolic steroids, antineoplastic agents, asparaginase, ascorbic acid, aspirin, atenolol, azathiorpil, basilixmin, benzthiazide, betaxolol, bisoprolol, bumetanide, busulfan, calcitriol, candesartan, capreomycin, carvedilol, chloridiane, chorambucil, chlorothiazide, chlortalidone, cimetidine, cispalton, clopidogrel, clozcape, cycloporsine, cytarabine, dextran, diazoxide, diclofenac, didanosine, dilatiazem, diuretics, donepezil, doxazosin, doroxurbin, epoetin αf, ethacrynic acid, ethambol, etoposide, fludarabine, furosemide, gentamicin, goserelin, granulocyte colony-stimulating factor, hydralazine, hydrochlorothiazide, hydrofluemethiazide, hydroxyurea, ibuprofen, indomethacin, irbesartan, isonizad, isosorbide dintrate, isoretinoin, lansoprazole, leuproide, levaterenol, levodopa, lisinopril, lomefloxacin, losartan, mechlorothamine, mercaptopurine, mesalamine, methicillin, methotrexate, methylothiazide, metomycin, metoprolol, mitomycin, nifdefine, nifdefine, nifdefine, nifdefine, oxandrolone, nifeprin, niai, niai, nac thromidine, nacitridine, nicotine, nifdefine, nifdefine, nifdefine, oxandrolone, pentoxyfylino, pindolol, pravastatin, prazosin, prednisolone, probucol, psylgium, simvastatin, stanozol, terazosin, trifalotazine, unfractionated heparin, verapamil

Decreased by:
- acarbose, ACE inhibitors, acetylsalicilc acid, amiodarone, amloidine, anabolic steroids, ascorbic acid, asparaginase, atorvastatin, azathioprine, bisoprolol, captopril, carvedilol, cervatistatin, chenolidol, chlortalidone, cholestyramine, clofibrate, colestopil, danazol, dexfenfuramine, diiltiazem, doxazosin, enalapril, estrogen therapy, fenofibrate, flaxseed oil, fuvastatin, gemfibrozil, glyburide, HMG CoA-reductase inhibitors, hydroxychlororouquine, hydroxyurea, indomethacin, insulin, ketoconazole, levodopa, levonorgestrel, levothyroxine, LMW heparins, lovastatin, medroxyprogesteron, metformin, metimazole, methadone, methylclobidz, micotin, nicardipine, nifdefine, nifdefine, nifdefine, oxandrolone, pentoxyfylino, pindolol, pravastatin, prazosin, prednisolone, probucol, psylgium, simvastatin, stanozol, terazosin, trifalotazine, unfractionated heparin, verapamil

Uric Acid (Urine)

Increased by:
- acetohexamide, acetaminophen, ascorbic acid, asparaginase, aspirin, busulfan, chlorothiazide, clofibrate, corticotropin, cortisone, coumadin, diethylthiobestrol, diflunisal, dobutamine, doxazosin, enalapril, ethacrynic acid, etodolac, fenofibrate, griseofulvin, guaiifenesin, ibuprofen, indomethacin, levodopa, lisinopril, lithium, mannitol, mechlorothamine, mfenamic acid, methotrexate, methylodopa, phenothiazines, prednisolone, probenecid, radiographic agents, salicylate, sertralin, spironolactone, sulfamethohxazole, sulfinpyrazine, ticarcilin/clavulanate, verapamil, vinblastine

Decreased by:
- acetohexamide, allopurinol, amiloride, amloidine, ascorbic acid, aspirin, azathiorpil, cannabis, cannola oil, chloridiazepoxide, chlorpromazone, cdiofovir, clofibrate, corticosteroids, corticotropin, cortisone, Coumadin, diethylthiobestrol, diflunisal, dobutamine, doxazosin, enalapril, ethacrynic acid, etodolac, fenofibrate, griseofulvin, guaiifenesin, ibuprofen, indomethacin, levodopa, lisinopril, lithium, mannitol, mechlorothamine, mfenamic acid, methotrexate, methylodopa, phenothiazines, prednisolone, probenecid, radiographic agents, salicylate, sertralin, spironolactone, sulfamethohxazole, sulfinpyrazine, ticarcilin/clavulanate, verapamil, vinblastine
Decreased by: acetazolamide, allopurinol, aspirin (low dose), azathioprine, bumetanide, chlorothiazide, chlorthalidone, diazoxide, ethacrynic acid, ethambutol, furosemide, hydrochlorothiazide, levaterenol, niacin, probenecid, pyrazinamide, salicylates (low doses), thiazides

**Urine Specific Gravity**

Increased by: dextran, isotretinoin, penicillin, radiographic agents

Decreased by: lithium

**Urobilinogen (Stool)**

Increased by: amyl nitrate

Decreased by: antibiotics, aspirin, chloramphenicol, chlordiazepoxide, chlorpromazine, erythromycin, methimazole, nalidixic acid, neomycin, oral contraceptives, oxymetholone, prochlorperazine, promazine, sulfamethoxazole, sulfinpyrazone, tetracycline, thiabendazole, thiazides, tolbutamide, trifluoperazine

**Vanillylmandelic Acid (VMA) (Urine)**

Increased by: aminosalicylic acid, chlorpromazine, disulfiram, guaifenesin, guanethidine, insulin, isoproterenol, labetalol, levaterenol, levodopa, lithium, methocarbamol, methyldopa, nalidixic acid, nifedipine, nitroglycerin, oxytetracycline, phenazopyridine, prazosin, prochlorperazine, rauwolfia, reserpine

Decreased by: clonidine, disulfiram, fluvoxamine, guanethidine, guanfacine, imipramine, levodopa, MAO inhibitors, methyldopa, morphine, phenoxyzine, phenothiazines, radiographic agents, reserpine

**Vitamin B₁₂ (Serum)**

Increased by: chloral hydrate, omeprazole

Decreased by: anticonvulsants, ascorbic acid, cholestyramine, chlorpromazine, colchicine, metformin, neomycin, octreotide, oral contraceptives, ranitidine, rifampin

**BIBLIOGRAPHY**

Appendix K Protocols for Hair, Nails, Saliva, Sputum, and Breath Specimen Collection

Specimen collection of hair, nails, and saliva are easy to obtain and noninvasive. Breath specimen collection methodologies can range from the simple to the complex, and sputum collection involves specific protocols. Sputum specimens are generally easy to obtain, noting that specific procedures are followed. Results from analysis of these samples aid in diagnosing several diseases and may be used as evidence in legal situations. It is imperative that all protocols are followed precisely to avoid interference in evaluating test results.

Samples of clean hair and nails can be analyzed for evidence of fungal infections, abnormal concentrates of toxic and nutrient minerals (selenium), heavy metals (mercury and lead), therapeutic drugs, and illegal drugs. High levels of some elements are caused by exposure to industrial wastes and by contaminated drinking water. The limit of detection of most drugs in hair is 0.1 ng/mL or higher.

Indications for testing of hair samples for drugs:

1. As a marker of toxin exposure
2. To monitor parolees and probationers
3. To validate drug self-reporting
4. To identify in utero drug use
5. To assess pattern of drug use (at 1-month intervals)
6. To aid in drug treatment programs
7. Testing in workplace
8. To evaluate parents’ drug use in child custody cases
9. As forensic evidence after death
10. Compliance with nonpsychotropics (eg, antihypertensives or antibiotics) for months to years

Oral fluid saliva specimens may be used to identify high levels of IgG in the gingival crevice. This crevice secretes a fluid or transudate that contains a relatively high concentration of IgG. The procedure for oral fluid saliva specimen collection involves obtaining a gingival cervical fluid (ie, mucosal transudate) specimen by rubbing the oral mucosa (ie, lining of the gum and cheek) in the mouth with a cotton collection pad on a stick.

Sputum specimens are examined to identify pathogens or conditions related to the respiratory system. Pertinent symptoms may include cough with or without sputum production, fever, chest pain, shortness of breath, and fatigue. Sputum specimens can also provide clues about antibiotic or drug sensitivity, the best course of treatment, and the effectiveness of treatment.

Exhaled breath specimens are obtained to identify *H. pylori* infection, detect alcohol, and monitor hormones and other byproducts of abnormal metabolism by using special analyzers or kits that may involve specific instructions.

REFERENCE VALUES

Normal

1. Hair and nails, negative for presence of:
   a. Fungus
   b. Heavy metals
   c. Therapeutic and illegal drugs
   d. Anabolic steroids
2. Sputum, negative for presence of:
   a. TB
   b. Streptococcus
   c. Staphylococcus
   d. Other pathogens
3. Saliva, negative for presence of:
   a. Alcohol
   b. Tobacco
c. Therapeutic and illegal drugs

d. Hepatitis A, B, C

e. Cancer

f. Diabetes

g. Autoimmune diseases and infections

h. Hormone levels outside normal ranges (see each specific hormone in Chapter 6).

4. Breath, negative for presence of:

a. Urease in *H. pylori* infection

b. Alcohol and other drugs and toxins

c. There is no significant change in breath specimens after ingestion of substance such as lactose.

**PROCEDURES**

1. Hair Sampling

a. Use extreme care in obtaining hair samples. Wear gloves and follow established protocols. Hair should be shampooed and free of oil, conditioners, hair spray, and gels. Clip the hair close to the scalp or skin. Color-treated hair is usually acceptable; in this instance, pubic hair is preferred.

b. Obtain hairs from the correct sites: beard, mustache, axilla, genital area, and scalp. Inform the patient not to use any deodorant, powder, or lotion after shampooing or bathing until sampling is done. Use sterile scissors or instruments when cutting hair or nails.

c. Obtain a 10-g specimen of hair so that drugs such as opiates, cocaine, methadone, and amphetamines can be extracted after washing and decontamination with substances such as acetone, methylene chloride, or methanol.

d. Be aware that hair is most often collected from the back of the head (nape of the neck).

e. Cut hair as close as possible to the scalp and store in a dry tube.

f. Tie a “pencil-thick” specimen of hair with string before cutting to a length of 6 cm (if possible).

2. Nail Sampling

a. Wear gloves and follow established protocols.

b. Clip nails close to the cuticle; toenails are preferred. Before clipping, thoroughly wash and dry toenails or fingernails.

3. Both Hair and Nail Samplings

a. Transfer specimens to the laboratory in a special envelope with a biohazard label or a metal-free, screw-top plastic container.

b. Document the type and amount of specimen, site of hair or nail sampling, tests ordered, disposition of specimen, hair color, and if chemically treated, condition of nails (eg, soft, gangrenous), appearance of follicle at hair shaft, the time collected, and the relevant skin conditions (eg, scaling, dermatitis, inflamed, reddened).

4. Saliva Sampling

a. Wear gloves. Obtain a special testing kit. The kit usually includes a specially treated cotton pad on a nylon stick and a vial containing a nontoxic preservative solution. Saline solution in the pad facilitates oral fluid sample absorption.

b. Follow the established procedure precisely. Place the pad between the lower cheek and gum, rub it back and forth gently but firmly until moistened, and then leave it in place for at least 2 minutes up to a maximum of 5 minutes. Avoid salivary gland injury. After the elapsed time, remove the treated pad from the mouth, and insert it all the way to the bottom of the specimen vial that contains special antimicrobial preservative solution. Hold the vial upright to prevent spillage. Bend the pad stick handle until the upper half snaps off. Discard the broken piece, leaving the sampling pad inside the vial.

c. When placing the stopper cap on the vial, be sure to press the cap completely into the vial to prevent leakage.

d. Affix a signed and dated tamper-evident tape over the cap and down both sides of the vial.

e. Place the specimen container in a resealable biohazard bag and label it properly.

f. Transport the specimen to the laboratory as soon as possible. In the laboratory, some saliva samples are quick frozen (ie, cortisol, estriol, estradiol, progesterone, and testosterone).

g. Document the patient's name, type of specimen, test ordered, date and time of collection, and other pertinent observations.

5. Procedures for Sputum Sampling

a. Remember that sputum specimens must come from the bronchi. Postnasal secretions or saliva are unacceptable. Expectoration; ultrasonic nebulization, chest physiotherapy, nasotracheal or tracheal suctioning, and bronchoscopy are methods used to obtain sputum and bronchial specimens. Early morning specimens are the best.

b. Instruct the patient to remove dentures, rinse the mouth with water, and gargle, if possible.

c. The patient should first clear the nose and throat, take three or four deep breaths, perform a series of short coughs, and then inhale deeply and cough forcefully to raise a sputum specimen.

d. The sputum should be expectorated into a sterile container with the proper preservative if indicated. A 2- to 3-mL sample is adequate. Place the sealed container into a leak-proof biohazard bag and transfer it to laboratory after labeling properly.

e. Sputum specimens are usually not refrigerated and should be taken to the laboratory as soon as possible. Include the pertinent information, such as type of specimen, appearance, preservative, tests ordered, date and time of collection, and disposition of specimen.

f. Document the specimen appearance and the patient's response to the procedure.

6. Breath Sampling

a. Wear gloves. Use special testing kit that is needed. Follow instructions carefully; these instructions are supplied with the kit. These procedures require special training for most types of tests.

b. Some kits are disposable, contain a breath analyzer scanner, and are designed for one-time use (eg, alcohol).

c. All collections, both single and multiple, are clinician supervised. The patient inhales normally and exhales into the tube-type mouthpiece.

d. The special kit for lactose deficiency testing contains a mouthpiece attached to a special bag, vacuum-sealed collection tubes, and lactose for oral ingestion and dilution.
e. In the collection procedure for the lactose breath test, the exhaled breath is collected in a special bag and/or vacuum-sealed tubes (baseline value) both before and after drinking a solution of lactose and water (1, 2, and 3 hours). The patient must drink the lactose solution within 5 minutes of the first sip, fasting as required for lactose tests.

f. Breath analysis has two phases:
   1. Gas phase <5cc/mL
   2. Liquid phase (condensate)

### Clinical Alert

1. Urine samples provide information about short-term drug use; hair provides information about long-term drug use.
2. A challenged urinalysis from an employee should be investigated with hair analysis.

### Clinical Alert

For all specimen collecting, wear gloves and use standard precautions and precise procedures. Document type and amount of specimens, site of hair and nail sampling, appearance of nails (soft, gangrenous), appearance of saliva, sputum, relevant symptoms (eg, skin inflamed, coughing, shortness of breath), time of collection (eg, breath alcohol at 10:12 a.m.).

### NOTE

Some devices (eg, OmniSal) employ a different collection method in which a cotton pad is placed under the tongue. An indicator in the collecting device changes color when an adequate amount of oral fluid has been collected.

### Clinical Alert

In some unionized workplaces, saliva specimens are tested once a month or as often as two times a day as part of “Last Chance” agreements, or when there is reasonable suspicion of alcohol abuse. If the workplace saliva alcohol outcome is 20 mg/dL (4.3 mmol/L), the employee is sent by taxi for breath alcohol test at an emergency room or prepared clinic.

### INTERFERING FACTORS

Unsatisfactory sputum samples include “dry” specimens (ie, saliva samples without actual sputum) or contaminated specimens.

### CLINICAL IMPLICATIONS

1. Hair and Nail Sampling:
   a. Abnormal findings reveal presence of chronic heavy metal exposure and monitored levels present in the body as part of a therapeutic regimen
   b. Presence of fatty acid ethyl esters (FAEEs) in hair is a marker of long-term alcohol abuse
   c. Presence of hair and nail fungal infections
   d. Identification of illegal drug use
   e. Mother and newborn infant hair tested corroborates indications of mother’s drug use during pregnancy
   f. In postmortem assays, drug use determined as part of history in determining the cause of death
   g. For pre-employment and ongoing evaluations in the workplace

2. Saliva Sampling: Normal findings include:
   a. Presence of alcohol and drugs of abuse (eg, amphetamines, barbiturates, benzodiazepines, caffeine, cocaine, inhalants, LSD, marijuana, opiates, tobacco)
   b. Presence of HIV infection
   c. Presence of hepatitis A, B, and C; *Helicobacter pylori* infection; autoimmune disease; cancer (eg, carcinoembryonic antigen, prostate-specific antigen, CA 125 antigen); diabetes (types 1 and 2); presence of therapeutic drugs and other drugs (ie, saliva drug concentrates are lower than those in urine or blood plasma); hormone levels above or below expected outcomes for patient, (eg, cortisol, testosterone, progesterone, prolactin, DHEA); anabolic steroids; ABO blood group typing identified
   d. Specimen contains findings or results that may be useful for research or investigational purposes (Table K.1).

### Table K.1 Saliva to Plasma Ratios for Common Drugs

<table>
<thead>
<tr>
<th>Drug</th>
<th>Saliva/Plasma Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amphetamine</td>
<td>2.76</td>
</tr>
<tr>
<td>Antipyrine</td>
<td>1</td>
</tr>
<tr>
<td>Buprenorphine</td>
<td>0.05–0.41</td>
</tr>
<tr>
<td>Caffeine</td>
<td>0.6–1.0</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>0.1–0.3</td>
</tr>
<tr>
<td>Chlorpromazine</td>
<td>0.4</td>
</tr>
<tr>
<td>Clorazepate</td>
<td>5.78</td>
</tr>
<tr>
<td>Cocaine</td>
<td></td>
</tr>
<tr>
<td>Crack, 42 mg</td>
<td>1.5–17.0</td>
</tr>
<tr>
<td>Intranasal, 42 mg</td>
<td>0.7–13</td>
</tr>
<tr>
<td>IV, 25 mg</td>
<td>0.5–3.5 (average, 1.3)</td>
</tr>
</tbody>
</table>
**Clinical Alert**

If the test purpose is to detect HIV, to identify alcohol, illegal drugs, or to establish paternity, a chain of custody protocol is implemented.

1. The patient must sign a consent form and the tamper-evident tape that protects the integrity of the specimen.
2. Provide the patient’s name, date of birth, and collection date on the vial.

**INTERVENTIONS**

1. Hair and Nail Sampling
   a. Pretest Patient Preparation
      1. Assess and document signs and symptoms of drug presence or toxic exposure. Include the geographic location, water supply sources and quality, pesticide use, industrial waste exposure, food contaminants, and current medications.
      2. Explain how keratin is laid down in bone, hair, and nails, and how this relates to the test purpose. Toenail may give more accurate measures of exposure to certain substances such as selenium. It takes 3 months for the nail on the little toe to grow out and a year for the big toe to grow out.
      3. Explain how mycotic organisms (eg, ringworm) attack the keratin.
      4. If testing is done for illegal or improperly used drugs, follow chain of custody protocols.
      5. Check to see if dandruff shampoos have been used; they contain selenium and can skew results.
      6. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
   b. Posttest Patient Care
      1. Inform the patient regarding anticipated times the test results reports are expected.
      2. Interpret test results and counsel appropriately regarding any follow-up testings or treatment intervention.

---

*Saliva pH of 6.8, serum pH of 7.4

3. Sputum Sampling

4. Breath Sampling:
   a. Abnormal results reveal presence of alcohol, toxins produced by bacteria, or hydrogen ($H_2$) after ingestion of lactose and electrolytes.
   b. A flat curve is seen in most persons with lactose deficiency who are not diabetic.
2. Saliva Sampling
   a. Pretest Patient Preparation
      1. Explain the purpose, procedure, and any interfering factors related to oral fluid specimen collection. There should be no candy or gum in the mouth. Recent food intake, smoking, use of oral hygiene, and treatment with anticholinergic drugs do not affect test results. Drug excretion in oral fluid depends on the pH and salivary flora.
      2. Reassure the patient that there is no discomfort with the procedure. Patients who fear dental work may be apprehensive about having anything done in the mouth.
      3. Usually, dentures or partials do not have to be removed. Assess for an allergy to cotton.
      4. Follow standard precautions, and wear gloves (see Appendix A).
      5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
   b. Posttest Patient Care
      1. Inform the patient regarding anticipated times the test results reports are expected.
      2. Interpret test results and counsel appropriately regarding any follow-up testings or treatment intervention needed.
      3. After collecting specimen, patient may resume activities (eg, gum chewing, eating candy, foods) immediately.
      4. After lactose ingestion, diarrhea and cramps may occur.
      5. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

3. Sputum Sampling
   a. Pretest Patient Preparation
      2. Obtain a sputum collection kit and supplies.
      3. Instruct the patient about all aspects of collection, including any specific body positioning or chest physiotherapy, such as chest clapping.
      4. Inform the patient not to touch inside of the sputum container.
      5. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
   b. Posttest Patient Care
      1. Evaluate patient outcomes, and counsel appropriately about treatment and self-care for respiratory illness.
      2. Monitor the respiratory status as necessary, and intervene appropriately when indicated.
      3. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

4. Breath Sampling
   a. Pretest Patient Care
      1. Explain test purpose and type of specimen collection. Inform the patient that during the act of breathing, certain substances may occur in the exhaled breath, which are then identified by means of a breath analyzer.
      2. Dietary restrictions may apply. Check with your laboratory (eg, fasting 6 to 12 hours prior to lactose testing).
      3. Activity is limited during complex procedures. No strenuous activity prior to tests.
      4. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.
   b. Posttest Patient Care
      1. Evaluate patient outcomes and counsel appropriately about results, further testing, and possible treatment.
      2. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

1. Ultrasonic nebulizers may be used for sputum induction when the cough is not productive. If this is the case, proper cleaning and disinfection of the nebulizer must be done.
2. Do not obtain a suctioned sample without first consulting the physician.
3. Specimens that are to be transported within an institution should be placed in a biohazard zip-lock bag. Ideally, the requisition should accompany the specimen but not be sealed inside the bag (typically there is a pocket on the outside of the bag for the requisition).

BIBLIOGRAPHY


Appendix L Protocols for Evidentiary Specimen Collection in Criminal or Forensic Cases

PROCEDURES FOR SPECIMEN COLLECTION AND PRESERVATION IN CRIMINAL OR FORENSIC CASES

Clinical Alert: Procedure for Sexual Assault
Clinical Alert: Procedure for Collecting Other Evidentiary Specimens in Violent and Trauma Cases

Important evidentiary specimens are collected from the living and the dead and can include blood, tissue, hair, nails, body fluids (eg, urine, semen, saliva, vaginal fluid, gastric fluid), and evidence generated by diagnostic procedures such as x-ray, CT scans, angiograms, endoscopy, and electrocardiograms. All collection procedures are to be followed precisely, with complete documentation with appropriate signatures and titles as required. These specimens and test results often become evidence for legal situations. Collaboration with other professionals is mandatory. Health professionals may interact with assault and abuse victims in settings other than emergency rooms and ambulances (eg, critical care areas, medical/surgical departments in hospitals, home environments and school settings).

PROCEDURES FOR SPECIMEN COLLECTION AND PRESERVATION IN CRIMINAL OR FORENSIC CASES

1. Obtain evidence from the victim or subject, including victims of predatory crimes, physical and sexual assault (both female and male), homicide, and child, elder, and spousal abuse, victims of date rape drugs, poisons, and toxic drugs. Specimens may be obtained from the patient, from the deceased person, the perpetrator, the suspect, the accused or falsely accused, drug addicts, or the public in general.

2. Collect evidentiary material required by law enforcement, the medical examiner, participants in the scientific investigation of death and criminal injury, and needed for public or legal proceedings.

3. Retrieve evidentiary items during diagnostic and therapeutic procedures (eg, head shot projections, shotgun wadding or bullets) that are often recovered from the victim or suspect's clothing, or the sheets/blankets on which the victim is transported. These items should be wrapped in gauze and placed in an envelope or cup.

4. Collect victim's clothing that may contain the victim's or perpetrator's blood, body fluid and/or tissue.

5. Obtain trace evidentiary specimens, such as traces of soot as in gunshot wounds, "tattooing" from gunpowder, punctate abrasions in intermediate-range gunshot wounds, or collect debris (eg, gravel, grass, soil, twigs, glass) that place the victim at the crime scene.

6. Record and have a witnessed record (ie, by a record person) of narcotics found on the victim, dangerous drugs, and money (ie, have two people count).

7. Place items in paper bags, not plastic.

8. Document patient's condition, signs and symptoms of "club" drugs ingestion, malignant hyperthermia, and/or odor of alcohol. Club drugs are colorless, tasteless, and odorless. Combination of these drugs can lead to muscle breakdown, cardiovascular failure, and death. (See Table L.1)

Table L.1 Date Rape and Common Club Drugs

<table>
<thead>
<tr>
<th>Drug Names (Chemical and Street)</th>
<th>Route of Administration</th>
<th>Physical Examination (findings, signs, and symptoms of use in unaware victims)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MDMA (I) 3–4 methylenedioxymethamphetamine Ecstasy, Adam, XTC, X, Hug Drugs, Beans, Love Drugs, Lover's, Speed</td>
<td>Smoked, snorted, swallowed, injected</td>
<td>Tachycardia, hypertension, hyperthermia, mydriasis, diaphoresis, stimulant, entacto-genic effects, visual hallucinations</td>
</tr>
<tr>
<td>GHB (I) Gamma hydroxybutyrate Liquid ecstasy, Liquid X, Everclear, Soap, Easy Lay, Goops, Georgia home boy LSD (I)</td>
<td>Liquid shots (capfuls), ften mixed with alcohol, or juices due to bitter taste; pills, capsules, injected intramuscularly</td>
<td>Bradycardia, respiratory depression, hallucinations, amnesia</td>
</tr>
<tr>
<td>Lysergic acid diethylamide Acid, L, Blotter, Trips, Cid, Tabs, Microdots Rohypnol (I) Flunitrazepam Rophies, Roofies, Roach, Rope, Circles, Mexican valium</td>
<td>Sniffed, snorted, mixed with alcohol or water, injected with heroin</td>
<td>Tachycardia, hypertension, hyperthermia, mydriasis, lacrimation, visual hallucinations, syntheses ataxia, tremors</td>
</tr>
</tbody>
</table>

Respiratory depression, Hypotension, dis-Orientation, dizziness, Visual disturbances, anterograde amnesia
Methamphetamine (II)
Smoked, snorted, swallowed, injected intravenously
Tachycardia, hypertension, hyperthermia, mydriasis, diaphoresis, visual hallucinations, "floating out-of-body" or "near-death" Experience

Methylamphetamine
Speed, Ice, Meth, Crystal, Crank, Fire, Glass

Ketamine (III)
Swallowed, snorted, smoked, injected intra-muscularly, combined with other drugs (eg, MDMA), smoked with marijuana and tobacco
Tachycardia, hypertension, nystagmus, respiratory depression, visual hallucinations, cataleptic state

Ketamine hydrochloride
Special K, Lady Kay, Vitamin K, Jet, K, Keets, Super C, Cat
valium, K-hole

Clinical Alert
1. Never leave evidence unattended.
2. Human bites have an infection rate of 15%–20%.

Procedure for Sexual Assault

1. Obtain consent using form in standard sexual assault kit.
2. Wear gloves and use standard precautions in conscious, cooperative, nonhallucinatory victims.
3. For obtaining specimens after sexual assault, use a Sexual Assault Evidence Collection Kit, which contains the items needed to collect samples as required by the area crime laboratory for alleged sexual assault cases. Samples can be obtained with the kit from males and females, victims and suspects. Photographs of assault areas are often taken.
4. Provide information on a form enclosed in the evidentiary specimen kit and have the individual and the examining medical professional sign and date the form.
5. Be aware that one-swab techniques are generally used for moist secretions and two-swab techniques are used for dry secretions for the following tests:
   a. Clothing samples
      1. Have the individual stand on clean pieces of examination paper if he or she is conscious and able to stand.
      2. Have the individual remove clothing, placing each article into a new, clean paper bag. Then fold the examination paper, placing it also into a clean paper bag. Date, seal, and initial each bag.
   b. Vaginal swab
      1. Using the four swabs in the kit, thoroughly swab the vagina.
      2. Prepare a smear using the four swabs; allow the swabs and smear to air dry.
      3. Place the swabs in a swab box, return the smear to the cardboard slid holder, and tape it shut. Seal and complete the information requested on the envelope provided.
   c. Cervical swab
      1. Using two swabs, thoroughly swab the cervix and immediately prepare a smear.
      2. Allow the swabs and smear to thoroughly air dry, return the swabs to the swab box, and return the smear to the cardboard slide holder. Seal and complete the information requested on the envelope provided.
   d. Rectal swab
      1. Using a single swab, moisten it with sterile water, gently insert into the rectum (to a depth of about 3 cm) and rotate.
      2. Allow the swab to air dry, place it in the envelope, and seal and label it.
   e. Penile swab
      1. Using a single swab, moisten it with sterile water, and thoroughly swab the external area of the entire penis.
      2. Use at least two (2) swabs, allow the swabs to air dry, place them in the envelope, seal, and label.
   f. Pubic hairs
      1. Using comb provided, obtain 20 to 30 pubic hairs, or obtain at least 20 to 25 plucked pubic hairs from the victim. Patients are given the option of plucking their own hairs.
      2. Be aware that these hairs are compared with suspect's hairs.
   g. Blood and/or urine samples for DNA testing
      1. Perform a venipuncture, and collect at least 5 mL of blood in the lavender-stoppered (EDTA) blood tube provided in the kit. Label the tube with the individual's name and the date.
      2. From this tube, withdraw 1 mL of blood, and fill each of the four printed circles on the DNA Stain Card.
      3. Allow the card to air dry, write the individual's name on the card, place it in the envelope, seal it, and complete the information requested. The blood tube should be placed in the Styrofoam blood tube holder.
      4. Seal the holder with the evidence tape, supply the information requested, and place the unit in the zip-lock bag provided. DNA can be identified in skin cells, dandruff, and perspiration stains on clothing. Urine specimen collection can be random witnessed or obtained by catheterization.
   h. Other samples
      1. Be aware that only certain commercial analyzers are approved for evidentiary breath testing (eg, breath alcohol).
      2. Remember that saliva samples for evidentiary use are usually not collected, as other specimens for DNA evidence are more important.
6. Before internal specimen examination, use Wood's lamp (long-wave ultraviolet light) to scan genital area. Record findings. Apply toluidine blue dye to external perineal area with cotton-tipped applicators. Area is then gently wiped dry and lubricating gel applied.
7. After all samples have been collected, place all specimens (except blood tube) back into the kit. The blood tube should be refrigerated and the kit kept at room temperature until picked up by the police. Items that are wet (eg, bloody clothing) are first dried and then placed in thick paper bags (not plastic) and then labeled as a biohazard. Each specimen is to be labeled and packaged separately.
8. In cases of rape, obtain samples for identification of sexually transmitted diseases (STDs).
   a. Record and report findings, the types of specimens saved, and report to proper authorities. Date rape or acquaintance rape facilitated by means of “club” drugs will exhibit varied signs and symptoms (eg, memory loss, confusion, vertigo, slurred speech, drowsiness, etc.).
   b. Assess and record evidence of trauma. Colposcopic examination is often done.
   c. Record use of weapons, forced drug use, frequency of sexual acts, and penetration of vagina or rectum by penis or object.

Clinical Alert

1. Specimens in suspected rape cases are tested for pregnancy, HIV, hepatitis B, syphilis, trichomoniasis, gonorrhea, and chlamydia.
2. Specimens of newborn cord blood and meconium may be examined for evidence of the mother's drug use during pregnancy. Nail clippings and nail scrapings are also examined.
3. Check level of consciousness and record vital signs in semiconscious, unconscious, and nonresponding persons.
4. Tachycardia, bradycardia, hypertension, and hyperthermia are signs of “club” drug administration. Treat immediately.
5. Examine, measure, photograph, and record evidence of bite marks, burns, abrasions, ecchymosis, and trauma.
7. Document whether trauma is present or absent.
8. Specimen collection after death is explained in Chapter 16.

Procedure for Collecting Other Evidentiary Specimens in Violent and Trauma Cases

1. Be certain not to cut through bullet holes or marks from other weapons that may have been used in a violent attack.
2. Do not give clothing or other items belonging to the victim back to the possible offender. The possible offender may have accompanied the victim.
3. Seal clothing in paper bag, as bacteria could destroy DNA evidence if stored in plastic bags.
4. Examine evidence carefully, such as looking for trauma, at spatter patterns on clothing, in hair fibers, presence of gunshot powder and identifying potential causes of wounds (gunshot, stabbing). Take photographs and be sure to include identifying information (eg, victim's name, date/time, case number and other relevant information) on the photograph, slide mount, or negative.
5. For example, in situations where children have been exposed to home methamphetamine labs, assess for possible methamphetamine ingestion, collect evidence, and prepare children for going into protective custody. To gain children's cooperation, develop a positive rapport with them and alleviate their fears.

Clinical Implications

1. Presence of sperm or semen—a negative finding in suspected rape cases is <10 U/L (or <167 nkat/L) of acid phosphatase, whereas a level that is >50 U/L (or >833 nkat/L) indicates a positive result for the presence of semen. Although the acid phosphatase levels of prostatic fluid can still be elevated in about 10% of women 72 hours after assault, low or no appreciable levels do not exclude recent penile penetration (eg, ejaculation may not have occurred).
2. DNA samples from blood, semen deposits, stains in clothing, etc. are taken and compared with venous blood sample from victim.
3. Positive or negative findings for pregnancy, STDs, HIV, hepatitis B, syphilis, chlamydia, gonorrhea, and trichomoniasis
4. Diffuse uptake or no uptake from toluidine blue dye, a negative finding for presence of microlacerations
5. No abnormal findings on colposcopy of trauma or genital injuries
6. No positive findings of date rape drugs, poisons, or other toxic substances
7. Clinical implications of these findings can be categorized as some concern, serious concern, and grave concern (see Table L.2.)

<table>
<thead>
<tr>
<th>Level of Concern</th>
<th>Clinical Indicators</th>
</tr>
</thead>
<tbody>
<tr>
<td>Some</td>
<td>Anal dilation (&gt; 2 cm) or fissures</td>
</tr>
<tr>
<td></td>
<td>Perianal bruising</td>
</tr>
<tr>
<td></td>
<td>Friable posterior fourchette</td>
</tr>
<tr>
<td></td>
<td>Presence of labial friability (in girls beyond the diaper-wearing age)</td>
</tr>
<tr>
<td>Serious</td>
<td>Anal scarring or tags (beyond the midline)</td>
</tr>
<tr>
<td></td>
<td>Disruptions of the hymenal border</td>
</tr>
<tr>
<td></td>
<td>Genital injury</td>
</tr>
<tr>
<td></td>
<td>Posterior scars</td>
</tr>
<tr>
<td>Grave</td>
<td>Pregnancy</td>
</tr>
<tr>
<td></td>
<td>Presence of semen, sperm, or acid phosphatase</td>
</tr>
<tr>
<td></td>
<td>Anogenital injury</td>
</tr>
<tr>
<td></td>
<td>Non-perinatally acquired STD</td>
</tr>
</tbody>
</table>

Adapted from Olshaker JS, Jackson MC, Smock WS: Forensic Emergency Medicine, p. 156. Philadelphia, Lippincott Williams & Wilkins, 2001
Interfering Factors

1. Samples should ideally be collected immediately after an alleged assault, because 66% of women examined within 6 hours of the incident do not show motile sperm. Consequently, estimating the time of assault may be hampered. Bathing can wash away evidence.
2. See examples of form used for victim information in Appendix H.

Interventions

Pretest Patient Care

1. To help reduce victim's acute psychological trauma, be especially sensitive and nonjudgmental when caring for victims, who often are filled with guilt, shame, and self-blame.
2. Explain the purpose and procedure of procuring samples as they relate to the alleged incident. Check to see whether informed consent must be signed before obtaining specimens. Whenever possible, enlist the aid of rape crisis professionals. Sedatives may be given to childhood victims of sexual assault and abuse.
3. Become familiar with effects of “street” drugs that may have been given to the victim of sexual assault.
4. Ensure that all forms are completed accurately and signed by the victim. (See examples of forms used in Appendix H.)
5. Use standard precautions to collect specimens.
6. Prepare to institute a witnessed chain of custody procedure, and follow these policies throughout when obtaining, handling, and preserving specimens.
7. Obtain names, addresses, and telephone numbers of witnesses or persons accompanying the victim.
8. Use forms (examples of typical forms in Appendix H) for female and male evidence collection.
9. Follow guidelines in Chapter 1 for safe, effective, informed pretest care.

Posttest Patient Care

1. Follow chain of custody (chain of evidence) when turning over evidentiary specimens to law enforcement officials.
2. Record all actions regarding specimen retrieval, including photographs and expected and unexpected samples (eg, bullets, drugs). Victim's name, identifying number, case and/or hospital number, and other relevant information should be written on the photograph, or slide mount or negative envelope.
3. Evaluate patient outcomes, monitor, and counsel the patient appropriately about further tests (pregnancy, syphilis, HIV at 6, 12, and 24 weeks) or treatment or follow-up for post-traumatic stress disorder. As appropriate: antibiotics, tetanus for human bites, hydration, coolants for hyperthermia, muscle relaxants, placing patient in a quiet area, activated charcoal to remove drugs from stomach, sedatives and anticonvulsants to control delirium. (See Table L.3, transmission risk of STDs)

Table L.3 Risk of STDs after Sexual Assault

<table>
<thead>
<tr>
<th>Sexually Transmitted Disease</th>
<th>Transmission Risk (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trichomoniasis</td>
<td>30–85 or 0.30–0.85</td>
</tr>
<tr>
<td>Chlamydia</td>
<td>30–70 or 0.30–0.70</td>
</tr>
<tr>
<td>Gonorrhea</td>
<td>20–90 or 0.20–0.90</td>
</tr>
<tr>
<td>Syphilis</td>
<td>30–60 or 0.30–0.60</td>
</tr>
<tr>
<td>HPV</td>
<td>60–70 or 0.60–0.70</td>
</tr>
<tr>
<td>HSV-2</td>
<td>&lt;5 or &lt;0.05</td>
</tr>
<tr>
<td>Hepatitis B</td>
<td>&lt;5 or &lt;0.05</td>
</tr>
<tr>
<td>HIV</td>
<td>&lt;5 or &lt;0.05</td>
</tr>
</tbody>
</table>

Adapted from Olshaker JS, Jackson MC, Smock WS: Forensic Emergency Medicine, p. 135. Philadelphia, Lippincott Williams & Wilkins, 2001

4. Follow guidelines in Chapter 1 for safe, effective, informed posttest care.

Clinical Alert

It is best if rape crisis professionals can evaluate the victim and collect specimens. Prior to the start of the Sexual Assault Nurse Examiner (SANE) programs, victims of sexual assault were taken to hospital ERs and may not have been triaged into the health system as a high priority for care. Assault victims need immediate care, but not necessarily technical care. SANE nurses are trained in forensic investigation, whereas most ER staff can handle medical screening appropriately, but not the forensic aspects of the situation.

How the SANE program works when a woman is assaulted:

1. Police officer or rape crisis line counselor notifies SANE on-call nurse.
2. Nurse meets victim in a designated private room away from the ER. Seriously injured persons or possible victims of date rape drugs would be treated first.
3. Forensic examination is performed, medical treatment provided, nonjudgmental support offered, and referrals for ongoing care completed.
4. SANE nurse provides evidence to police department and may later testify in court.
5. Specially trained professionals may also aid in identifying missing children and their abductors.
BIBLIOGRAPHY


International Association of Forensic Nurses. (Online.) Available at:  http://www.forensicnurse.org/

Olshaker JS, Jackson MC, Smock WS: Forensic Emergency Medicine, Philadelphia, Lippincott Williams & Wilkins, 2001


Williams RH: Club drugs—what's all the rave?. Clin Lab News, 10–13, December 2001