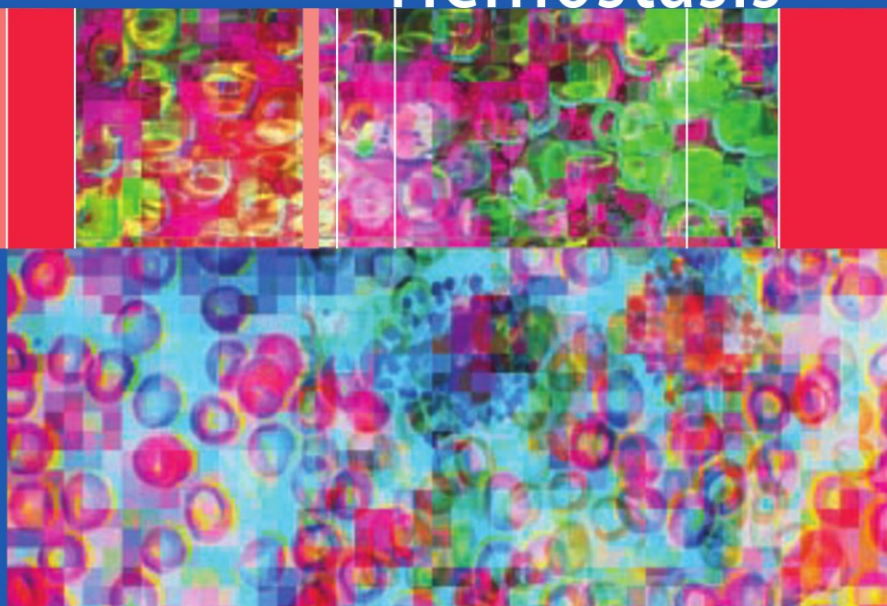


Sterling T. Bennett
Christopher M. Lehman
George M. Rodgers
Editors

Laboratory Hemostasis



A Practical Guide for Pathologists

 Springer

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Sterling T. Bennett, M.D., M.S.

*Medical Director, Intermountain Healthcare Central
Laboratory, and Volunteer Adjunct Associate Professor
of Pathology, University of Utah Health Sciences Center
Salt Lake City, Utah*

Christopher M. Lehman, M.D.

*Associate Professor of Pathology
University of Utah Health Sciences Center
and
Assistant Director of Hospital Clinical Laboratories
ARUP Laboratories
Salt Lake City, Utah*

George M. Rodgers, M.D., Ph.D.

*Professor of Medicine and Pathology
Division of Hematology
University of Utah Health Sciences Center
and
Medical Director, Coagulation Laboratory
ARUP Laboratories
Salt Lake City, Utah*

Editors

With Contributions by Catherine Thompson, B.S.,
MT (ASCP), and Robert C. Blaylock, M.D.



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Sterling T. Bennett, M.D., M.S.
Medical Director
Intermountain Healthcare Central
Laboratory
Volunteer Adjunct Associate Professor
of Pathology
University of Utah Health
Sciences Center
Salt Lake City, UT 84143
USA

Christopher M. Lehman, M.D.
Associate Professor of Pathology
University of Utah Health
Sciences Center
Salt Lake City, UT 84132
and
Assistant Director of Hospital
Clinical Laboratories
ARUP Laboratories
Salt Lake City, UT 84132
USA

George M. Rodgers, M.D., Ph.D.
Professor of Medicine and Pathology
Division of Hematology
University of Utah Health
Sciences Center
Salt Lake City, UT 84132
and
Medical Director, Coagulation Laboratory
ARUP Laboratories
Salt Lake City, UT 84132
USA

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*To Elizabeth Bennett, Julie Monahan and
other family and friends for their unfailing support,
and to the numerous residents and fellows
and our faculty colleagues who have
stimulated our interest in coagulation
and inspired this book*

Preface

In the course of our practice as coagulation laboratory directors, we have acquired personal libraries of hemostasis and thrombosis texts that serve as excellent references for the basic science and clinical aspects of hemostasis. What is missing, though, is a single reference text that provides practical information and guidance on the topics relevant to directing a coagulation laboratory. We are sure we are not alone in wishing for such a reference. After talking about it for several years, we have now attempted to fill the void with a handbook containing the information we provide to the residents and fellows that we are helping train to become laboratory directors.

In writing this book we have tried to cite the consensus recommendations of authoritative bodies whenever possible, and otherwise, have cited reviews, chapters, and research articles of experts in their respective fields. These references should act as a starting point for a more detailed study of the various aspects of coagulation testing.

It is our intent that pathologists, clinical laboratory scientists, and other physicians serving as laboratory directors will find this book helpful to understand and carry out their responsibilities. Another intent is that residents and fellows will find this book to be a useful tool for learning the basics of coagulation testing and for studying for board examinations. We also hope that coagulation laboratory supervisors, technologists, and technicians find this to be a helpful reference for the day-to-day operation of the laboratory. Finally, clinicians may find aspects of this book helpful in understanding the role of the coagulation laboratory in patient evaluation and monitoring.

STERLING T. BENNETT
CHRISTOPHER M. LEHMAN
GEORGE M. RODGERS

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1

Role and Responsibilities of the Laboratory Director

Sterling T. Bennett

Role and Responsibilities of the Laboratory Director

Clinical laboratories provide essential services and information for the practice of medicine. The majority of electronic medical data consists of laboratory results and laboratory results are vital information for many clinical decisions. Accordingly, laboratories have the potential of making great contributions to patient safety and medical outcomes when their services are timely and results are accurate.

Serving as a laboratory director is an interesting, challenging, and rewarding job, and one that carries great responsibility. The director is accountable for all aspects of the laboratory service and is the key bridge between laboratory operations and clinical practice. As such, the laboratory director needs to have a broad knowledge of clinical medicine, pathology disciplines, basic medical sciences, clinical laboratory sciences, laboratory operations, and quality management systems. Knowledge of and skills in informatics, data analysis, and business management are also important assets. The breadth of knowledge and skills required of a laboratory director is one aspect that makes the job so attractive.

In the midst of all this knowledge, it is vital for the success of the laboratory that the director clearly understands his or her role and responsibilities. Definitions of the laboratory director's responsibilities have been published by the College of American Pathologists, American Association of Blood Banks, International Organization for Standards (ISO), Clinical Pathology Accreditation (UK) Ltd., National Pathology Accreditation Advisory Council, European Society of Pathology, and Canadian Association of Pathologists, to

name a few¹⁻⁸. In addition, the United States' Code of Federal Regulations defines the laboratory director's responsibilities under the Clinical Laboratory Improvement Amendments of 1988 (CLIA-88)⁹. Table 1.1 provides a summary listing of the laboratory director's responsibilities.

For any given laboratory, the laboratory director's responsibilities are defined in large measure by prevailing regulations and the standards of accreditation bodies. Responsibilities may be further defined by the organizational structure of the institution and contractual arrangements between the institution and laboratory director; however, organizational structure and contracts do not absolve the laboratory director of responsibilities defined by regulations and accreditation standards.

Each set of regulations or accreditation standards is unique, with its own emphasis on the director's responsibilities. Some define detailed responsibilities while others provide few specifics. Tables 1.2-1.8 illustrate these differences by listing the duties specifically attributed to the laboratory director in four sets of regulations or standards.

These tables are not intended to be a comprehensive cross-reference between various standards and regulations, but to illustrate the commonalities and differences in the level of detail that different organizations give to the specific responsibilities of the director. Many other regulations and standards exist beyond those listed. These tables include only those items that are specifically identified as responsibilities of the "laboratory director" by the regulations of the Clinical Laboratory Improvement Amendments of 1988 (US)⁹ and the College of American Pathologists¹, the "person providing supervision and direction" by the National Pathology Accreditation Advisory Council (Australia)⁷, or "laboratory management" by Clinical Pathology Accreditation (UK) Ltd.⁴. The one point on which all agree is that the laboratory director is responsible for assuring the laboratory complies with all provisions of the regulations or standards. An abstraction from the various definitions suggests the laboratory director's responsibilities may be broadly categorized into general responsibilities, personnel, facilities and safety, test procedures, quality management, consultation and education, communication, and operational management. This chapter will discuss specific responsibilities within each of these categories.

Table 1.1. Laboratory Director's Responsibilities

| General Responsibilities | Personnel | Facilities & Safety | Test Procedures | Quality Management | Consultation & Education | Communication | Operational Management |
|------------------------------------|---|---|---|--|--|---|--|
| Overall operation & administration | Needs assessment Job descriptions Hiring standards Staff selection | Biosafety Chemical hygiene Ergonomics | Test menu selection Method selection Method validation Quality control program | Quality program Quality standards Quality assessment | Accessible to medical staff & laboratory staff Test ordering Result interpretation | Clinicians Patients Administration Lab staff | Strategic planning Capital budgets Operational budgets |
| Definition of others' duties | Assignment of duties | | Harmonization or standardization of methods | Proficiency testing | Therapeutic recommendations | Risk management Regulators | Research and development |
| Delegation of authority | Training standards Staff training | | Requisitions/ ordering processes | Inter-method comparisons | Troubleshooting Utilization patterns | Accreditation or certification bodies | |
| Retention of responsibility | Competency standards Competency assessment | | Specimen acceptance & rejection criteria | Audits Surveys | Test menu changes Staff continuing education | | Contracting Marketing |
| Licensure & accreditation | Performance standards | | Specimen collection & handling | Inspections Problem reports | Clinician education Institutional education | | Billing Organizational goals |
| Medical staff privileges | Performance reviews Remedial actions | | Transportation Result reporting Standard operating procedures | Remedial actions Quality improvement | Users' needs assessment | | Metrics |
| | | | Reference laboratory selection & monitoring | | | | |

Table 1.2. General Responsibilities of the Laboratory Director; Examples of Regulations and Standards^a

| Category | CLIA | CAP | NPAAC | CPA |
|------------------------------------|--|--|---|---|
| Overall Operation & Administration | Take responsibility for overall operation and administration of lab, including employment of personnel competent to perform test procedures, record and report test results promptly, accurately and proficiently. | Take responsibility for overall operation and administration of lab. Provide supervision. Assume professional, scientific, consultative, organizational, administrative, and educational responsibilities. | Take responsibility for professional, scientific, consultative, organizational, administrative and educational requirements of lab. | N/A |
| Duties & Delegation | If desired, delegate duties of technical supervisor, technical consultant, clinical consultant, general supervisor, and testing personnel to qualified personnel. Specify, in writing, responsibilities and | If desired, delegate administrative, medical and technical responsibilities to qualified individuals. Retain consultative services, as needed, for each laboratory discipline. | If desired, delegate elements of responsibilities to qualified and experienced pathologists or scientists. | Define organization and management of lab and its place in parent organization. Specify roles, responsibilities, authority and interrelationships of all personnel. |

(Continued)

Table 1.2. (Continued)

| | | | | |
|-----------------------------|---|--|-----------------------------------|---|
| | duties of each consultant, supervisor, and person engaged in any phase of testing. | | | |
| Retention of Responsibility | If responsibilities are reapportioned, remain responsible for ensuring all duties are properly performed. | If duties are delegated, remain responsible to assure quality patient services are provided. | N/A | N/A |
| Licensure & Accreditation | Assure compliance with applicable regulations. | Implement and maintain CAP Standards. | Implement and maintain standards. | Commit to continuing compliance with accreditation standards. |
| Medical Staff Privileges | N/A | As applicable, serve as an active member of medical staff for facilities served. | N/A | N/A |

^a Abbreviations: CLIA – Clinical Laboratory Improvement Amendments of 1988 (US), CAP – College of American Pathologists, NPAAC – National Pathology Accreditation Advisory Council (Australia), CPA – Clinical Pathology Accreditation (UK).

Table 1.3. Personnel: Responsibilities of the Laboratory Director; Examples of Regulations and Standards^a

| Category | CLIA | CAP | NPAAC | CPA |
|-------------------|---|---|--------------|--|
| Supervision | Ensure on-site supervision of high complexity testing. Identify supervision required for specimen processing, test performance or result reporting. Identify supervisory or director review required prior to reporting patient test results. | N/A | N/A | N/A |
| Adequate Staffing | Employ sufficient personnel with education, training, and experience to provide consultation, supervise and perform tests, and report test results. | Ensure sufficient qualified personnel with documented training and experience to meet needs of lab. | N/A | Ensure appropriate staff, with education and training to meet service demands and regulations. Ensure registration of staff in accordance with applicable regulations. Ensure staffing includes roles in quality management, training, education, health and safety. |

(Continued)

Table 1.3. (Continued)

| | | | | |
|--------------------------------------|---|-----|-----|--|
| Personnel Management & Authorization | Identify which examinations and procedures each individual is authorized to perform. | N/A | N/A | Ensure personnel procedures include recruitment, selection, orientation, job descriptions, contracts, records, reviews, meetings, communication, training, education, grievance and disciplinary action. Ensure confidentiality of personnel records. |
| Training & Competency Assessment | Ensure all personnel have appropriate education and experience, appropriate training for type and complexity of services, and demonstrated they can perform testing reliably and report accurate results. | N/A | N/A | Appoint a training officer. Ensure all staff are oriented to lab, parent organization, conditions of employment, patient confidentiality, data protection, health, safety, occupational health services, job description, organizational chart, salaries and wages, and staff facilities. |
| Monitoring Competency | Ensure policies and procedures for monitoring individuals to assure competency and to assure identification of needs for remedial training or continuing education. | N/A | N/A | N/A |

^a Abbreviations: CLIA – Clinical Laboratory Improvement Amendments of 1988 (US), CAP – College of American Pathologists, NPAAC – National Pathology Accreditation Advisory Council (Australia), CPA – Clinical Pathology Accreditation (UK).

Table 1.4. Facilities and Safety: Responsibilities of the Laboratory Director; Examples of Regulations and Standards^a

| Category | CLIA | CAP | NPAAC | CPA |
|---------------------|---|---|-------|---|
| Facilities & Safety | Ensure physical plant and environmental conditions are appropriate and provide a safe environment in which employees are protected from physical, chemical, and biological hazards. | Implement a safe lab environment in compliance with good practice and applicable regulations. | N/A | <p>Ensure a safe working environment.</p> <p>Provide personal protective equipment.</p> <p>Delegate day to day management of health and safety to health and safety officer.</p> <p>Provide model rules for staff and visitors.</p> <p>Provide for regular reporting of communicable diseases to government agencies.</p> <p>Define and implement procedures for actions in the event of fire, major spill of dangerous chemicals or clinical material, or inoculation accident; reporting and monitoring of accidents and incidents; risk assessments; disinfection processes; decontamination of equipment; chemical handling; and storage and disposal of waste.</p> |

^a Abbreviations: CLIA – Clinical Laboratory Improvement Amendments of 1988 (US), CAP – College of American Pathologists, NPAAC – National Pathology Accreditation Advisory Council (Australia), CPA – Clinical Pathology Accreditation (UK).

Table 1.5. Test Procedures: Responsibilities of the Laboratory Director; Examples of Regulations and Standards^a

| Category | CLIA | CAP | NPAAC | CPA |
|--|--|--|-------|---|
| Quality in Preanalytic, Analytic & Postanalytic Phases | Ensure testing systems provide quality services for all aspects of testing. | N/A | N/A | Define and implement procedures for specimen collection, handling, reception, transportation, and rejection. |
| Test Methods | Ensure test methodologies can provide quality required for patient care. | N/A | N/A | Establish procedures for procurement and management of equipment. |
| Verification of Performance Characteristics | Ensure verification procedures determine accuracy, precision, and other pertinent performance characteristics of test methods. | N/A | N/A | N/A |
| Quality Control | Ensure quality control program is established and maintained to assure quality and identify failures in quality as they occur. | Define overall quality control program, including goals, policies, procedures, delegation of responsibilities, and regular review. Define, implement, and monitor standards of performance in quality control. | N/A | Ensure availability of quality control material. Establish procedures for management of quality control material. |

(Continued)

Table 1.5. (Continued)

| Category | CLIA | CAP | NPAAC | CPA |
|-------------------------------|--|---|------------|---|
| Analytical Performance Levels | <p>Ensure personnel perform test methods as required for accurate and reliable results.</p> <p>Ensure establishment and maintenance of acceptable levels of analytical performance for each test system.</p> <p>Ensure patient test results are reported only when the system is functioning properly.</p> | <p>Define program that monitors and demonstrates the proper calibration, function, and preventive maintenance of instruments and equipment.</p> | <p>N/A</p> | <p>Ensure availability of required reagents, calibration and quality control material.</p> <p>Establish procedures for management of reagents, calibration and quality control material, including selection, purchasing, ordering, supplier assessment, receipt, verification of identity and condition, inventory management, risk assessment, assignment of handling precautions, and safe disposal.</p> |
| Remedial Actions | <p>Ensure remedial actions whenever significant deviations from established performance characteristics are identified.</p> | N/A | N/A | N/A |

(Continued)

Table 1.5. (Continued)

| | | | | |
|------------------------|---|---|-----|---|
| Reports | Ensure reports of test results include pertinent information required for interpretation. | N/A | N/A | Establish procedures for reporting results, giving reports by telephone, and issuing amended reports. |
| Data Management | N/A | N/A | N/A | Ensure availability of data and information to meet user needs. Establish procedures for management of data and information, including security, access, confidentiality, data protection, backup systems, storage, archive, retrieval and secure disposal. Ensure compliance with regulations for data protection. |
| Procedure Manual | Ensure approved procedure manual is available to all personnel. | N/A | N/A | N/A |
| Anatomic Pathology | N/A | Perform anatomic pathology procedures as appropriate. | N/A | N/A |
| Reference Laboratories | N/A | Select and monitor reference labs. | N/A | Define and implement procedures for referral to other labs. |

^a Abbreviations: CLIA – Clinical Laboratory Improvement Amendments of 1988 (US), CAP – College of American Pathologists, NPAAC – National Pathology Accreditation Advisory Council (Australia), CPA – Clinical Pathology Accreditation (UK).

Table 1.6. Quality Management: Responsibilities of the Laboratory Director; Examples of Regulations and Standards^a

| Category | CLIA | CAP | NPAAC | CPA |
|---------------------|--|---|-------|-----|
| Proficiency Testing | <p>Ensure enrollment in approved proficiency testing program.</p> <p>Ensure proficiency testing samples are tested in same manner as patients' specimens.</p> <p>Ensure results are returned within timeframes established by proficiency testing program.</p> <p>Ensure proficiency testing reports are reviewed to evaluate performance and identify problems that require corrective action.</p> <p>Ensure approved corrective action plan is followed when any proficiency testing result is unacceptable or unsatisfactory.</p> | <p>Monitor results of proficiency testing and participate in documentation of corrective actions.</p> <p>Develop a mechanism for determining accuracy and reliability of procedures for which a formal external audit program is not available.</p> | N/A | N/A |

(Continued)

Table 1.6. (Continued)

| | | | | |
|-----------------|--|---|-----|--|
| Quality Program | Ensure quality assurance program is established and maintained to assure quality of lab services and identify failures in quality as they occur. | Systematically monitor and evaluate quality and appropriateness of the laboratory's contribution to patient care. Ensure a process to address any systematic problem or any problems of individual competence that have been identified. | N/A | Establish quality management system. Establish quality policy that includes commitments to full implementation of quality management system. Ensure quality policy is signed by a person with authority, understood and implemented throughout the laboratory, reviewed annually, and consistent with other policies in the organization. Set quality objectives and undertake quality planning. Prepare quality manual. Appoint quality manager. Establish procedures for document control, control of process and quality records, and control of clinical material. Conduct management reviews of quality program. |
|-----------------|--|---|-----|--|

(Continued)

Table 1.6. (Continued)

| Category | CLIA | CAP | NPAAC | CPA |
|---------------------|------|--|-------|---|
| Quality Improvement | N/A | Assume responsibility for implementation of quality improvement plan. Define, implement, and monitor standards of performance in quality improvement. Ensure department participates fully in institutional quality improvement plans that deal with relevant areas and outcomes of patient care. Participate as members of various quality improvement committees of the institution. | N/A | Establish procedures for assessment of user satisfaction and complaints, internal audit of quality management system, internal audit of examination processes, external quality assessment, reports from external assessment bodies, and quality improvement. Assess clinical relevance of lab investigations and reliability of interpretive reports in conjunction with users. Participate in evaluation of clinical effectiveness, audit and risk management activities of the parent organization or external bodies. |

^a Abbreviations: CLIA – Clinical Laboratory Improvement Amendments of 1988 (US), CAP – College of American Pathologists, NPAAC – National Pathology Accreditation Advisory Council (Australia), CPA – Clinical Pathology Accreditation (UK).

Table 1.7. Communication, Consultation & Education: Responsibilities of the Laboratory Director; Examples of Regulations and Standards^a

| Category | CLIA | CAP | NPAAC | CPA |
|--|--|---|---|--|
| Interaction with Others | N/A | Relate and function effectively with accrediting regulatory agencies, administrative officials, medical community, medical device industry, and patient population. | N/A | N/A |
| Accessible to Laboratory | Be accessible to lab to provide onsite, telephone or electronic consultation as needed. | N/A | N/A | N/A |
| Medical Significance, Interpretation & Correlation of Data | Ensure consultation is available to lab's clients on matters relating to quality of test results reported and interpretation concerning specific patient conditions. | Provide consultations about medical significance of clinical lab data. Interpret, correlate, and communicate lab data to clinical requestors. | Provide advice on clinical significance of results, interpretation and correlation of lab data, suitability of tests and procedures in various clinical situations, and appropriate further tests and procedures. | Ensure that advice on examinations and interpretation of results is available to meet the needs and requirements of users. |

(Continued)

Table 1.7. (Continued)

| Category | CLIA | CAP | NPAAC | CPA |
|-----------------|-------------|---|--------------|------------|
| Education | N/A | Provide educational programs for medical and lab staff, and participate in educational programs of the institution. | N/A | N/A |

^a Abbreviations: CLIA – Clinical Laboratory Improvement Amendments of 1988 (US), CAP – College of American Pathologists, NPAAC – National Pathology Accreditation Advisory Council (Australia), CPA – Clinical Pathology Accreditation (UK).

Table 1.8. Operational Management Responsibilities of the Laboratory Director: Examples of Regulations and Standards^a

| Category | CLIA | CAP | NPAAC | CPA |
|-----------------------------|-------------|---|--------------|--|
| Strategic Planning | N/A | Perform planning for setting goals and developing and allocating resources appropriate to medical environment. | N/A | N/A |
| Administration & Management | N/A | Provide effective and efficient administration, including budget planning and control with responsible financial management. Define, implement, and monitor standards of performance in cost-effectiveness of lab services. | N/A | Have regular meetings, with records of agreed action points, and ensure actions are discharged within an appropriate and agreed timescale. Perform management reviews. Ensure availability of necessary resources. |
| Research & Development | N/A | Plan and direct research and development appropriate to the facility. | N/A | N/A |
| Users' Needs | N/A | N/A | N/A | Determine users' needs and requirements with users. Specify users' needs and requirements as objectives for the organization and management of the lab. |

^a Abbreviations: CLIA – Clinical Laboratory Improvement Amendments of 1988 (US), CAP – College of American Pathologists, NPAAC – National Pathology Accreditation Advisory Council (Australia), CPA – Clinical Pathology Accreditation (UK).

General Responsibilities

The laboratory director is responsible for the overall operation and administration of the laboratory. As the sign on the late USA President Harry Truman's desk read: "The buck stops here." Although it takes many people in a variety of roles to make a laboratory run well, the ultimate accountability for success or failure rests with the laboratory director. It is imperative that the laboratory director assure that he or she has adequate authority to discharge this accountability. When the laboratory director is a private contractor rather than an employee of the organization that owns the laboratory, it is common that the laboratory director's span of authority is ambiguous. Even when the laboratory director is an employee of the organization, others in management may see the director as a figurehead with limited authority in day-to-day operations. But in the event of regulatory infractions or failure to meet accreditation standards, the laboratory director will be held accountable, so it is important for the director to establish, both contractually and functionally, the authority necessary to truly direct the overall operation and administration of the laboratory.

With the complexity of contemporary clinical laboratories, it is not practical (or even possible) for the laboratory director to personally perform every facet of his or her responsibilities. The laboratory director may not even have the expertise to fulfill some responsibilities. Fortunately, the laboratory director is allowed to delegate responsibilities to other qualified individuals. To maintain order and clarity—and, in some cases, to meet regulatory or accreditation requirements—delegation should be specific and in writing, and the laboratory director should be certain that designees understand the director's expectations. The director also needs to assure that the laboratory is organized in a manner that allows designees to fulfill their responsibilities.

The laboratory director retains accountability even for delegated responsibilities. In the event of an untoward event, the director cannot "pass the buck" to a designee, so the director needs to implement a method for assuring that designees are performing their delegated functions and for detecting unsatisfactory performance.

The laboratory director is accountable for licensure and accreditation. It is important for the director to be intimately familiar with applicable regulations and accreditation standards. The director's involvement is also required for licensure or accreditation applications, laboratory inspections, and correspondence with government agencies and accreditation bodies.

Time devoted to studying regulations, reading standards documents, consulting other laboratory directors, and conducting self-inspections or external inspections is time well spent. Time discussing clinical needs and laboratory issues with other members of the medical staff or physician clients is also time well spent. Understanding the role of the laboratory in clinical care and understanding the director's responsibilities in the laboratory will enable the director to help the laboratory succeed and will help the director to avoid the pitfalls that will be discussed later in this chapter.

Personnel

Personnel constitute the single most important asset of any laboratory. No matter how high the level of instrumentation, automation, computerization, or other technology, a laboratory cannot function well without competent and dedicated people. From the highs of exceptional service to the lows of errors that jeopardize patients, the fruits of the laboratory service are attributable to its people. Accordingly, the laboratory director has many responsibilities related to human resources.

The director must assure that the laboratory has an appropriate number of trained and competent staff with adequate supervision to meet the demands of the laboratory service, regulations, and accreditation standards. The director needs to assure that job descriptions accurately reflect duties to be performed, that staff are selected in an equitable process from a pool of qualified applicants, and that each member of the laboratory staff is provided with a clear description of expectations and responsibilities.

The director is responsible for the implementation of a training program that includes training standards, acceptable methods of instruction, and a system of documentation. The program must be operated in a manner that assures all staff members are adequately trained before working in the laboratory. The director is accountable for ongoing evaluation and improvement of the training program.

Closely related to training is competency, and the director is responsible for implementation of a competency assurance program, including competency standards, methods of assessment, and documentation systems. The program must assure that all personnel are competent to perform their duties and that any deficiencies in competency are detected and corrected through additional training, adjustments in supervision, reassignment of duties, or other means. The director is

accountable for ongoing evaluation and improvement of the competency assurance program.

The director is also accountable for periodic reviews of staff to evaluate additional aspects of performance such as attendance, punctuality, interpersonal skills, customer service, and so forth. A minimum frequency for reviews may be set by the organization or by regulations, but reviews should be held as often as necessary to develop attributes of staff members in accordance with the laboratory's needs and objectives.

Facilities and Safety

The director is accountable for assuring that laboratory staff, visitors, and patients have a safe and healthy environment and that the facilities are appropriate for the services provided by the laboratory. The safety program needs to encompass biological safety, chemical hygiene, and ergonomics. The director assures that safety policies and procedures are consistent with regulations and good laboratory practices and personnel are well trained in safety.

The director assures that all necessary personal protective equipment is available to and used appropriately by everyone in the laboratory. The director assures that fire extinguishers, eyewash stations, emergency showers, chemical spill kits, biohazard spill kits, and other equipment are in good working order and personnel are adequately trained in their use.

The director is accountable for the proper decontamination of equipment and work areas, the proper handling, storage, and disposal of hazardous chemicals, and the proper handling and disposal of contaminated waste.

The director assures that accidents and violations of safety policies are promptly evaluated and appropriate corrective measures are instituted, including reporting to government agencies when required. The director assures the ongoing evaluation and improvement of the laboratory's safety program.

Test Procedures

The laboratory director is accountable for all aspects of testing, including the preanalytical, analytical, and post-analytical phases. Selection of the test menu is one of the key functions of the laboratory

director. In this regard, important guidance may be derived from discussions with the medical staff or physician clients, consideration of the needs of other services in the institution, proximity of other laboratory services, requisite expertise, and evaluation of cost impact on the laboratory and other services.

The laboratory director is responsible for test methods used in the laboratory. The director assures the selection of suitable analyzers, reagents, supplies, calibrators, and control materials, so that test methods have performance characteristics that meet the needs of laboratory users¹⁰. The director assures that test methods are validated against established acceptance criteria before being placed into service for testing patients' specimens. The director assures that when two or more methods are used for the same analysis that results are equivalent.

The director is responsible for the quality control program, including types of controls, frequency of quality control testing, control limits, interpretation of control results, and procedures for out-of-range control values. The director assures that all assays are in control when patients' results are reported.

The director assures that only suitable specimens are analyzed. The director approves the types of specimens and anticoagulants that are acceptable for the methods employed and defines rejection criteria for unacceptable specimens. Rejection criteria include not only improper specimen types but also ordering, labeling, and handling issues that render a specimen unsuitable for testing.

The director is responsible for the reporting of test results. The director assures that reference intervals are appropriate for the laboratory's patient population and that any other information needed for proper interpretation of results is provided to clinicians. The director is accountable for the methods of reporting, whether by electronic means, printed reports, fax, telephone, etc., to assure that test results are accurately transmitted and available when needed by clinicians caring for the patients.

The director is responsible for standard operating procedures for all aspects of testing. The director assures that standard operating procedures are valid, updated as necessary, and accessible to laboratory personnel.

The director is accountable for the selection of reference laboratories for tests that are not performed in the laboratory. The director assures that reference laboratories are licensed and accredited, if required, that reference laboratories provide timely services, and that the laboratory receives the reference laboratories' results and reports them properly to its users.

Quality Management

The laboratory director is accountable for the quality of all services of the laboratory, including the implementation of a quality management system. The quality management system includes an overarching quality plan, quality standards, and a quality manual with policies and procedures. The quality system addresses all aspects of the laboratory service including organization, personnel, equipment, purchasing and inventory, process control, documents and records, information management, error and incident management, assessments, process improvement, customer service and satisfaction, facilities, safety, and so forth.

The director is responsible for quality assessment methods including proficiency testing, inter-method comparisons, audits, surveys, inspections, and incident or problem reports. The director assures that results of assessments are evaluated, opportunities for improvement are identified, and corrective or remedial actions are instituted. The director is responsible for the identification, prioritization, and implementation of quality improvement activities.

Consultation and Education

A visible role of the laboratory director is in the area of consultation and education. As both a laboratory and clinical expert, the laboratory director is in a unique position to help maximize the clinical value of laboratory testing. To do this, the director must be accessible to the medical and laboratory staff.

The director consults with clinicians about test selection and ordering, interpretation of test results, and therapeutic decisions. The director educates clinicians about changes in test methods or test menu, the utility of new tests, and opportunities to improve usage of existing tests. The director fields complaints or concerns from clinicians and assesses clinicians' needs. The wise director pays careful attention to astute clinicians' observations about laboratory results or services. The director participates in the institution's education program, as applicable.

The director consults with laboratory staff on issues related to test methodology, quality control, client satisfaction, vendor relations, and so forth. The director evaluates the clinical and regulatory ramifications of laboratory problems or incidents. The director assures that laboratory staff receive education necessary to perform their duties and enhance the laboratory's services.

Communication

Although most communication is made by managers, supervisors, and frontline staff, the laboratory director, by virtue of his or her position of accountability and authority, plays a critical role in communication with clinicians, patients, lab staff, administration, risk management, government regulators, inspectors, vendors, and others. The director's personal communications can often facilitate the resolution of troublesome situations.

Operational Management

Depending on prevailing regulations, accreditation standards, and contractual provisions, the laboratory director may or may not have direct accountability for operational management; however, the impact of operational management on the director's ability to discharge his or her other duties cannot be ignored and creates a virtual or implied accountability. By virtue of background, training, expertise, and responsibilities, the laboratory director is in a unique and critical position with respect to strategic planning, organizational goal setting, capital and operational budgeting, research and development, marketing, and vendor contracting. The director must champion patients and clinicians in deliberations and decisions related to operational management because, in many cases, no one else is in a position to do so.

Resources

It should be obvious by this point in this chapter that the responsibilities of the laboratory director are broad and deep. The knowledge and skills required are tremendous. Fortunately, laboratory directors are not left without resources.

Other laboratory directors are a great potential resource for practical ideas. Former mentors and respected laboratory directors in the community are generally more than willing to provide feedback or ideas. Professional societies at the local, regional, national, or even international level provide networking opportunities and educational forums for enhancing one's knowledge and skills. Private organizations offer training programs for laboratory directors. Some government agencies also offer training for laboratory directors.

Many publications address one or more aspects of the laboratory director's responsibilities. For example, the Clinical and Laboratory Standards Institute has published guidelines for quality system modeling, quality improvement, training and competency assessment, laboratory design, laboratory safety, technical procedure manuals, proficiency testing, reference laboratory selection and evaluation, cost accounting, assessment of test accuracy, method comparisons, and others¹¹⁻²¹. The CLIA-88 regulations in the Code of Federal Regulations⁹ and the standards of the College of American Pathologists, American Association of Blood Banks, International Organization for Standardization, and others contain much information about contemporary good laboratory practices¹⁻⁸.

Common Pitfalls

Before concluding this chapter, a final note about common pitfalls seems in order. Because the responsibilities of the laboratory director are so numerous, because most responsibilities can be delegated, and because the laboratory director often has other service responsibilities, it is easy for a lab director to fall into one or more traps. The nature and implications of the traps and the means of extrication are self-evident, so the traps are listed without elaboration.

- The laboratory director doesn't understand the scope of his or her responsibilities.
- Hospital administrators, lab managers, lab supervisors, and lab staff don't understand the role and responsibilities of the laboratory director.
- Regulators have a broader interpretation of the laboratory director's responsibilities than the employer or contracting facility.
- The laboratory director's delegation of responsibilities is informal or ambiguous.
- The laboratory director is unaware of problems or issues.
- The laboratory director's follow up is inadequate or non-existent.
- The laboratory director accepts multi-laboratory system directives that conflict with or fail to meet local needs.
- The laboratory director and subordinates are unaware of or don't understand regulations or accreditation standards.
- The laboratory director passively accepts the institution's failure to devote adequate resources to meet service needs and comply with regulations and accreditation standards.

Summary and Key Points

The position of laboratory director bestows many responsibilities requiring a broad set of knowledge and skills in clinical medicine, pathology, clinical laboratory sciences, basic sciences, operations, and quality management. Some key points for the director to keep in mind are:

- The laboratory director is accountable for all aspects of the laboratory.
- Specific accountabilities may be further defined by prevailing regulations, accreditation standards, organizational models, and contracts.
- The laboratory director is the key bridge between laboratory operations and clinical practice.
- Delegation to others is required to fulfill the laboratory director's responsibilities.
- Common pitfalls can be avoided.

By understanding and fulfilling his or her responsibilities, the laboratory director not only assures that the laboratory makes a great contribution to clinical care and patient safety but also obtains the personal benefits of an interesting, challenging, and rewarding position.

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2

Collection of Coagulation Specimens

Sterling T. Bennett

Collection of Coagulation Specimens

Although specimen integrity is important for every laboratory test, coagulation testing that requires plasma specimens seems to be particularly sensitive to even minor deviations from standard practices regarding anticoagulant concentration, container materials, collection technique, centrifugation, and storage. The purpose of this chapter is to describe current recommendations for coagulation specimens and points of ambiguity to assist the laboratory director in establishing standard operating procedures and evaluating the acceptability of non-ideal specimens.

This chapter will emphasize citrated plasma specimens, which comprise the majority of specimens used for routine and special coagulation testing, but will also cover other types of specimens used for hemostasis- and thrombosis-related testing.

Anticoagulant

The recommended anticoagulant for coagulation specimens is 105 to 109 mmol/L (3.13% to 3.2%) trisodium citrate ($\text{Na}_3\text{C}_6\text{H}_5\text{O}_7 \cdot 2\text{H}_2\text{O}$), with or without buffer¹. This is commonly called 3.2% citrate. A higher concentration of citrate (3.8%) used in the past is no longer considered acceptable because it produces longer prothrombin time (PT) and partial thromboplastin time (PTT) results², lower activated protein C (APC) resistance results³, and discrepant International Normalized Ratio (INR) values⁴⁻⁵. Furthermore, International Sensitivity Index (ISI) assignments, for calculating INRs, based on the World Health Organization protocol, are made using specimens collected in 3.2% citrate and are not validated for specimens collected in 3.8% citrate⁵.

The ratio of blood to anticoagulant is important because a relative excess of citrate, from under-filling specimen tubes, prolongs routine coagulation tests⁶. The ideal volume ratio of blood to citrate anticoagulant is 9:1¹, but there is some degree of tolerance for deviations from this ratio⁷. A note of caution is in order. Studies demonstrating acceptable results with specimen volumes less than 90% of the ideal volume have been conducted on only a very limited number of reagents, analyzers, and assays. It seems prudent, therefore, to require tubes to be filled in the range of 90% to 110% of the ideal volume unless published or in-house studies with the laboratory's assay systems support different criteria.

Another circumstance that yields a relative citrate excess is with a high hematocrit, due to the relatively low plasma volume. The Clinical and Laboratory Standards Institute (CLSI) has defined high hematocrit as being greater than 0.55 L/L (55%)¹. In this case, an alteration in the blood-to-anticoagulant ratio in the specimen is required to maintain a suitable citrate concentration in the plasma. One convenient approach is to remove a specified volume of citrate from a coagulation tube, then fill the tube with blood to the usual fill volume. The volume to remove, as a function of hematocrit and tube size, is shown in Table 2.1. For example, if a 3.0 mL tube (0.3 mL citrate and 2.7 mL blood) is to be used and the patient has a hematocrit of 60%, first remove 0.08 mL citrate, leaving 0.22 mL citrate in the tube. Add enough blood to bring the total volume of blood and citrate to 3.0 mL (i.e., add 2.78 mL blood).

It should be noted that CLSI's recommended hematocrit threshold of 0.55 L/L was based on indirect data¹; however, Marlar and colleagues validated this threshold for their laboratory methods by directly comparing paired specimens with adjusted and unadjusted citrate concentration⁸. The effect was more clinically significant with PT and PTT tests than with fibrinogen, factor VIII, protein C activity, or protein C antigen tests. Given the difficulty, expense, and time delays involved in the preparation and use of citrate-reduced tubes, laboratories with a large number of high-hematocrit patients may wish to consider conducting a study to see whether a hematocrit threshold higher than 0.55 L/L can be validated for the laboratory's test methods.

Glass Vs. Plastic Tubes

Over the past several years, the use of plastic specimen tubes throughout the laboratory has been on the rise for several reasons, including lower risk of breakage, which reduces biohazard exposure risk, and lighter

Table 2.1. Citrate Reduction in Specimens with High Hematocrit

| Hematocrit, % | Ideal fill volume of tube (blood + citrate), mL | | | |
|---------------|---|------|------|------|
| | 5.0 | 3.0 | 2.0 | 1.0 |
| 56 | 0.09 | 0.06 | 0.04 | 0.02 |
| 57 | 0.10 | 0.06 | 0.04 | 0.02 |
| 58 | 0.11 | 0.07 | 0.04 | 0.02 |
| 59 | 0.12 | 0.07 | 0.05 | 0.02 |
| 60 | 0.13 | 0.08 | 0.05 | 0.03 |
| 61 | 0.13 | 0.08 | 0.05 | 0.03 |
| 62 | 0.14 | 0.09 | 0.06 | 0.03 |
| 63 | 0.15 | 0.09 | 0.06 | 0.03 |
| 64 | 0.16 | 0.10 | 0.06 | 0.03 |
| 65 | 0.17 | 0.10 | 0.07 | 0.03 |
| 66 | 0.18 | 0.11 | 0.07 | 0.04 |
| 67 | 0.19 | 0.11 | 0.08 | 0.04 |
| 68 | 0.20 | 0.12 | 0.08 | 0.04 |
| 69 | 0.21 | 0.12 | 0.08 | 0.04 |
| 70 | 0.21 | 0.13 | 0.09 | 0.04 |
| 71 | 0.22 | 0.13 | 0.09 | 0.04 |
| 72 | 0.23 | 0.14 | 0.09 | 0.05 |
| 73 | 0.24 | 0.14 | 0.10 | 0.05 |
| 74 | 0.25 | 0.15 | 0.10 | 0.05 |
| 75 | 0.26 | 0.16 | 0.10 | 0.05 |
| 76 | 0.27 | 0.16 | 0.11 | 0.05 |
| 77 | 0.28 | 0.17 | 0.11 | 0.06 |
| 78 | 0.29 | 0.17 | 0.11 | 0.06 |
| 79 | 0.30 | 0.18 | 0.12 | 0.06 |
| 80 | 0.31 | 0.18 | 0.12 | 0.06 |

This table shows the volume of citrate (mL) to remove from common sizes of coagulation specimen tubes prior to specimen collection. After removal of the specified volume of citrate, enough blood is added to fill the tube to the ideal volume.

The volume to remove is given by the formula

$$R = V \left(0.1 - \frac{100 - H}{595 - H} \right)$$

where

R is the volume of citrate to be removed, in mL,

V is the ideal fill volume of the tube (blood + citrate), in mL, and

H is the hematocrit, in %.

Source: Clinical and Laboratory Standards Institute¹.

weight, which reduces shipping and disposal expenses. Regulatory agencies have also strongly encouraged the use of plastic tubes and tube manufacturers have discontinued the production of many types of glass tubes.

Coagulation testing is known to be sensitive to the composition of specimen tubes⁹, and historically, coagulation specimens have been collected in siliconized glass tubes to limit contact activation of clotting factors. Several studies comparing coagulation test results on plastic versus glass tubes found statistically significant differences in PT results between tube types, but in most cases the differences were deemed to be clinically insignificant¹⁰⁻¹¹; however, the differences were deemed clinically significant in one recent study⁹. In another study, Rodgers and colleagues evaluated the effect of plastic tubes on esoteric coagulation tests, including factor assays, thrombin time, and tests for lupus anticoagulant, von Willebrand factor, protein C, protein S, APC resistance, and antithrombin, with normal donor plasma specimens. They found that the only test significantly affected was the thrombin time¹².

Without clear consensus on the suitability of plastic tubes, it is advisable for laboratories to evaluate the effect of plastic tubes on the results of their coagulation assays prior to placing these tubes into routine use.

Collection Techniques

Several options are available for collecting blood specimens for coagulation testing, including venipuncture blood collection systems into evacuated tubes, winged needles and tubing, syringes, vascular access devices (VADs), and capillary specimens. Of these, the recommended method is venipuncture collection directly into tubes containing anticoagulant¹. The other methods all have potential problems that may affect specimen quality. Winged collection systems have a length of tubing whose dead-space may result in under-filled tubes or activation of clotting factors or platelets if not properly managed. Syringe draws may result in activation of clotting factors when there is a delay in transferring blood from the syringe to the anticoagulated specimen tube. Syringe draws may also cause hemolysis if blood is drawn into or expelled from the syringe with too much force. VADs may lead to heparin contamination if the line is heparin-impregnated or has been flushed with heparin or had heparin administered through it. VADs

may also lead to specimen dilution or partially clotted blood if the dead space is not managed appropriately. Capillary specimens are subject to activation of clotting factors if blood is not flowing freely, dilution with extracellular fluid if the skin puncture site is squeezed too vigorously, and hemolysis if the puncture site is overly “milked.”

Historically, a discard sample (or “pilot tube”) was collected prior to the specimen for coagulation testing, due to concern about activation of clotting factors or platelets by the venipuncture procedure. Studies have shown that this practice is not necessary for PT and PTT testing^{1,13-14}. Data do not exist for other coagulation tests, so the need for a discard sample remains controversial, particularly with factor assays. Two caveats apply to the omission of a discard sample. First, with winged collection sets, the long tubing contains a dead space that must be cleared of air to avoid under-filling the specimen tube. A discard sample may be drawn to fill the dead space with blood, but the discard sample itself need not be of any particular volume¹. Second, the dead space of VADs may contain IV fluids, flush fluids, heparin, micro-clots, bacteria, and other undesirables. For this reason, the use of VADs for coagulation specimen collection should be discouraged. If a VAD has ever had heparin in it, it should not be used for collecting coagulation specimens. When a VAD must be used, it should be flushed with at least 5 mL saline, then 5 mL or 6 dead-space volumes should be drawn and discarded prior to specimen collection¹. Discard samples should be collected in non-additive or coagulation tubes or in non-additive syringes. Other additives may contaminate the collection device and affect coagulation test results.

Needle gauge should be based on the amount of blood to be drawn, age of patient, and vein size¹. Small gauge needles on syringes create a temptation to use excessive force when drawing blood into or expelling blood from a syringe, leading to hemolysis. Small gauge needles on winged collection sets may result in slow blood flow through the tubing, with resultant activation of clotting factors and platelets. However, with proper technique, needle gauge need not be an impediment to collection of suitable specimens.

The order in which blood collection tubes should be filled has been defined to avoid problems with cross-contamination of additives between tubes. Cross-contamination of additives can result from drawing additive tubes before non-additive tubes resulting in spurious test results (e.g., contamination of a serum tube with potassium EDTA

will falsely increase the serum potassium measurement and falsely decrease the serum calcium measurement)¹⁵. The recommended order of draw is:

- Blood culture tube
- Coagulation tube
- Serum tube with or without clot activator, and with or without gel separator
- Heparin tube, with or without gel separator
- EDTA tube

A summary of coagulation specimen collection techniques, issues, and cautions is provided in Table 2.2.

Centrifugation

The accepted goal of centrifugation of coagulation specimens is to produce platelet-poor plasma with a platelet count $< 10 \times 10^9/L$ ($< 10 \times 10^3/\mu L$)¹. This may be accomplished by centrifuging specimens at 1500 g for 15 minutes or longer at room temperature. Other centrifugation protocols may be used, but should be validated to assure the production of platelet-poor plasma and comparable results on coagulation assays.

Some coagulation tests are less sensitive to the presence of platelets than others. PT, PTT, and thrombin time (TT) test results are not significantly affected by platelet counts up to $200 \times 10^9/L$ in fresh specimens. However, this does not hold true for specimens containing heparin¹⁶. Tests for lupus anticoagulant, antiphospholipid antibodies, and heparin monitoring require platelet-poor plasma. Data are not available for other tests. Specimens to be frozen must be platelet-poor; otherwise, the disruption of platelet membranes during the freeze-thaw process may affect coagulation test results.

Given the possibility that other tests may be added to specimens initially tested for PT, PTT, or TT and given that laboratories are often unaware of the presence or absence of therapeutic heparin, particularly in specimens from hospital inpatients, it seems prudent to retain the goal of producing platelet-poor plasma with all coagulation specimens.

Table 2.2. Specimen Collection Techniques and Cautions

| Method | Dead-space issues | Discard sample | Other Limitations & Cautions |
|--------------------------------|--|---|---|
| Venipuncture collection system | None. | None. | |
| Winged collection system | Dead-space air needs to be cleared prior to specimen collection. | Needed only to clear air from the tubing. | Slow blood flow through tubing may lead to activation of clotting factors or platelets. Air leaks in connections may cause hemolysis or under-filling. |
| Syringe with hypodermic needle | None. | None. | Excessive force when drawing into or expelling from syringe may cause hemolysis. Blood must be transferred from syringe to anticoagulant tube within 1 minute of collection. Air leaks in connections may cause hemolysis or under-filling. |
| Vascular access device | Dead space may contain IV fluids, micro-clots, heparin, etc. | After VAD is flushed with saline, discard 5 mL or 6 dead-space volumes, whichever is greater. | Due to high risk of specimen contamination, use of VADs should be discouraged. VADs that have ever had heparin in them should not be used. Air leaks in connections may |

(Continued)

Table 2.2. (Continued)

| Method | Dead-space issues | Discard sample | Other Limitations & Cautions |
|-----------|-------------------|-------------------------------------|---|
| Capillary | None. | Do not use the first drop of blood. | <p>cause hemolysis or under-filling.</p> <p>Free flow of blood is needed to avoid activation of clotting factors or platelets.</p> <p>Excessive squeezing or “milking” may cause hemolysis or contamination with extracellular fluid.</p> |

Specimen Stability

Specimen stability varies with the test to be performed, type of specimen required, presence of cells, presence of heparin, and temperature. Stability may also be affected by test methods. Stability data for routine and special coagulation tests are shown in Table 2.3. Laboratories should verify the stability data on their own test systems and adjust their standard operating procedures accordingly.

Problem Detection and Actions

As the preceding sections indicate, a number of serious problems can (and do!) occur with coagulation specimens. It is important to have procedures for detecting and addressing problems that may affect the integrity of test results.

- *Underfilled tubes.* At the time of receipt in the laboratory, the fill level of each tube should be assessed for adequacy. Many laboratories find it helpful to have a chart or marked tube for comparison. If more than one size of tube is used in the institution or by laboratory clients, then it will be necessary to have a comparison chart or tube for each

Table 2.3. Specimen Stability for Coagulation Tests

| Test | Specimen type | Storage Conditions | | |
|-------------------------------|-----------------------|-------------------------|----------------------------|-------------------|
| | | Ambient (18 to 24°C) | Refrigerated (2 to 4°C) | Frozen (-20°C) |
| ACT | Whole blood | 1 min | Unacceptable | Unacceptable |
| Anticardiolipin antibodies | Serum | 2 days | 2 weeks | 1 year |
| Antithrombin | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Anti-Xa heparin assay | Plasma (citrate) | 4 hours | 8 hours | 6 months |
| APC resistance | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| D-dimer | Plasma (citrate) | 4 hours | 8 hours | 6 months |
| Factor V | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Factor V Leiden mutation | Whole blood (EDTA) | 3 days | 1 week | Unacceptable |
| Factor VII | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Factor VIII | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Factor IX | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Factor X | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Factor XI | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Factor XII | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Fibrinogen | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Lupus anticoagulant | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Plasminogen | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Protein C | Plasma (citrate) | 4 hours | 4 hours | 6 months |
| Protein S | Plasma (citrate) | 4 hours | 4 hours | 6 months |

(Continued)

Table 2.3. (Continued)

| Test | Specimen type | Storage Conditions | | | |
|---|---|--|--|---|--|
| | | Ambient (18 to 24°C) | Refrigerated (2 to 4°C) | Frozen (-20°C) | Frozen (-70°C) |
| Prothrombin Prothrombin mutation | Plasma (citrate) Whole blood (EDTA, ACD, citrate, heparin) Plasma (citrate) | 4 hours 3 days 24 hours ^a | 4 hours 1 week Not recommended ^b | 2 weeks Unacceptable 2 weeks | 6 months Unacceptable 6 months |
| PT/INR | | | | | |
| PTT | Plasma (citrate) | 4 hours ^c | 4 hours ^c | 2 weeks | 6 months |
| Ristocetin cofactor TEG [®] | Plasma (citrate) Whole blood Whole blood (citrate) | 4 hours 4 hours 6 min 2 hours | 8 hours Unacceptable Unacceptable | 2 weeks Unacceptable Unacceptable | 6 months Unacceptable Unacceptable |
| Thrombin time | Plasma (citrate) | 4 hours | 4 hours | 2 weeks | 6 months |
| VWF antigen | Plasma (citrate) | 4 hours | 8 hours | 2 weeks | 6 months |
| VWF multimers | Plasma (citrate) | 4 hours | 4 hours | 2 weeks | 6 months |

^a Long stability is for unopened tubes at room temperature, either centrifuged or uncentrifuged.

^b Refrigeration is not recommended due to cold activation of factor VII.

^c Specimens containing heparin should be centrifuged within 1 hour of collection.

Abbreviations: ACT = activated clotting time; APC = activated protein C; PT = prothrombin time; INR = international normalized ratio; PTT = partial thromboplastin time; TEG[®] = thromboelastograph; VWF = von Willebrand factor

Source: Clinical and Laboratory Standards Institute. Collection, transport, and processing of blood specimens for testing plasma-based coagulation assays; approved guideline—fourth edition, document H21-A4. Wayne, PA:CLSI, 2003, and ARUP Laboratories 2005–2006 User's Guide.

size. Underfilled tubes should be rejected and new specimens should be collected.

- *High hematocrit.* After centrifugation, the height of the cell fraction should be used to estimate hematocrit. It is helpful to have a chart or marked tube for comparison. A supplemental approach is to have the hematology laboratory notify the coagulation laboratory of patients with high hematocrits. When the hematocrit is too high, a new specimen should be obtained with a different blood-to-citrate ratio, as described in the anticoagulant section of this chapter. If the hematocrit is borderline high and the coagulation tests are within the reference range, it may be unnecessary to obtain a reduced-citrate specimen.
- *Heparin contamination.* A markedly prolonged PTT with a normal or slightly prolonged PT is the most common clue of heparin contamination, particularly with specimens collected by nursing staff or through VADs. Even if clinicians claim that a patient is not on heparin and that the VAD has never had heparin in it, suspicious results should be evaluated for heparin contamination before a major workup is initiated for factor deficiency or coagulopathy. The presence of heparin may be presumptively made if 1) a heparin assay indicates a concentration of heparin consistent with the test results, 2) test results revert to normal after the specimen is treated with a heparinase or a heparin filtration procedure, or 3) results are normal with a new specimen collected by venipuncture.
- *Clots in specimen.* After centrifugation, each specimen should be observed for the presence of clots and, if present, testing should not be performed.
- *Lipemia.* Lipemia may interfere with optical clot detection methods. If a specimen is severely lipemic or the analyzer indicates interference, the specimen should be retested by a method with mechanical or electromechanical clot detection. Alternatively, the specimen may be ultracentrifuged and the supernatant retested.
- *Icterus.* Icterus may also interfere with optical clot detection. If this occurs, the specimen should be retested by a method with mechanical or electromechanical clot detection.
- *Hemolysis.* After centrifugation, each specimen should be observed for hemolysis. Hemolysis may be either an *in vitro* or *in vivo* phenomenon. *In vitro* hemolysis indicates a problem with specimen collection or handling, and the specimen should not be used because of possible activation of clotting factors¹. The *in vitro* nature of the hemolysis may be inferred if other types of specimens from

the same patient do not show visible hemolysis. A new specimen is required for coagulation testing. *In vivo* hemolysis may occur with hemolytic anemia, disseminated intravascular coagulation, crush injuries, and other conditions. Coagulation testing may be informative, even though the specimen shows visible hemolysis. Communication with the ordering clinician may be useful for determining whether to proceed with testing. If the level of hemolysis interferes with optical clot detection, then mechanical or electromechanical clot detection must be employed to obtain valid results.

Coagulation Testing on Specimens Other than Citrated Plasma

This chapter has focused on collection, preparation, handling, and storage of citrated platelet-poor plasma specimens, but a few coagulation tests require other specimen types. The activated clotting time (ACT), thromboelastograph (TEG[®]), and point-of-care testing PT and PTT tests use fresh whole blood without anticoagulants. The TEG[®] may also be run on citrated whole blood. Specimens for these tests are usually collected by skin puncture, syringe venipuncture or from VADs. All the cautions described above apply with the additional note that the presence of hemolysis or clots is not likely to be detected because the specimens are not centrifuged. So specimen collection techniques need to be even more pristine.

Some fibrin/fibrinogen degradation products (FDP) or fibrin/fibrinogen split products (FSP) tests require serum prepared from whole blood collected in a tube with thrombin/soybean trypsin inhibitor. In this case, it is important that personnel who collect specimens are aware of the special tube requirement to avoid delays in testing and the need for specimen re-collections.

Antiphospholipid antibody testing, including anticardiolipin antibodies, uses serum. Routine specimen collection procedures are appropriate.

Genetic tests, including factor V Leiden mutation and prothrombin G20210A mutation, require EDTA-anticoagulated whole blood, although ACD, citrate, or heparin anticoagulants may also be acceptable. It is important that personnel understand to retain these specimens as whole blood (i.e., uncentrifuged) even though they fall in the domain of the coagulation laboratory.

Common Pitfalls

The potential for problems with coagulation specimens is great. Common hazards that may lead to inaccurate test results include:

- Specimen tubes in an institution have different citrate concentrations.
- Specimen tubes near or beyond their expiration date do not fill adequately due to partial loss of vacuum force.
- Laboratory staff do not have clear criteria for rejecting under- or over-filled tubes.
- New tube types are introduced without validation studies to assess their equivalence with existing tubes.
- Phlebotomy or nursing staff are not adequately trained in collection techniques.
- Specimens are collected through lines that have had heparin in them.
- The discard volumes for line collections are insufficient.
- Centrifugation protocols do not produce platelet-poor plasma with all specimens.
- Changes in centrifuge settings go unnoticed.

Summary and Key Points

Coagulation testing is particularly sensitive to deviations from standard practices regarding anticoagulant concentration, container materials, collection technique, centrifugation, and storage. Some key points for the director to keep in mind are:

- Specimen integrity is vital for accurate coagulation test results. Accordingly, standard operating procedures need to be established for specimen collection and handling as well as for specimen acceptance and rejection.
- Specimen problems are commonplace and serious. When unrecognized, these problems lead to inaccurate or invalid test results. It is vital for laboratories to establish processes for detection of and responding to specimen problems.
- Well-founded recommendations exist and, if followed, will prevent many problems. The data behind the recommendations also provide the laboratory director with guidance for dealing with non-ideal specimens.

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3

Instrumentation for the Coagulation Laboratory

Christopher M. Lehman and Catherine Thompson*

Instrumentation for the Coagulation Laboratory

A broad spectrum of coagulation analyzers is available for purchase on the market today. Laboratories can choose from manual, semi-automated or fully automated, moderate or high-throughput analyzers with narrow or broad assay menus depending upon clinical and test volume requirements. These laboratory-based instruments require plasma prepared from spun, anticoagulated whole blood for analysis. In addition, a number of point of care (POC) devices designed to analyze fresh whole blood are available for use on hospital patient care units, in clinics and doctors' offices, and even in patients' homes. For a current summary of marketed devices, the reader is referred to the annual College of American Pathologists summary list of laboratory-based coagulation analyzers and their attributes published on the College's web site (www.cap.org). A separate list of point of care analyzers can also be found on the site.

Laboratory Instruments

Test Methodologies

Coagulation test methodologies available on current laboratory analyzers vary according to the analyte or process being measured. Routine clot-based assays include the prothrombin test (PT)/International Normalized Ratio (INR), the partial thromboplastin test (PTT), the thrombin time (TT), the activated clotting time (ACT),

* Research and Development Specialist, ARUP Laboratories

coagulation factor assays (including fibrinogen), and lupus anticoagulant tests. Chromogenic assays, tests based on the production of a chromophore secondary to the intrinsic enzymatic activity of the analyte (e.g., antithrombin, proteins S and C, factor VIII), or the inhibition of that activity (e.g., heparin anti-factor Xa activity) are available on many instruments. Finally, immunologic-based testing is also available on some platforms (e.g., D-dimer, von Willebrand factor, protein S). Most laboratory analyzers have an open reagent system that allows use of reagents not produced by the instrument manufacturer. However, the vendor may indicate that the system is optimized for their reagents, and therefore will not guarantee results using other vendors' reagents. Regardless of the source of reagents, each individual reagent-instrument test combination must be validated due to the known variability of test results between reagents – even for different lots of the same reagent, and from analyzer to analyzer – even of the same make and model. In the U.S. market, coagulation instruments, unlike chemistry analyzers, generally don't have the capability for calibration, or don't provide reagents for the purpose of calibration.

Detection Methods

A key factor in assessing the clinical utility of a coagulation analyzer is a consideration of the detection modes available (Table 3.1). Detection methods in laboratory instruments can be classified into two general categories: photo-optical and electro-mechanical. Clot-based, chromogenic and immunologic assays may all utilize a photo-optical detection method, while electro-mechanical detection of clot formation is obviously limited to clot-based assays. Photo-optical detection of clot formation involves measurement of light that has emanated from a source (generally monochromatic) and has passed through, been absorbed by, or scattered from a reaction vessel containing a mixture of reagent, patient sample and fibrin clot. In the case of the PT, PTT, TT and ACT tests, the time required to reach a pre-defined, optical endpoint determines the clotting time. Any factor, other than fibrin clot, that diminishes the relative amount of light passing through the test mixture can adversely affect the result. This includes interfering colored substances (e.g., hemoglobin, bilirubin) that may absorb light at defined frequencies, or suspended particles such as lipoproteins that scatter light. The same substances may also interfere with chromogenic assays that use spectrophotometric detection, as well as immunologic methods that employ turbidometry. Therefore, as noted in Chapter 2,

Table 3.1. Automated Laboratory Coagulation Analyzers

| Vendor | Instrument | Clot Detection Method | Compensation for Interfering Substances |
|----------------------|-----------------------|------------------------------|--|
| Beckman Coulter | ACL TOP, Advance | Photo-optical | Increase in light source intensity |
| BioMerieux | MDA II | Photo-optical | Multiple wavelengths; Flagging |
| Dade Behring | Symex CA-7000 | Photo-optical | Blank measurement |
| | BCS | Photo-optical | Blank measurement; Second wavelength |
| Diagnostica Stago | STA Compact, STA-R | Mechanical | Not Applicable (see text) |
| Helena | Cascade M4 | Photo-optical | Blank measurement |
| Trinity Biotech | Destiny, Destiny Plus | Mechanical | Not Applicable (see text) |
| | | Photo-optical | Use mechanical mode |

Collection of Coagulation Specimens, lipemic, hemolyzed or icteric specimens may interfere with assays employing photo-optical detection methods. The simplest approach to dealing with samples containing interfering substances is to re-collect the specimen to avoid the interference (e.g., avoid hemolysis, avoid post-prandial lipemia). However, this may be either impractical (e.g., patient availability) or impossible (e.g., hyperlipidemic or hemolytic disease). Ultracentrifugation of samples to remove lipoprotein particles is a common approach to dealing with lipidemia. Use of a blank measurement to zero out the interference, measurement at a less susceptible wavelength, or measurement at multiple wavelengths are other potential approaches to modify photo-optical methods to compensate for interferences. An alternative approach is to use a mechanical clot detection system.

Mechanical detection methods employed in automated laboratory analyzers detect clot formation through changes in viscosity (movement of a metal ball), and are generally not considered to be susceptible to interference from colored solutes or suspended particles. However, it is worth noting that large concentrations (> 3 g/dL) of dissolved hemoglobin, in the form of hemoglobin-based oxygen carriers, can cause artificial prolongation of clotting times when employing a

mechanical-based detection method¹. In addition, high concentrations of fibrin degradation products act to impede fibrin strand polymerization, and may result in falsely low fibrinogen measurements^{2,3}.

Point of Care Instruments

Portable coagulation analyzers have been developed primarily to meet the need for whole blood monitoring of out-patient warfarin therapy (PT/INR) and moderate- to high-dose heparin anticoagulant management (ACT) during invasive procedures in the hospital setting. Over time, the manufacturers have extended the test menus of the professional instruments to encompass more of the routine coagulation assays available from the Laboratory (e.g., PTT, TT, fibrinogen, heparin). In addition, portable monitors that measure the viscoelastic properties of clot formation have also been developed for the management of intra-operative hemostasis. More recently, analyzers designed for platelet function analysis have entered the market and are undergoing clinical outcome validation.

Waived PT/INR Instruments

POC monitors intended primarily for warfarin management (Table 3.2) provide whole blood PT/INR testing, and are used primarily for clinic and home monitoring. PT/INR monitors employ unit-use reagents that contain thromboplastin reagents that are assigned an International Sensitivity Index (ISI) based on a comparison between a whole blood device and a reference plasma method. Theoretically, this indirect assignment (plasma to whole blood) could present difficulty in comparing results between plasma and whole blood tests using reagents with comparable ISI assignments. However, comparability is generally quite good when the ISI values are similar⁴. The preferred specimen type is fresh capillary whole blood, though untreated venous blood collected in a plastic syringe is also acceptable for some analyzers. These instruments detect clot formation through electrochemical (impedance) or mechanical (cessation of movement of iron particles; change in rate of blood/reagent mixture movement) methods (Table 3.2). Patients who are being treated with heparin or who have tested positive for a lupus anticoagulant should not be monitored with these devices due to the potential for interference with the test.

Table 3.2. CLIA Waived PT/INR Monitors

| Instrument | Vendor | ISI | Clot Detection | Integrated QC |
|-------------------|-------------------|------------|------------------------|----------------------|
| ProTime | ITC | 1.0 | Microcapillary flow | Yes |
| CoaguChek S | Roche Diagnostics | 2.0 | Iron particle movement | No |
| CoaguChek XS | Roche Diagnostics | 1.0 | Electrochemical | Yes |
| INRatio | Hemosense | 1.0 | Sample impedance | Yes |

Published comparisons of INR values produced by PT monitors and laboratory coagulation analyzers on the same specimens generally demonstrate good agreement in the therapeutic range⁴. POC INR values above 3.0 may diverge from the laboratory instrument value, and the degree and spread of disparity may increase with increasing POC INR. Some clinicians and/or laboratories may decide to confirm POC INR values by the laboratory method at some value above an INR of 4.0. This approach seems prudent since the laboratory analyzer is expected to be more precise, and the POC value is generally more susceptible to sample collection error. However, published data supports the safety of managing patients solely by POC INR⁴⁻⁷. Nonetheless, POC coagulation analyzers must be validated against the laboratory reference method, and clinicians should be made aware of the analytical relationship between the methods, since patients may be tested by different modalities during the course of their routine care.

Unlike the laboratory analyzers, the waived POC PT analyzers cannot interface to a laboratory or hospital information system. This poses a significant risk of transcriptional errors. Hopefully the current emphasis on patient safety and the establishment of a standard format for data transfer⁸ will prompt vendors to add the capacity for electronic data transfer to these analyzers.

POC Analyzers Designed for Coagulation Monitoring during Invasive Procedures

The primary clinical application of these instruments is real time monitoring of temporary heparinization during invasive procedures. They may also be used to direct fresh frozen plasma transfusion therapy during surgical procedures. Real-time heparin management is employed in a variety of settings including cardiac bypass surgery,

percutaneous coronary therapeutic interventions, radiological procedures and extracorporeal membrane oxygenation. Instruments used to monitor heparin are generally less specialized than the PT monitors, frequently offering the PT, the PTT and the ACT tests on the same device. Fresh whole blood is the predominant sample employed by these analyzers. They detect clot formation by mechanical methods (detection of cessation of movement of fluid or iron particles; detection of displacement of a magnet; measurement of resistance to movement of a plunger through the sample), and in one case, by electrochemical (amperometric) detection (Table 3.3).

POC PTT tests can be used to monitor low- to intermediate-intensity heparinization (0.2–1.0 U/mL) or to detect the presence of low levels of residual heparin after cardiovascular surgery. The POC PTT test is a one-stage test, unlike the two-stage test employed in laboratory instruments. Correlations between POC and laboratory instruments have generally been reported to be low enough that results cannot be interpreted interchangeably. This is particularly true for patients treated

Table 3.3. POC Analyzers Designed for Coagulation Monitoring During Invasive Procedures

| Instrument | Vendor | Test Menu | Clot Detection |
|------------------------|-------------------------------|--|-----------------------|
| i-Stat [®] 1 | Abbott | PT/INR, ACT | Amperometry |
| Actalyte XL | Helena | ACT, Max ACT | Magnet Position |
| Actalyte Mini II | Helena | ACT, Max ACT | Magnet Position |
| GEM PCL Plus | Instrumentation Laboratory | PT, PTT, ACT | Fluid Movement |
| Hemochron Signature | ITC | PT, PTT, PTT, ACT | Fluid Movement |
| Hemochron Response | ITC | PT, PTT, PTT, ACT, TT, Protamine dose assay | Magnet Displacement |
| HMS Plus | Medtronic | ACT, Heparin Dose Response, Heparin Protamine titration | Plunger Motion |
| ACT Plus | Medtronic | ACT | Plunger Motion |

Abbreviations: PT = prothrombin time; PTT = partial thromboplastin time; ACT = activated clotting time; INR = international normalized ratio.

with heparin. Therefore, POC-specific cutoffs and ranges should be established for clinical use⁴.

The primary test employed to manage heparin therapy is the ACT test, since this test, unlike the PTT, can be designed to be sensitive to a wide range of blood levels of unfractionated heparin (0.5–8.0 U/mL). Test reagents use either celite or kaolin to activate clotting, and clotting times vary depending upon the activator in the reagent. Celite-based tests generally produce longer clotting times than kaolin-based tests. Clotting times may also vary between ACT analyzers manufactured by different (or the same) vendors depending upon the source of the activator, the formula of the activator, the amount of activator relative to the sample volume, the mixing method, the response to hypothermia and hemodilution, and the response to therapeutic drugs (e.g., aprotinin) administered to the patient⁹. Analyzers generally achieve a level of precision less than 10% (coefficient of variation); however repeat values in individual patients can vary considerably, particularly at higher ACT values¹⁰. Like the PTT, ACT levels are highly variable in their correlation with heparin concentration (anti-factor Xa levels). Published correlations range from approximately $r = 0.2$ to 0.9, depending upon the device, the heparin level and the clinical indication for heparin therapy¹¹. Therefore, there is no reference method that can be used to validate the ACT test. Instrument-specific protocols must be established and validated for each type of clinical procedure, and revalidated if a new ACT system is put into use⁴.

Viscoelastic Analyzers

Analyzers that measure the viscoelastic properties of fibrin polymerization in whole blood have been developed to address the criticism that standard coagulation tests (PT, PTT, and platelet count) are artificial, non-cellular, surrogate measures of an *in vivo*, cell-based (platelet) hemostatic process. In different variations of this type of analyzer (TEG[®], ROTEM[®], and the Sonoclot[®]), blood is allowed to clot in a cup or cuvette with a pin or hollow probe suspended into the patient whole blood sample. The cup (TEG[®]) or pin (ROTEM[®]) is rotated through a narrow angle (~ 4 degrees), or the probe oscillates 1.0 μM up and down (Sonoclot[®]), and the viscoelastic forces generated between the clot and the pin or probe are recorded as a tracing. The shapes of the tracings represent the initial formation of fibrin, the strengthening and

stabilization of the clot, and in the case of the Sonoclot[®], clot retraction (Figures 3.1 & 3.2). Clot lysis can also be recorded in hyperfibrinolytic states.

TEG[®] and ROTEM[®] analyses produce five comparable routine parameters that are derived from the tracing (Figure 3.1): R and CT reflect the latent period between initiation of the test and initial clot formation; K and CFT measure the time from initial clot formation to a defined level of clot strength; α is a measure of the rate of clot strengthening; MA and MCF are a measure of maximum amplitude of the tracing and are related to clot strength; and finally, CL30 and LY30 are a measure of clot lysis 30 minutes after maximum amplitude (MA). Some, but not all studies have found small but significant correlations between these 5 parameters and traditional measures of coagulation such as the PTT (R/CT, K/CFT and α), fibrinogen concentration (R/CT, MA/MCF and α), platelet count (MA), and clot lysis time (euglobulin lysis time)¹⁰. Consequently, protocols for the transfusion of platelets, cryoprecipitate and fresh frozen plasma have been developed around these tracing parameters. Though it might be expected that standard TEG[®] analysis, and specifically the MA parameter, would be sensitive

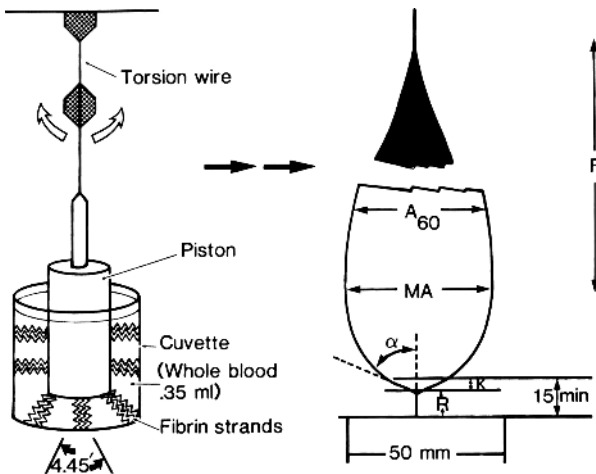


Figure 3.1. Schematic diagram of the Thromboelastograph[®] analyzer and an example of a normal tracing. See text for a discussion of the measured parameters. By permission of Lippincott Williams & Wilkins, *Anesth Analg* 1989;69:69–75.

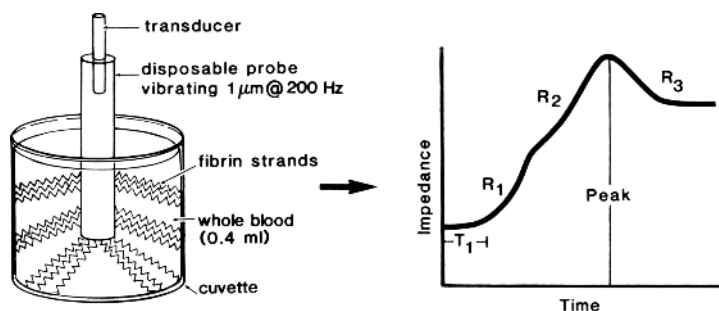


Figure 3.2. Schematic diagram of the Sonoclot[®] analyzer and an example of a normal tracing. See text for a discussion of the measured parameters. By permission of Lippincott Williams & Wilkins, *Anesth Analg* 1989;69:69–75.

to changes in platelet function, TEG[®] has been shown to be insensitive to the platelet effects of aspirin and unable to differentiate the *in vitro* effects of added platelets versus platelet fragments¹².

A number of modifications have been developed to standard TEG[®] analysis. Activators can be added to the whole blood sample to accelerate clotting (e.g., celite) or to eliminate the effect of heparin (e.g., heparinase). Addition of tissue factor dramatically shortens R/CT, and increases clot strength (MA/MCF) presumably through platelet membrane GPIIb/IIIa receptor-fibrin interaction. This modification has been experimentally applied to the evaluation of GPIIb/IIIa inhibitor effects on platelet function^{13,14}.

The Sonoclot[®] tracing yields 5 primary parameters: the onset time, T1 (time to initial clot formation), the R1 slope (the rate of fibrin formation), the R2 slope (reflecting further fibrin production and polymerization plus platelet-fibrin interaction and initial clot retraction), time to peak amplitude (overall rate of fibrin production), and the R3 slope (a result of clot retraction, and clot lysis in cases of hyperfibrinolysis) (Figure 3.2). In the presence of sufficient clot activator, the onset time behaves like an ACT test. The coagulation tests that have been correlated with the Sonoclot[®] parameters are the PT (R1, R2), the PTT (ACT, R1), and the fibrinogen concentration and platelet count (R1 and peak amplitude). Like thromboelastography, Sonoclot[®] analysis is insensitive to the platelet-effects of aspirin, but can detect GPIIb/IIIa receptor inhibition¹⁵. Gender and age have been identified as significant contributing factors to some Sonoclot[®] parameters¹⁶.

Platelet Function Analyzers

While laboratory platelet aggregation analysis remains the gold standard for the evaluation of platelet function, it is not routinely available and is not amenable to bedside analysis. While the template bleeding time has served as a surrogate evaluation of platelet function for many years, the recognition of the bleeding time's poor reproducibility, sensitivity and specificity¹⁷ has led to the development of a number of platelet-function POC analyzers (Table 3.4). As noted above, viscoelastic analyzers can be used to monitor some, but not all platelet function defects.

PFA-100

The PFA-100 is the most studied of this class of analyzers. It is designed to assess platelet function under conditions of high shear. Blood is aspirated through a capillary tube in a test cartridge that contains a membrane coated with either collagen plus epinephrine (CEPI) or collagen plus ADP (CADP). The membranes induce platelet adhesion and aggregation resulting in occlusion (closure) of the central aperture. Results are reported as closure times (CT) in seconds. The analysis requires < 1.0 mL of citrated whole blood for each cartridge tested, and pneumatic tube sample transport is not recommended due to platelet activation. Closure times are not affected by deficiencies of coagulation factors; however, the CT is highly dependent upon the interaction between von Willebrand factor (VWF) and platelet membrane GPIb

Table 3.4. Point of Care Platelet Function Analyzers

| Device | Endpoint | High Shear Flow Measurement |
|-----------------|-----------------------------------|------------------------------------|
| PFA-100® | Closure of membrane aperture | Yes |
| HemoSTATUS™ | Ratio of ACT times | No |
| Plateletworks® | Platelet counts | No |
| VerifyNOW™ | Platelet aggregation | No |
| HAS | Clot generated force | No |
| DiaMed Impact-R | Platelet adhesion and aggregation | Yes |

Abbreviations: HAS = hemostasis analysis system.

and GPIIb/IIIa. Consequently, the PFA-100 lacks specificity in that it can't distinguish between a decrease in VWF levels (i.e., von Willebrand disease) and platelet function disorders. In addition, the PFA-100[®] is insensitive to mild forms of inherited platelet disorders and von Willebrand disease. Therefore, the International Society on Thrombosis and Haemostasis has determined that the PFA-100[®] has insufficient sensitivity and specificity for use in screening patients for platelet disorders¹⁸.

Aspirin ingestion by normal individuals prolongs the CEPI CT, but not the CADP CT, presumably allowing differentiation of drug-induced platelet dysfunction from other causes. However, the CEPI CT is not consistently prolonged in patients taking aspirin for coronary or peripheral vascular disease. GPIIb/IIIa inhibitors prolong both closure times, while the two closure times are not particularly sensitive to the effects of either clopidogrel or ticlopidine¹⁹.

Patient and pre-analytical factors that can prolong the CT include thrombocytopenia ($< 80 \times 10^9/L$), anemia (hematocrit $< 30\%$), blood type O (due to decreased VWF levels), and collection in 3.8% citrate versus 3.2% citrate anticoagulant. Closure times are shortened in newborns^{19,20}, and in the morning versus the evening. Flavenoid-rich foods prolong the CEPI CT. Inconsistency of results between different cartridges lots has been reported to be an issue²¹.

HemoSTATUSTM

This is a kaolin ACT test modified by the addition of platelet activating factor (PAF) to assess relative shortening of the clotting time at 4 levels of PAF. Six ACT tests are performed simultaneously – two controls and the four PAF-added reactions. Clot ratios for the PAF-added tests versus controls are reported, as well as percentage of maximal normal response. The HemoSTATUSTM is designed primarily for monitoring surgical patients, and has been demonstrated to detect a reduction in clot ratio after administration of protamine post-cardiopulmonary bypass, and after desmopressin acetate (DDAVP) or platelet administration post-operatively. However, correlation with post-operative blood loss has not been consistently demonstrated. The HemoSTATUSTM is insensitive to the effects of aspirin and VWF-platelet interactions, but can detect the effects of GPIIb/IIIa receptor inhibitors. The test is influenced by platelet counts less than $50,000/\mu L$ and WBC counts $< 4000/\mu L$ or $> 9000/\mu L$ ^{22,23}.

Plateletworks®

The Plateletworks® assay is based upon the comparison of electronic impedance platelet counts in an EDTA anticoagulated sample versus samples containing one of two platelet agonists – collagen or ADP. The decrease in platelet count in the agonist-containing tubes is directly proportional to the degree of aggregation induced by the agonist. The test output includes a complete blood count including platelet counts, % aggregation and % inhibition. Plateletworks® collagen and ADP aggregation results correlate with standard platelet aggregometry. The analyzer is designed for evaluation of platelet function in patients undergoing cardiovascular procedures.

VerifyNow™

The VerifyNow™ system measures platelet-mediated aggregation of fibrinogen-coated microparticles induced by a platelet agonist. Aggregation is measured as an increase in light transmittance, and results are reported in platelet aggregation units. The test specimen is anticoagulated whole blood. The test cartridges are designed to measure the *in vivo* effect of specific drugs or drug classes on platelet function through the use of appropriate platelet agonists. Current test (agonist) cartridges available assess the platelet function effects of aspirin (arachidonic acid), GPIIb/IIIa receptor blockade (thrombin receptor activating peptide), and the thienopyridine class of drugs that includes clopidogrel (ADP plus prostaglandin E1). Analysis of a pre-drug administration sample is only required for the GPIIb/IIIa test. A limitation of these tests is a lack of evaluation of the effects on test results of platelet counts less than 100,000/ μL , and the presence of inherited platelet disorders.

Hemostasis Analysis System

The Hemostasis Analysis System measures the force generated by a clot as it forms in a narrow gap between the surface of a heated sample cup with a cone-shaped depression and the surface of a suspended, form-fitting sample cone. Three parameters are measured as a function of time. The platelet contractile force (PCF™), clot elastic modulus (CEM), and thrombin generation time (TGT™). The PCF™ measures the force exerted on the sample cone at any point in time by platelet clot retraction. It is dependent upon platelet count, platelet function, temperature, calcium concentration, antithrombin (heparin) activity and

thrombin formation. The CEM is derived from the PCF™ but is sensitive to fibrinogen concentration, platelet count and function, the rate of thrombin generation, and the flexibility of red blood cells. The TGT™ is measured as the time from recalcification of the citrated blood sample to the time when PCF™ begins to increase, and identifies the initiation of thrombin formation. The TGT™ is sensitive to inherited clotting factor deficiencies, and the effects of heparin and warfarin administration²⁴.

DiaMed Impact-R

This device evaluates platelet adhesion and aggregation under more physiologic conditions by simulating laminar flow conditions at arterial pressures. The analyzer operates under the cone and plate principal which involves a cone suspended over a flat, circular plate (like a child's top spinning on a floor), with a narrow angle between the plate and the sides of the cone. Spinning either the cone or the plate produces laminar flow over the surfaces. The anticoagulated sample undergoes laminar flow over a polystyrene sample well. Plasma proteins stick to the polystyrene forming a surface amenable to platelet adhesion and aggregation. The sample well is washed, the adherent platelets are stained, and the instrument automatically records and analyzes the image. No studies evaluating this analyzer have been published to date.

Summary and Key Points

There are a variety of instruments available for coagulation testing in the laboratory. Sensitivity to interfering substances and the levels at which interference occurs are important considerations when selecting a laboratory analyzer. POC instruments that measure the PT/INR, the PTT and the ACT have been integrated into routine patient monitoring during invasive procedures and in the clinic setting. Viscoelastic analyzers have found limited application in the operating room to date. With the exception of the PFA-100®, POC platelet function analyzers are relatively new on the scene, and not well studied. The clinical utility of the POC platelet function analyzers and the viscoelastic analyzers is discussed further in Chapter 7, Testing for Acquired Platelet Function.

- Mechanical detection is less susceptible to commonly encountered interfering substances than photo-optical detection.

- Waived POC PT/INR monitors are suitable for use in oral anticoagulant clinics and, with careful selection of patients, for home monitoring.
- Instrument-specific ACT protocols must be established for each category of invasive procedure requiring heparinization.
- The role of viscoelastic analyzers in platelet function analysis has not yet been defined.
- POC platelet function analyzers have the potential for targeted clinical applications, but data supporting use is preliminary.

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4

Validation of Coagulation Assays, Instruments and Reagents

Sterling T. Bennett

Validation of Coagulation Assays, Analyzers, and Reagents

A key responsibility of the laboratory director is to assure that information and services provided by the laboratory meet the needs of its users and conform to established regulations and standards. With coagulation assays, this assurance should be in place before clinical specimens are tested and results are generated for patient care. The process of providing this assurance is commonly referred to as validation.

Validation Overview

In general terms, validation is the process of assessing whether something meets expectations. In a more formal sense, validation is the documented process of showing that a system is stable and capable of producing pre-determined attributes¹. The need for validating assays, analyzers, and reagents in the coagulation laboratory is obvious, yet there are no universal approaches to or standards for validation. Ultimately, it is the laboratory director who is responsible for the validity of test systems and laboratory processes. The laboratory director has much latitude and must exercise judgment in establishing validation procedures and assessing results of validation studies. In this chapter we will provide guidelines for validation of coagulation test systems and references for additional information.

Validation Tools

Validation needs to be thorough and systematic. Each laboratory should create or obtain a set of tools to assist in this process. Checklists are effective tools for assuring that all important issues are considered. Checklists may also be used to document the results and the laboratory director's approval of validation studies. An example of a checklist for coagulation laboratory validation studies is shown in Figure 4.1. Forms, spreadsheets, and statistical software are additional tools that help assure consistent recording and analysis of data.

Validation Studies

In the coagulation laboratory, validation studies encompass the establishment of analytical performance expectations; determination of quality control ranges, reference intervals, and therapeutic ranges; method comparisons; verification of analyzer and computer set-up; evaluation of reports, both electronic and printed; verification of implementation readiness of laboratory staff and clinicians; and a "green light" from the laboratory director. This set of studies may be organized in a variety of ways, and varies to some extent by the type of assay, level of familiarity with the assay, analyzer, and reagents, and number of components involved. A generic set of validation studies is presented in this section.

Validation Plan

A written validation plan should be developed and approved by the laboratory director or designee prior to the initiation of validation studies. A written plan helps assure that decisions about the extent of validation testing have been made, expectations are clear for all personnel who conduct validation studies, and at the conclusion of validation testing the director will have the information necessary to confidently approve the implementation of the new test or method.

Calibration

Many coagulation assays do not require calibration, but for those that do, it is important that calibration and calibration verification be performed prior to other validation studies. Having a properly calibrated assay helps assure the validity of other validation studies.

| Coagulation Assay, Analyzer, or Reagent Validation | | | | |
|--|--|------------|-----------------------|-----------|
| Laboratory | | | | |
| Assay | | | | |
| Analyzers | Manufacturer | Model | Serial No. | |
| | | | | |
| Reagents | Manufacturer | Name | T. of No. | Exp. Date |
| | | | | |
| Category | Step | Completed? | Document(s) Attached? | |
| Calibration | Assay calibrated | Yes N/A | Yes | No N/A |
| | Calibration verified | Yes N/A | Yes | No N/A |
| Quality Control | Control ranges established | Yes N/A | Yes | No N/A |
| | Control ranges correct in analyzers | Yes N/A | Yes | No N/A |
| | Control ranges correct in LIS | Yes N/A | Yes | No N/A |
| Reference Intervals | Reference intervals established | Yes N/A | Yes | No N/A |
| | Reference intervals correct in analyzers | Yes N/A | Yes | No N/A |
| | Reference intervals correct in LIS | Yes N/A | Yes | No N/A |
| | Critical value limits are correct in LIS | Yes N/A | Yes | No N/A |
| Performance Specifications | Accuracy verified | Yes N/A | Yes | No N/A |
| | Precision determined | Yes N/A | Yes | No N/A |
| | Interferences determined | Yes N/A | Yes | No N/A |
| | Analytical measurement range verified | Yes N/A | Yes | No N/A |
| | Reagent stability determined | Yes N/A | Yes | No N/A |
| | Specimen stability determined | Yes N/A | Yes | No N/A |
| | Factor sensitivities determined | Yes N/A | Yes | No N/A |

Figure 4.1. Example of a validation checklist. (Continued)

| | | | |
|-------------------------------|---|-----------|------------|
| | Lupus anticoagulant sensitivity determined | Yes N/A | Yes No N/A |
| | Heparin sensitivity determined | Yes N/A | Yes No N/A |
| | Carryover determined | Yes N/A | Yes No N/A |
| Method Comparisons | New & current results compared (crossover) | Yes N/A | Yes No N/A |
| | Primary & secondary methods compared | Yes N/A | Yes No N/A |
| INR | ISI and MNPT correct in analyzers | Yes N/A | Yes No N/A |
| | ISI and MNPT correct in LIS | Yes N/A | Yes No N/A |
| | INR calculations verified in analyzers | Yes N/A | Yes No N/A |
| | INR calculations verified in LIS | Yes N/A | Yes No N/A |
| Heparin Therapeutic Ranges | Therapeutic range determined for unfractionated heparin | Yes N/A | Yes No N/A |
| | Therapeutic range determined for low-molecular-weight heparin | Yes N/A | Yes No N/A |
| Instrument Interfaces | Instrument interfaces with LIS working | Yes N/A | Yes No N/A |
| Reports | Results display correctly in LIS | Yes N/A | Yes No N/A |
| | Results display correctly on reports printed from LIS | Yes N/A | Yes No N/A |
| | Results display correctly on reports printed from analyzer | Yes N/A | Yes No N/A |
| | Results display correctly on interfaced information systems | Yes N/A | Yes No N/A |
| | Therapeutic range comments display correctly on all reports | Yes N/A | Yes No N/A |
| Implementation Readiness | Standard operating procedures are correct and approved | Yes N/A | Yes No N/A |
| | Laboratory personnel have been notified and trained | Yes N/A | Yes No N/A |
| | Clinicians have been notified | Yes N/A | Yes No N/A |
| Unresolved Issues or Concerns | | | |
| Approval | Name | Signature | Date |
| Document prepared by | | | |
| Approved by | | | |

Figure 4.1. (Continued)

- Calibrate the assay.

Calibration should be performed in accordance with manufacturer's instructions.

- Verify the calibration.

Calibration verification should be performed in accordance with manufacturer's instructions using verification samples that are different from the calibrators used in the calibration step.

Quality Control Ranges

Validation of quality control ranges consists of verifying that quality control range studies are appropriately conducted, control ranges (or control limits) are properly selected, and control ranges are correctly entered into the analyzers, laboratory information system (LIS), or quality control (QC) application, as applicable.

- Establish quality control ranges.

Some guidelines for establishing quality control ranges are as follows:

- Test controls a minimum of 20 times spread out over at least 3 days. A larger number of results generated over a greater number of days will provide even better control limits. When fewer than 40 results are used to establish a control range, it is a good idea to recalculate control limits later with a larger number of results. It is inadvisable to generate all control values in a single run on one day because the variance of the assay will be underestimated, perhaps grossly underestimated, resulting in control ranges that are unrealistically tight. It is also advisable to simulate the working environment rather than ideal conditions when conducting control range studies. For example, if a control sample may be used for 24 hours after preparation, control studies should use control samples from throughout their "lifespan" rather than only newly prepared samples. Similarly, control studies should encompass the lifespan of reagents.
- Review a scatter plot or run chart of values to evaluate for trends or outliers. If trends are obvious or there are other indications of instability in the assay, then the control range study should be repeated

after the assay has been stabilized. Caution should be exercised in classifying values as outliers. Some good working definitions of an outlier are²:

- 1) An observation whose difference from its closest observation is more than 1/3 of the range of all observations.
 - 2) An observation that exceeds the 3rd quartile (75th percentile) by more than 1.5 times the interquartile range (i.e., the difference between the 75th and 25th percentiles).
- Calculate the mean and standard deviation for each control, and compare these statistics with the manufacturer's information, if any. Substantial deviations from target means or excessive imprecision may indicate problems with the analyzer, reagents, controls materials, or testing procedures, and should be evaluated.
 - Select control limits. Parametric methods are usually employed. For example, the control limits are often set at the mean plus or minus 2 standard deviations³. Other multiples of the standard deviation may be suitable based on the precision of the assay and clinical tolerance for bias and variance in test results³.

See Figure 4.2 for a sample report for evaluating control range studies and selecting control ranges.

A rigorous discussion of quality control may be found in Clinical and Laboratory Standards Institute (CLSI) document C24-A2⁴.

- Verify that control ranges are properly entered into the analyzers, LIS, and QC application.

When analyzers have quality control capabilities, laboratories may find it convenient to use these capabilities for alerting testing personnel of out-of-range results, documenting corrective actions, preparing run charts, and calculating summary statistics. Many laboratories use their LIS for QC management to consolidate quality control from throughout the laboratory and to implement more complex quality control rules. Some laboratories use stand-alone applications for quality control, either in place of or as a supplement to their LIS or because they don't have an LIS. It is not uncommon for quality control to be managed with a combination of applications. It is vital that control ranges be entered correctly and consistently in all applications used.

PT, PTT, and Fibrinogen Control Ranges Study May 2006

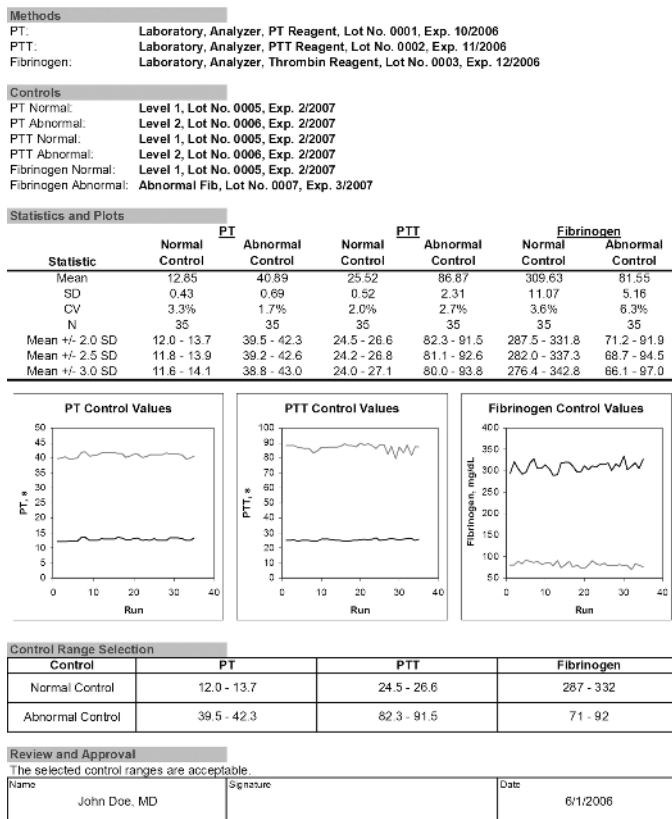


Figure 4.2. Example of a spreadsheet for evaluating quality control results and assigning control ranges.

Reference Intervals

Reference intervals are the principal means by which clinicians interpret test results. Valid reference intervals help assure appropriate triage, therapy, and additional diagnostic studies. Inappropriate reference intervals have been identified as a source of misguided therapy and over-utilization of laboratory tests and other medical services^{5,6}. Validation of reference intervals consists of verifying that reference interval

studies are appropriately conducted, reference intervals are properly selected, and reference intervals are correctly entered into the analyzers and LIS.

- Establish reference intervals.

A number of important steps are required to establish valid reference intervals. Here are some guidelines:

- ◆ Select an equal number of female and male reference subjects. Some laboratories also obtain an equal number of subjects above and below age 45. The minimum number of subjects is 20, but using 40 or more provides a higher level of confidence in the reference intervals. Laboratories that are unable to obtain a satisfactory number of reference subjects may find it valuable to obtain reference specimens from another laboratory, although a dose of caution is in order for tests such as the PTT that may be unreliable with frozen specimens unless they are prepared and handled meticulously.
- ◆ Exclusion criteria should include a history of bleeding or clotting disorders, systemic inflammatory conditions, pregnancy, hormonal contraceptives (oral, implant, or patch), and acute illness. With coagulation testing, the reference population is presumed to be those individuals with “normal” hemostasis. Because of the relatively high prevalence of acquired defects, including prescription and non-prescription medications that affect components of the hemostatic system, use of exclusion criteria with potential reference subjects is critical.
- ◆ Review a scatter plot of the reference values (i.e., test results from the reference subjects). Be cautious about classifying values as outliers, as discussed above.
- ◆ Calculate the mean and standard deviation of the reference values, and calculate the 2.5th and 97.5th percentiles. Compare these statistics with historical values, published reference intervals, and information from the manufacturer. Significant discrepancies from comparable methods and populations should provoke an evaluation of the reference subjects, specimen collection and handling procedures, and assay.
- ◆ Select reference intervals. Either parametric or non-parametric methods may be used, depending on the distribution of test results from the reference subjects.

See Figure 4.3 for a sample report for evaluating reference intervals.

One approach for establishing reference intervals that is often overlooked is that of transference. This is particularly valuable for small laboratories that have a difficult time getting an adequate number of reference subjects to establish valid reference intervals. It may be feasible for the small laboratory to establish comparability of results with a larger laboratory, particularly if the same analytical methods are employed. In this case, it is acceptable for the small laboratory to adopt or “transfer” the larger laboratory’s reference intervals.

An excellent discussion of reference intervals may be found in CLSI document C28-A2².

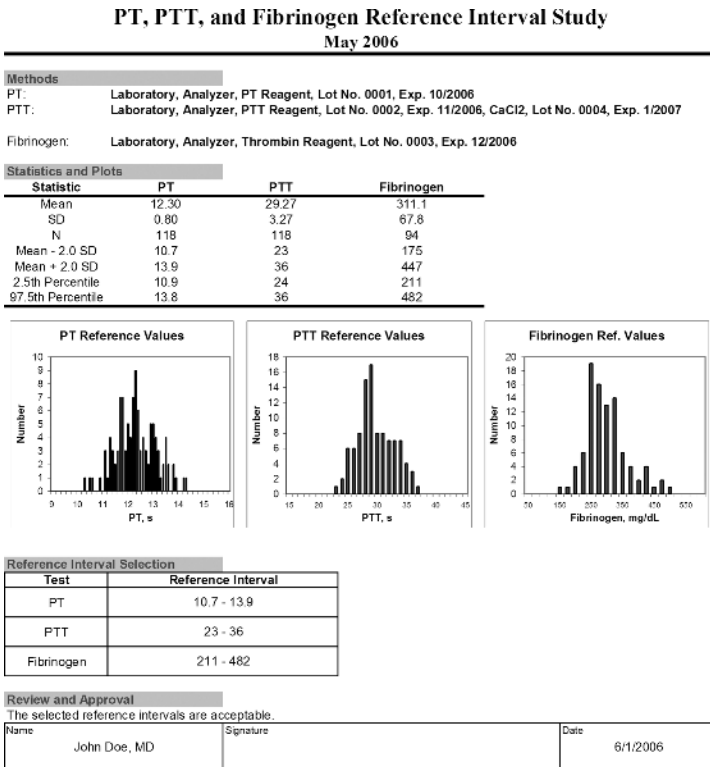


Figure 4.3. Example of a spreadsheet for establishing reference intervals.

- Verify that reference intervals are correct in analyzers and LIS.

Some laboratories find it useful to have the analyzer display reference intervals, when the analyzer has this capability. The analyzer may then be used to flag abnormal results and display reference intervals on analyzer-generated reports. The LIS provides reference intervals on reports and may transmit reference intervals to other interfaced information systems. If interfaced information systems maintain their own reference interval tables, then these also need to be verified. It is critical that reference intervals be entered correctly and consistently in the analyzers, LIS, or other information systems.

Performance Characteristics

Knowing the performance characteristics of an assay is good laboratory practice and, in some countries, is required by regulation. Performance characteristics include analytical accuracy and precision, interferences, analytical sensitivity and measurement range, and methodological limitations or issues such as carryover.

- Verify accuracy.

Accuracy is vital for test results to meet clinical requirements. Detailed procedures have been developed for accuracy verification⁷⁻¹⁰, but for most purposes the combination of calibration verification and analytical measurement range verification will provide laboratories with sufficient evidence of accuracy.

- Determine precision.

Information about precision helps laboratories assess whether differences in replicate results on a specimen indicate problems with the assay. Information about precision helps clinicians assess whether differences in test results on a given patient indicate a change in the patient's status. Detailed methods are available for determination of precision⁹⁻¹¹, but for most purposes, laboratories can gain adequate information about precision from manufacturer's information and quality control statistics.

- Determine interferences.

Determining interferences with an assay is important to avoid spurious results, particularly when the interferences are not detected by the assay system and a result is produced. Full determination of interferences is beyond the scope of this chapter, but for most purposes, laboratories can adequately determine interferences from the medical literature and manufacturer's information.

- Verify the analytical measurement range.

Analytical measurement range verification is required to establish the range over which analyses are acceptably accurate. For some assays, including the prothrombin time (PT), International Normalized Ratio (INR), partial thromboplastin time (PTT), activated clotting time (ACT), and thrombin time (TT), the concept of analytical measurement range is not applicable because there is no standard of "trueness." However, for many other coagulation tests standards do exist and the analytical measurement range can be verified.

Here are some guidelines for analytical measurement range verification:

- ◆ Obtain standards of a suitable matrix with target values or assayed values that cover the range of values over which testing is desired.
- ◆ Test each standard in duplicate and calculate the average of each set of duplicates. A higher number of replicates is useful for imprecise assays.
- ◆ Calculate the absolute and relative (percent) deviations of the averages from the target values.
- ◆ Compare the deviations with defined tolerance limits. Tolerance limits may be defined by the manufacturer, regulations, or published data. In their absence, tolerance limits may be defined by the judgment of the laboratory director.
- ◆ The analytical measurement range is verified at and between those levels at which the deviations are within the tolerance limits.

It is generally considered that if an assay has a theoretical measurement range beyond that for which standards are available, the working analytical measurement range should be limited to that covered by the standards.

- Determine reagent stability.

All reagents degrade over time, potentially leading at some point to incorrect test results. In many cases, laboratories may safely use manufacturer's information or published studies to determine reagent stability. When information does not exist or is not credible, it is incumbent on the laboratory to determine the period of reagent stability.

Here are some guidelines for one approach:

- ◆ Prepare fresh reagent and place on the analyzer.
 - ◆ Prepare and test fresh controls in duplicate initially and at defined intervals. The chosen interval is influenced by the expected stability and intended use of the reagent. For example, if a reagent is expected to be stable and intended to be used over a period of, say, 12 hours, then a suitable interval for testing fresh controls might be 1 or 2 hours. If a reagent is expected to be stable and intended to be used over a period of several days, then an interval of 8 or 12 hours might be chosen.
 - ◆ Prepare a run chart of the averages of duplicates.
 - ◆ The maximum stability is defined by the last time point before any value exceeds control limits or a clear and significant drift in results is evident.
 - ◆ Repeat the above steps with several preparations of fresh reagent. Depending on the analyzer, it may be possible to evaluate multiple preparations simultaneously.
- Determine specimen stability.

As with reagents, specimens also degrade over time, leading at some point to incorrect test results. In most cases, laboratories may safely use manufacturer's information or published studies to determine specimen stability. When information does not exist or is not credible, or the laboratory wishes to extend the period during which specimens can be tested, it is incumbent on the laboratory to determine the stability period.

Here are some guidelines for one approach:

- ◆ Sequentially or randomly select patients' specimens after completion of ordered testing. Hold specimens under defined storage conditions (e.g., ambient or refrigerated).

- ◆ Retest at defined intervals.
 - ◆ Assess the change in results as a function of time from collection.
 - ◆ The maximum stability is defined by the last time point before any value exceeds tolerance limits or a clear and significant drift in results is evident.
- Determine clotting factor sensitivities.

For PT and PTT tests, it is important to know the sensitivity of the assay to various factor deficiencies because insensitive assays are poor screens for clotting factor deficiencies. Accuracy is vital for test results to meet clinical requirements. Often, laboratories can gain adequate information about factor sensitivity from manufacturer's information and published studies. When information does not exist or is suspect in some regard, it may be necessary for a laboratory to determine the sensitivity of an assay to one or more clotting factor deficiencies.

Here are some guidelines for one approach:

- ◆ Prepare a set of specimens with a range of factor levels by mixing different ratios of factor-deficient plasma and normal plasma. Normal plasmas with assayed factor levels are commercially available and readily suited for this type of study. A suitable set of test specimens will have target factor concentrations of 100%, 50%, 40%, 30%, 20%, 10%, 5%, and 2.5%. The actual values may vary from these targets if the factor concentration in the normal plasma is not exactly 100%.
- ◆ Test each specimen in duplicate.
- ◆ Prepare a scatter plot of test results as a function of predicted factor concentration and fit a smooth curve to the data points.
- ◆ Identify the factor concentration at which the result exceeds the reference interval. This is the factor sensitivity of the assay.

See Figure 4.4 for a sample report for identifying the factor sensitivity of an assay from this type of study.

- Determine lupus anticoagulant sensitivity.

PTT reagents differ in their sensitivity to lupus anticoagulants. No consensus exists about the appropriate level of sensitivity; some laboratory directors favor insensitive reagents while others prefer more sensitive reagents. In laboratories that perform lupus anticoagulant

Factor VIII Sensitivity

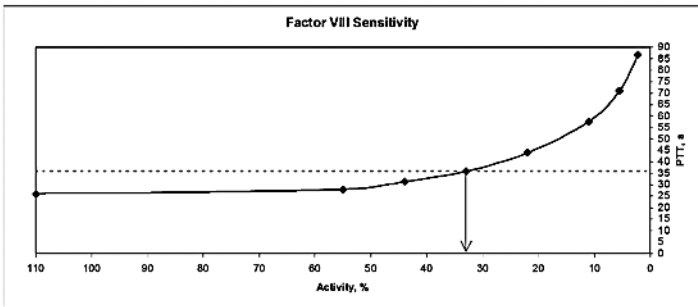
May 2006

Methods

PTT: Laboratory, Analyzer, PTT Reagent, Lot No. 0002, Exp. 11/2006

Data

| Factor VIII Activity, % | PTT | | Mean | Reference Interval Upper Limit |
|----------------------------|-------------|-------------|------|-----------------------------------|
| | Replicate 1 | Replicate 2 | | |
| 110.0 | 26.0 | 26.0 | 26.0 | 36.0 |
| 55.0 | 28.1 | 27.9 | 28.0 | |
| 44.0 | 30.8 | 31.7 | 31.3 | |
| 33.0 | 35.7 | 36.0 | 35.9 | |
| 22.0 | 44.1 | 43.7 | 43.9 | |
| 11.0 | 58.2 | 58.9 | 57.6 | |
| 5.5 | 71.5 | 70.5 | 71.0 | |
| 2.2 | 87.2 | 86.5 | 86.8 | |

**Factor VIII Sensitivity**

| Sensitivity |
|-------------|
| 33% |

Review and Approval

The selected control ranges are acceptable.

| Name | Signature | Date |
|--------------|-----------|----------|
| John Doe, MD | | 6/1/2006 |

Figure 4.4. Example of a report for factor sensitivity determination.

testing, it is feasible to store frozen plasma aliquots from patients with lupus anticoagulants. Specimens with a range of strengths of lupus anticoagulant should be tested with new PTT reagents or reagent lots¹². It is particularly important to include specimens with weak lupus anticoagulants to detect changes in sensitivity. Laboratories without ready access to lupus anticoagulant specimens may be able to obtain information from manufacturers about relative sensitivities.

- Determine heparin sensitivity.

The importance of laboratory monitoring of heparin therapy requires that laboratories assess heparin sensitivity of new PTT reagents and reagent

lots. This may be achieved by comparison with existing reagents or lots or by comparison with an anti-Xa assay¹³. Refer to the Unfractionated Heparin section in Chapter 10 for a detailed discussion.

- Determine carryover.

Many analyzers use a single probe for aspirating all specimens. Carryover from one specimen to another may produce inaccurate test results, so it is important to verify that significant carryover does not occur. Carryover of heparin neutralizers from PT reagents may artificially shorten PTT values in specimens containing heparin. Carryover of heparin may artificially prolong PT or TT results.

Here are some guidelines for one approach to assessing carryover:

- ◆ Establish a baseline PTT or TT on a normal specimen.
- ◆ Test a specimen spiked with a high concentration of heparin.
- ◆ Again test the normal specimen and evaluate for prolongation of the result compared to the baseline.

A more rigorous procedure has been described by Adcock and colleagues¹².

The procedures outlined in this section are suitable for and can be performed in most settings. Rigorous statistical methods for determining performance characteristics may be found in CLSI documents EP5-A2¹¹, EP9-A2⁷, AP10-A2⁸, EP15-A2⁹, and EP21-A¹⁰.

Method Comparisons

Before a laboratory implements a methodological change to an assay, it is important to assess how the change will affect patients' results. A methodological change may be as minor as changing a reagent lot or as major as bringing up a different manufacturer's analyzer and reagent. In either case, there is a potential for patients' results to change. Method comparison studies help assess the potential for clinical impact from the change.

- Perform a method comparison study (crossover) between the current and future methods.

A method comparison study, comparing the current method with the future method, will help answer the question of whether clinicians will

see a change and, if so, the magnitude of the change. This type of study is often referred to as a “crossover.”

Here are guidelines for crossover studies:

- ◆ Test 40 or more patients’ specimens over several days by the current and new methods. Select specimens that cover the range of results typical for the laboratory. A smaller number of specimens may be acceptable and useful if the test results are approximately uniformly spread over a broad range.
- ◆ Review a scatter plot of results. Be cautious about classifying values as outliers.
- ◆ Calculate the correlation coefficient of the pairs of results. Most spreadsheet programs have built-in functions for calculating correlation coefficients.
- ◆ Calculate the orthogonal regression line. Orthogonal regression is preferred over simple least squares regression because it allows for random error to be present in the results of both methods. See Figure 4.5 for an illustration of orthogonal regression. Unfortunately, most spreadsheet programs do not have built-in functions for orthogonal regression, but the calculations are relatively straightforward. The slope of the orthogonal regression line is calculated as follows: Given two data sets $\{x_1, x_2, x_3, \dots, x_n\}$ and $\{y_1, y_2, y_3, \dots, y_n\}$, $slope = u + \sqrt{u^2 + 1}$, where $u = \frac{\sum(y_i - \bar{y})^2 - \sum(x_i - \bar{x})^2}{2 \sum(x_i - \bar{x})(y_i - \bar{y})}$ and $intercept = \bar{y} - slope * \bar{x}$. In addition, most spreadsheet programs contain functions that can be used to simplify the orthogonal regression calculations. For example, the orthogonal regression slope for two sets of values can be calculated in Microsoft Excel (Microsoft Corp., Redmond, WA) using the following formula:

$$= ((\text{VARP}(Y) - \text{VARP}(X)) / (2 * \text{COVAR}(Y, X))) \\ + \text{SQRT}(((\text{VARP}(Y) - \text{VARP}(X)) / (2 * \text{COVAR}(Y, X)))^2 + 1)$$

where X and Y are spreadsheet ranges.

The intercept is found in the usual manner using the equation above.

- ◆ Evaluate the results for systematic shifts and concordance of clinical classifications relative to the reference intervals. Some individuals prefer difference plots, also known as Bland-Altman plots, over scatter plots and regression lines for detecting biases¹⁴. Ideally, there

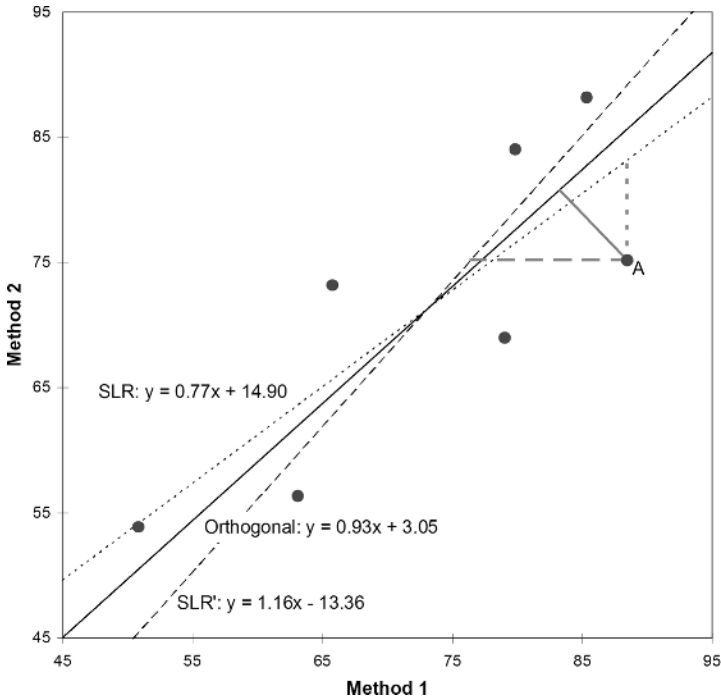


Figure 4.5. Illustration of orthogonal regression versus simple least squares (SLR) regression. The SLR line (---) is calculated under the assumption that the x-values are known without error and the y-values contain random error. The alternate SLR line, SLR' (— — —), is calculated under the assumption that the x-values contain random error but the y-values do not. The orthogonal regression line (——) is calculated under the assumption that both the x- and y-values contain random error. The line segments from point A illustrate that SLR minimizes the sum of the squared vertical distances between the data points and regression line, SLR' minimizes the sum of the squared horizontal distances between the data points and regression line, and orthogonal regression minimizes the sum of the squared perpendicular (or orthogonal) distances between the data points and the regression line. Note that orthogonal regression yields a line between the SLR and SLR' lines.

will be no significant shifts in results or changes in clinical classifications, so the change is invisible to clinicians. However, if there are changes, the method comparison studies provide information about the magnitude of the changes that is invaluable when informing clinicians about the new method.

See Figure 4.6 for a sample report of a crossover study.

- Perform a comparison of the secondary analyzer versus the primary analyzer.

A second type of method comparison is between analyzers in the same laboratory. This is necessary to assure that results with the new method

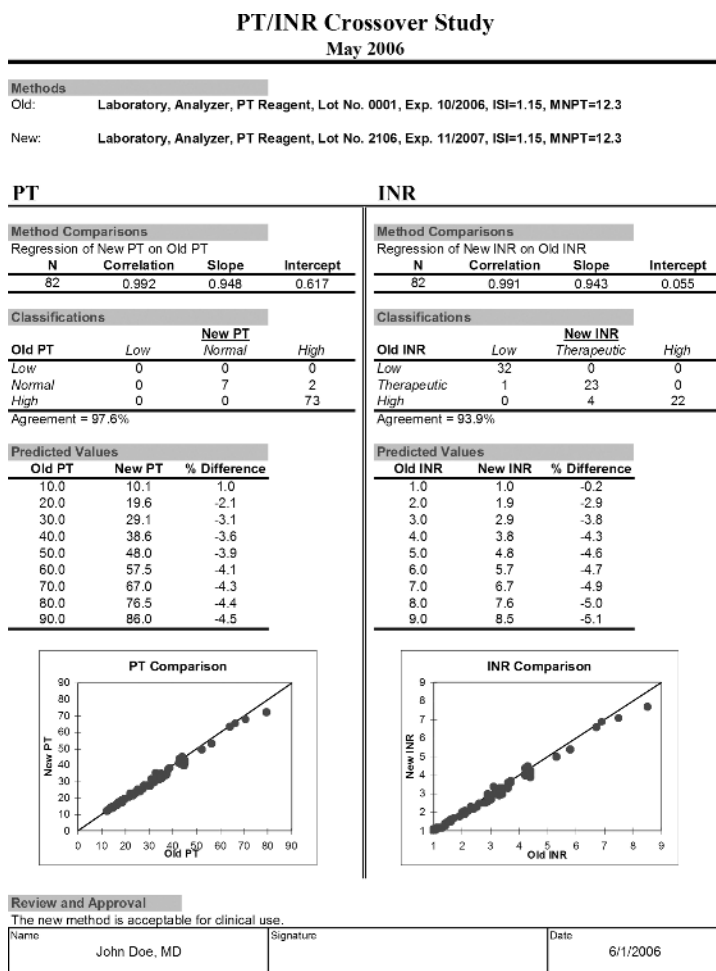


Figure 4.6. Example of a report for method comparison studies comparing a new PT thromboplastin lot with the current lot.

will be equivalent between analyzers. If the results are not equivalent, the method comparison studies provide information to guide the implementation of steps to prevent a negative impact on patient care.

- ◆ Follow all the steps of the method comparison study described above.
- ◆ Assess whether results are sufficiently equivalent. No universal criteria exist for accepting results as equivalent. Criteria may vary with the analytical capabilities of the analyzers, clinical tolerance for differences, and preferences of the laboratory director. Thresholds of 3%, 5%, 10%, and 15% are often used in practice.
- ◆ If results are not equivalent but are highly correlated, adjust the secondary analyzer's results using the regression equation and re-evaluate results. This step may be used to assess whether the secondary analyzer's results should be adjusted mathematically to make them equivalent to the primary method's results.

See Figure 4.7 for a sample report of a study comparing a secondary analyzer with a primary analyzer.

INR

The INR has become the standard for reporting PT values for warfarin anticoagulation therapeutic monitoring. The INR is calculated from the PT using the thromboplastin's ISI (international sensitivity index) and the laboratory's MNPT (mean normal prothrombin time). Refer to Chapter 10, *Monitoring Anticoagulant Therapy*, for a more detailed discussion of the INR. Validation of the INR involves verifying that the correct ISI and MNPT values are used and that calculations are performed correctly.

- Verify that the ISI and MNPT values are correct in the analyzers and LIS.

Some laboratories use the coagulation analyzer to perform the INR calculations, others use the LIS to do the calculations, and some use both. Advantages of using the analyzer to do the calculation are that INRs are viewable on the analyzer at the time of analysis and INRs continue to be calculated when the LIS is down. An advantage of using the LIS to do the calculations is that the ISI and MNPT may need to be entered only once for multiple analyzers. The ISI and MNPT values must be verified in the analyzers and LIS, as applicable.

Primary and Secondary PTT Method Comparison

May 2006

| Methods | | | |
|------------|---|--|--|
| Primary: | Laboratory, Analyzer 1, PTT Reagent, Lot No. 0002, Exp. 11/2006, CaCl ₂ , Lot No. 4, Exp. 1/2007 | | |
| Secondary: | Laboratory, Analyzer 2, PTT Reagent, Lot No. 0002, Exp. 11/2006, CaCl ₂ , Lot No. 4, Exp. 1/2007 | | |

| Raw Data | | | | Adjusted Data | | | |
|--|-------------|---------------|-----------|--|-------------|------------------------|-----------|
| Method Comparisons | | | | Method Comparisons | | | |
| Regression of Secondary PTT on Primary PTT | | | | Regression of Adjusted Sec. PTT on Primary PTT | | | |
| N | Correlation | Slope | Intercept | N | Correlation | Slope | Intercept |
| 92 | 0.981 | 1.091 | -3.806 | 92 | 0.981 | 1.002 | -0.131 |
| Classifications | | | | Classifications | | | |
| | | Secondary PTT | | | | Adjusted Secondary PTT | |
| Pri. PTT | Low | Normal | High | Pri. PTT | Low | Normal | High |
| Low | 0 | 0 | 0 | Low | 0 | 0 | 0 |
| Normal | 0 | 23 | 0 | Therapeutic | 0 | 23 | 0 |
| High | 0 | 0 | 69 | High | 0 | 0 | 69 |
| Agreement = 100.0% | | | | Agreement = 100.0% | | | |
| Predicted Values | | | | Predicted Values | | | |
| Pri. PTT | Sec. PTT | % Difference | | Pri. PTT | Adj. S. PTT | % Diff. | |
| 30.0 | 28.9 | -3.6 | | 30.0 | 29.9 | -0.2 | |
| 50.0 | 50.7 | 1.5 | | 50.0 | 50.0 | -0.1 | |
| 70.0 | 72.6 | 3.7 | | 70.0 | 70.0 | 0.0 | |
| 90.0 | 94.4 | 4.9 | | 90.0 | 90.0 | 0.0 | |
| 130.0 | 138.0 | 6.2 | | 130.0 | 130.1 | 0.1 | |

| Adjustment Procedure Selection | |
|-------------------------------------|--|
| <input type="checkbox"/> | No adjustment is needed. |
| <input type="checkbox"/> | Use the PTT-to-PTT adjustment. |
| <input checked="" type="checkbox"/> | * AdjSecondaryPTT = (SecondaryPTT + 3.806)/1.091 * Set up a worksheet or table to convert secondary PTTs. |

| Review and Approval | | |
|--|-----------|----------|
| The secondary method is acceptable for clinical use. | | |
| Name | Signature | Date |
| John Doe, MD | | 6/1/2006 |

Figure 4.7. Example of a report for method comparison studies between secondary and primary analyzers.

- Verify that INR values are calculated correctly by the analyzers and LIS.

Even if the ISI and MNPT are input correctly, it is possible for the calculations to be set up incorrectly. It is critical that calculations made

by the analyzers and LIS be checked against manual or spreadsheet calculations over a range of values.

Figure 4.8 shows a sample form for documenting the verification of calculations.

Another aspect of INR validation is the verification of its accuracy¹⁵. In most cases, laboratories cannot verify the accuracy of their INRs because they do not have access to reference thromboplastins and reference test methods. In this case, laboratories should at least verify the comparability of their INR values with previous methods or other laboratories. Refer to the Method Comparisons section

| INR Calculation Verification | | | |
|--|--------------|--------------|---------|
| Laboratory | | | |
| Methods | Make & Model | ISI | MNPT |
| Primary Analyzer | | | |
| Secondary Analyzer | | | |
| Test Results, Primary Analyzer | | | |
| PT | Expected INR | Analyzer INR | LIS INR |
| | | | |
| | | | |
| | | | |
| | | | |
| Test Results, Secondary Analyzer | | | |
| PT | Expected INR | Analyzer INR | LIS INR |
| | | | |
| | | | |
| | | | |
| | | | |
| Are all INR values correct? Yes No | | | |
| If no, explain: | | | |
| | | | |
| Approval | Name | Signature | Date |
| Document prepared by | | | |
| Approved by | | | |

Figure 4.8. Example of a form for documenting verification of INR calculations.

of this chapter for additional discussion. In some countries, certified INR plasmas are available and laboratories can assess the accuracy of their INRs over a range of values. Refer to the Performance Characteristics section of this chapter, the INR section of Chapter 10, Monitoring Anticoagulant Therapy, and CLSI document H54-A¹⁶ for additional information.

Heparin Therapeutic Ranges

Laboratory monitoring plays a critical role in the efficacy and safety of unfractionated heparin (UFH) therapy. The PTT assay is used in virtually every laboratory that supports facilities administering UFH therapy. The anti-Xa UFH assay is also available in many laboratories. Laboratory monitoring of low-molecular-weight heparin (LMWH) therapy with the anti-Xa LMWH assay is important for selected patient groups.

- Verify that the standard curve for the anti-Xa UFH assay has been established.

A suitable UFH preparation needs to be used to establish the standard curve. Refer to Chapter 10 for more information about the anti-Xa assay.

- Verify that the PTT therapeutic range has been established for UFH therapeutic monitoring.

The therapeutic range for UFH needs to be established for each PTT reagent/analyzer combination used in the laboratory. With each reagent lot change, the range needs to be verified or re-established. Refer to the Unfractionated Heparin section in Chapter 10, Monitoring Anticoagulant Therapy, for a more detailed discussion.

- Verify that the standard curve for the anti-Xa LWMH assay has been established.

The anti-Xa response of LMWH differs from that of UFH, so separate standard curves of the anti-Xa assay need to be established for LMWH and UFH. The standard curve for the LWMH assay should be established with a LMWH preparation in therapeutic use in the facilities served by the laboratory. Refer to Chapter 10, Monitoring Anticoagulant Therapy, for additional discussion.

Instrument Interfaces

When in place, instrument interfaces form the communication link between the LIS and the analyzers. The LIS transmits orders and patient information to the analyzer; the analyzer transmits results back to the LIS. If the interface is not functioning properly, orders may be missed or results may go unreported.

- Verify that instrument interfaces are working as expected.

Verification of instrument interfaces involves ordering tests on the LIS, tracing these orders to the analyzer, running the tests on the analyzer, and then tracing results back to the LIS. It is wise to challenge both arms of the interface. The ordering arm should be challenged by sending a variety of order types, for example, PT only, PT and PTT, fibrinogen only, fibrinogen and D-dimer, etc. The resulting arm should be challenged with normal and abnormal results, results with a > or < sign, and so forth.

Reports

Laboratories exist essentially to provide information, so reports are the laboratory's product. Reports may be issued in a number of ways, including by print, fax, and electronic display. Validation of reports involves not only test values but also reference intervals, high/low or abnormal flags, comments, and any other information provided in reports that may influence the interpretation of test values.

- Verify that results display correctly on reports printed from the LIS.

Many LISs provide a variety of report options and formats. Each type of report that is used should be validated.

- Verify that results display correctly in the LIS and interfaced information systems.

Electronic display is the primary means for the conveyance of laboratory results in many institutions. The displays on interfaced systems need to be carefully reviewed for their handling of reference intervals, > and < signs, comments, high/low flags, and modified results. When mappings of patient identification numbers or test codes are involved, these also need to be validated.

- Verify that results display correctly on reports faxed from the LIS.

Some LISs have the capability of directly generating faxes, rather than requiring faxing from a printed report. The output of these faxes should be validated for content and readability.

- Verify that results display correctly on reports printed from the analyzer.

Analyzers may be used to generate reports when the LIS is down or in laboratories without an LIS. The displays on interfaced systems need to be carefully reviewed for their handling of reference intervals, > and < signs, comments, high/low flags, and modified results.

- Verify that therapeutic range comments display correctly on all reports.

Many laboratories report target therapeutic ranges for warfarin and heparin therapy along with PT/INR and PTT results, respectively. The handling of this information should be reviewed for all report types and modalities.

Implementation Readiness

Once an assay is validated, a number of other activities need to occur before it is implemented.

- Validate standard operating procedures.

Methodological changes often require modifications to standard operating procedures (SOPs). For example, reference intervals and therapeutic ranges may need to be updated. For major changes, SOPs may need to be largely or entirely rewritten. Each modified SOP needs to undergo a technical validation to assure that following the SOP as written produces the desired result. SOP changes need to be approved prior to implementation of the assay changes.

- Notify and train lab personnel.

Laboratory personnel are the key to a smooth change. They should be notified of the upcoming change far enough in advance to prepare for

the change. Preparation may involve only becoming aware of minor changes or may involve significant hands-on training for major changes. In the validation phase, the important point is to verify that lab personnel are ready to implement the change.

- Notify clinicians.

Clinician notification is vital for any changes that may affect interpretation of results, such as new reference intervals, a systematic shift in results, a change in heparin sensitivity or therapeutic ranges, changes in factor sensitivity, or changes in lupus anticoagulant sensitivity. Laboratory personnel may not be aware of existing clinical protocols that use specified laboratory values as targets or triggers, such as heparin dosing algorithms. Notice should be given far enough in advance that clinicians have time to update protocols prior to the laboratory's implementing a change.

Figure 4.9 shows an example of a clinician notification letter.

Approval

Before implementation of a change, validation studies and documents should be reviewed and approved by the laboratory director or designee.

- Obtain approval of the laboratory director or designee.

Some laboratory directors like to see only summary data and others prefer more detail. Validation processes and documentation can be tailored to the laboratory director's preferences.

“Full” Versus “Partial” Validations

The scope of validation testing depends on the magnitude of the change. Implementing a brand new system obviously requires more extensive validation than changing a reagent lot. It is wholly appropriate for the laboratory director to exercise judgment in the extent of validation that he or she requires. For example, in a reagent lot change, if crossover studies do not show any significant differences in results, it may not be necessary to conduct reference interval studies, therapeutic range studies, or heparin sensitivity studies.

To: <<Facility>> Physicians and Nurse Managers
 From: <<Pathologist, Lab Manager, or Coagulation Supervisor>>
 Clinical Laboratory
 Date: <<Date>>
 Subject: **PT and PTT Coagulation Testing Changes**, Effective <<Starting Date>>

Please note the following information and share it with your staff.

On <<Starting Date>>, the Clinical Laboratory will be changing analyzers and reagents for prothrombin time (PT) and activated partial thromboplastin time (PTT) testing. Our studies comparing the new test methods with the current test methods show the following:

Reference Ranges

The reference ranges will change for both the PT and PTT.

| Test | New Reference Range | Old Reference Range |
|------|---------------------|---------------------|
| PT | nn.n – nn.n sec | nn.n – nn.n sec |
| PTT | nn – nn sec | nn – nn sec |

Therapeutic Ranges

Therapeutic ranges for the INR (for monitoring coumadin therapy) will not change. PTT values corresponding to unfractionated heparin concentrations of 0.2 and 0.4 U/mL are listed below.

| Test | To Monitor | Therapeutic Ranges |
|----------|------------------------|---|
| INR (PT) | Warfarin | 2.0 - 3.0 (moderate intensity) 2.5 - 3.5 (high intensity) Individual patients may require higher or lower ranges. |
| PTT | Unfractionated Heparin | nn – nn sec (heparin levels of 0.3 and 0.7 U/mL anti-Xa activity) |

Comparison of Results

With the new methods, testing for most patients is expected to show the following trends (individual patients may show larger or smaller differences):

- PT results will
- INR results will
- PTT results will

If you have questions or comments about these changes, please contact <<local contact>>.

Figure 4.9. An example of a clinician notification letter.

Other Considerations

Introduction of a new assay or modification of an existing assay may have other ramifications than those discussed in this chapter^{12,17}. For example, proficiency testing orders may need to be altered, client manuals may need to be updated, or regulatory agencies may need to be notified. These items may be added to the laboratory’s validation checklist or handled another way deemed appropriate by the laboratory director.

Common Pitfalls

Although the concept of validation is intrinsically understood, in practice it may be difficult to remember everything that needs to be done unless a systematic approach is developed and followed. Some of the more common problems with validation are:

- One of more of the validation studies are overlooked and not performed.
- Validation studies are inadequate.
- Calculations are not validated.
- Information system validation is not performed.
- Changes are not communicated effectively to clinicians.

Summary and Key Points

Validation is the process of showing that a system is stable and performs according to expectations. In the coagulation laboratory, validation encompasses establishing analytical performance expectations, determining limits for quality control, reference intervals, and therapeutic monitoring, assessing the clinical effects of methodological changes, verifying the setup of computer systems and analyzers, verifying the content and readability of reports, and assuring readiness for implementation. Some key aspects of this process are:

- Validation is critical to assure intended results.
- “Validation” means different things to different people. Defined procedures and analytical tools help assure the director’s definition of validation is met.
- Checklists help “error-proof” the validation process.

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5

Hemostasis Screening Assays

George M. Rodgers and Christopher M. Lehman

Hemostasis Screening Assays: Use and Interpretation

The cornerstone assays of coagulation testing are the prothrombin time (PT) and partial thromboplastin time (PTT). When combined with results of the platelet count obtained from the complete blood count (CBC), differential diagnoses can be generated to assist in the evaluation of patients with bleeding disorders. This chapter will summarize methodologies for the PT and PTT assays, as well as other clinically useful tests that many hospital laboratories provide. An approach to diagnosing bleeding disorders using results of the PT and PTT assays (and the platelet count) will also be presented. Lastly, limitations of these assays will be discussed in the context of evaluating patients for bleeding.

The CBC

The CBC is discussed in this chapter because the platelet count component of the CBC is a useful parameter in the evaluation of hemostasis. Details of platelet count testing are discussed in Chapter 7, Testing for Acquired Platelet Disorders. Typically, EDTA-anticoagulated blood is obtained for analysis in an automated particle counter. The reported platelet count is usually quite precise ($CV \sim 5\%$). In asymptomatic patients in whom thrombocytopenia is reported, the possibility of pseudothrombocytopenia or EDTA-induced thrombocytopenia should be considered, especially in patients without a history of bleeding. This phenomenon occurs in 0.1–1% of normal people; it results from EDTA modifying platelet membrane proteins which

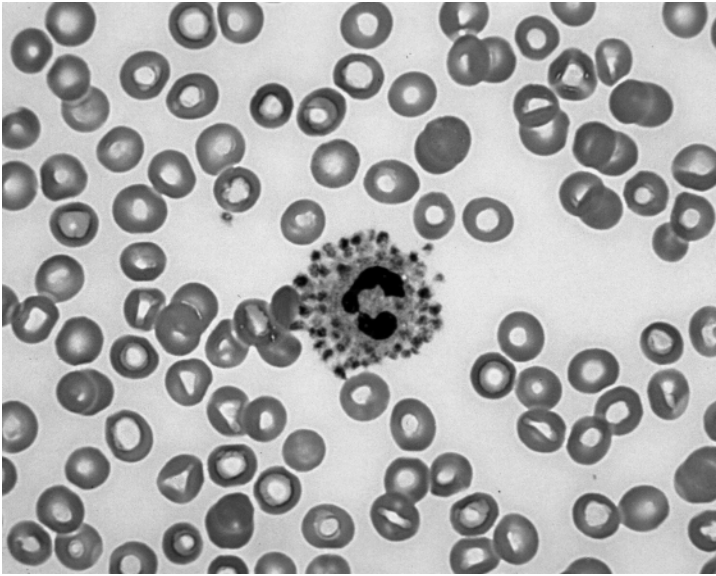


Figure 5.1. Platelet satellitism and platelet clumping in a blood smear from a patient with EDTA-pseudothrombocytopenia (Courtesy Sherrie L. Perkins, MD, University of Utah Medical Center [Wright's stain, oil immersion]). From Rodgers GM, *Case Studies in Hemostasis*, ASCP Press, 2000, p. 161, with permission.

then react with preexisting antibodies present in patient blood that recognize the modified platelet proteins, producing platelet clumping or satellitism (Figure 5.1). It should be routine laboratory policy for technical personnel to review peripheral blood smears of patients with newly diagnosed thrombocytopenia to determine whether the thrombocytopenia is true or spurious. If EDTA-induced thrombocytopenia is suspected, the CBC should be repeated using blood collected in a citrate or Acid-Citrate-Dextrose collection tube (see Chapter 7, Testing for Acquired Platelet Disorders). Figure 5.2 illustrates an algorithm that suggests one strategy to evaluate thrombocytopenia.

In terms of hemostasis evaluation, one limitation of the CBC is that even though it is usually a reliable indicator of the platelet count, the CBC does not measure platelet function. The bleeding time test was originally thought to perform this function, but as discussed below, is not uniformly reliable in assessing platelet function.

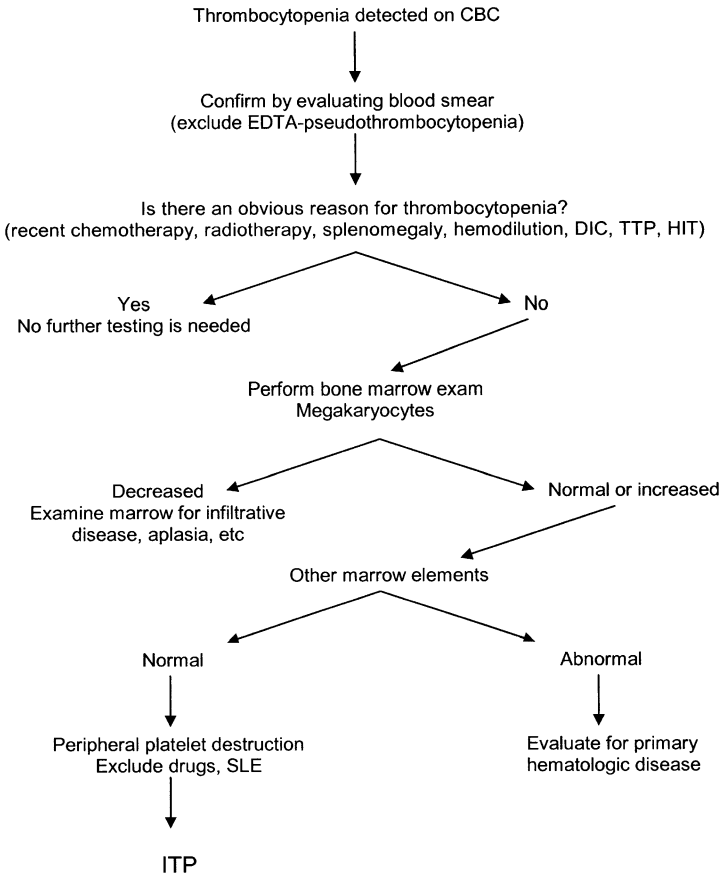


Figure 5.2. A strategy to evaluate thrombocytopenia. Algorithm for evaluation of thrombocytopenia. Abbreviations: DIC = disseminated intravascular coagulation; TTP = thrombotic thrombocytopenic purpura; SLE = systemic lupus erythematosus; HIT = heparin-induced thrombocytopenia; ITP = idiopathic thrombocytopenic purpura. The laboratory evaluation for DIC, TTP, etc. is discussed in Chapter 8, Acquired Coagulation Disorders and TTP and Chapter 9, Testing for Inherited and Acquired Thrombotic Disorders. Modified from Rodgers GM. Thrombocytopenia. In Kjeldsberg CR, ed. Practical Diagnosis of Hematologic Disorders. ASCP Press. Chicago, 2006, Fourth ed., with permission.

The PT Assay

The PT assay has two purposes: to screen for inherited or acquired deficiencies in the extrinsic and common pathways of coagulation (Figure 5.3) and to monitor oral anticoagulant therapy. The PT is affected by decreased levels of fibrinogen, prothrombin, factors V, VII, or X. Since 3 of the 5 coagulation factors measured by the PT are vitamin K-dependent proteins (prothrombin, factors VII and X), the PT assay is useful in detecting vitamin K deficiency from any cause including liver disease, malnutrition, or warfarin therapy. The PT does not measure factor XIII activity or components of the intrinsic pathway¹.

The PT assay is performed by mixing patient plasma with thromboplastin. Thromboplastin is a commercial tissue factor/phospholipid/calcium preparation, which is derived either from animal tissue or from recombinant methods. Tissue factor in the thromboplastin preparation binds factor VII in patient plasma to initiate coagulation. The clotting time is measured in seconds using instruments with mechanical or photo-optical endpoints that detect fibrin formation². Thromboplastin preparations can vary in their sensitivities, resulting in different clotting times. This is discussed further in Chapter 10, Monitoring Anticoagulant Therapy. A typical PT reference range is 10–15 sec. Most PT assays are automated, with the instrument adding reagents and patient plasma samples per protocol. Manual finger stick methods to measure PT/INR values are available and are discussed in Chapter 3, Instrumentation for the Coagulation Laboratory.

In general, the PT assay is more sensitive in detecting low levels of factors VII and X than low levels of fibrinogen, prothrombin or factor V. In particular, different thromboplastin reagents may exhibit variable sensitivities to these factor deficiencies. Mild factor deficiency (i.e., 40–50% of normal) may not be detected by many thromboplastin reagents. The PT assay is less affected by heparin than is the PTT assay, but therapeutic doses of unfractionated heparin will usually prolong the PT by a few seconds unless a heparin neutralizer is present in the PT reagent^{1,2}.

Shortened PT values may result from either a poor quality venipuncture (activated sample), chronic disseminated intravascular coagulation (in vivo activation), or cold activation of the sample (in vitro activation from factor XII activation of factor VII) that occurs if the plasma sample is stored at cold temperatures (above freezing) for several hours. Administration of recombinant factor VII_a also will decrease the PT.

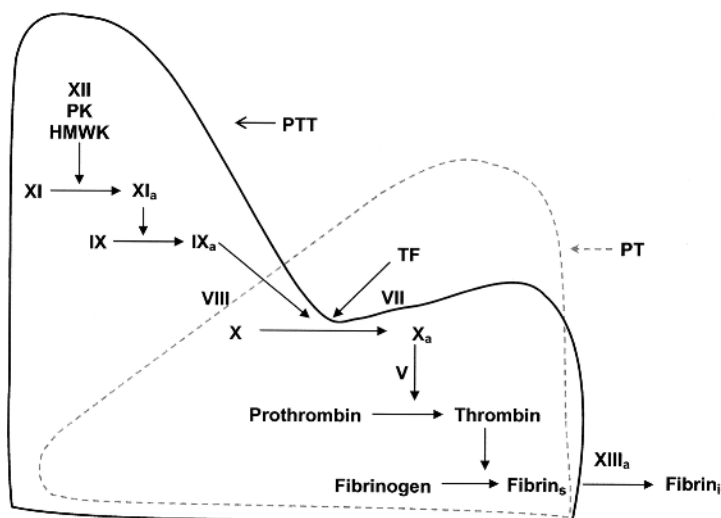


Figure 5.3. The coagulation mechanism as measured by the PT and PTT assays. The PT assay measures the extrinsic (tissue factor) and common pathways, and is performed in the laboratory by mixing patient plasma with a commercial preparation of tissue factor and calcium (thromboplastin). This results in tissue factor-factor VII activation of factor X, then factor X_a-mediated conversion of prothrombin to thrombin. This reaction requires factor V as a cofactor. Thrombin converts fibrinogen to soluble fibrin, which polymerizes into fibrin strands, the endpoint detected by the coagulation instrument. Factor XIII_a cross-linking of fibrin is not measured by the PT assay. The PTT assay measures the intrinsic and common pathways, and is performed by adding patient plasma to the PTT reagent (contact activator). This preincubation step initiates contact activation of plasma in which factor XII and factor XI are activated in the presence of prekallikrein and high-molecular-weight kininogen. Factor XI_a then activates factor IX to IX_a. Calcium is then added to the sample. This results in factor IX_a-mediated activation of factor X in the presence of the cofactor, factor VIII, with subsequent activation of prothrombin and fibrinogen as described above. As with the PT assay, the endpoint of the PTT assay is generation of polymerized fibrin strands, so that factor XIII activity is not measured by the PTT assay. (Abbreviations: PK = prekallikrein, HMWK = high-molecular-weight kininogen, TF = tissue factor, fibrin_s = soluble fibrin, fibrin_i = insoluble fibrin) Modified from Rodgers GM. *Diagnosis of Bleeding Disorders*. In Kjeldsberg CR, ed. *Practical Diagnosis of Hematologic Disorders*. ASCP Press. Chicago, 2006, Fourth ed., with permission.

The PTT Assay

The PTT assay is useful for three reasons – as a screening test for inherited or acquired deficiencies of the intrinsic pathway (Figure 5.3), to detect the lupus anticoagulant, and to monitor heparin therapy. The PTT is affected by decreased levels of intrinsic pathway components (factors VIII, IX, XI, XII, prekallikrein, and high-molecular-weight kininogen), as well as decreased levels of common pathway components (fibrinogen, prothrombin, factor V and X). Factors VII and XIII are not measured by the PTT assay¹.

To perform the PTT assay, patient plasma is preincubated with the PTT reagent (crude phospholipid and a surface-activating agent such as silica or kaolin). This preincubation initiates contact activation (intrinsic pathway activation) in which factors XII and XI are activated in the presence of cofactors, prekallikrein and high-molecular-weight kininogen. Factor XI_a then converts factor IX to IX_a. Calcium is then added to the preincubation mixture; this results in factors IX_a/VIII activation of factor X, then factor X_a/V-mediated activation of prothrombin to thrombin followed by conversion of fibrinogen to soluble fibrin that polymerizes into fibrin strands, the endpoint of the PTT assay^{1,2}. As with the PT assay, most PTT assays are automated.

Mild factor deficiency (i.e., 30–50% of normal) may not be detected by most PTT reagents, but deficient levels of 10–20% should be detected. The usual PTT reagent is less sensitive to factor IX than to factors VIII, XI, and XII. The ability to detect mild factor deficiency is reagent-dependent, and this is an important consideration when choosing the laboratory PTT reagent (see Chapter 4, Validation of Coagulation Assays, Instruments and Reagents). For example, a hospital laboratory that evaluates a large population of bleeding disorder patients may prefer a PTT reagent that is more sensitive to detection of factor deficiency than to detection of lupus anticoagulants. In contrast, a hospital laboratory that evaluates large numbers of obstetrical patients may prefer a PTT reagent that has the opposite characteristics. The PTT may be affected by high levels of factor VIII, an acute-phase response protein; high factor VIII levels may mask co-existing mild intrinsic coagulation deficiencies. A typical PTT reference range is 25–36 sec.

Shortened PTT values may result from a poor-quality venipuncture (activated sample), chronic disseminated intravascular coagulation (*in vivo* activation), or increased factor VIII levels.

It should be emphasized that the PT and PTT assays are *screening* tests, that normal PT/PTT results do not exclude a bleeding disorder, and that many patients with mild factor deficiency will have normal results for these assays¹.

The Mixing Test (Inhibitor Screen)

Abnormal PT or PTT results are due either to deficiency of a factor measured by the assay, or by an inhibitor such as an antibody or heparin. Uncommon inhibitors include fibrin degradation products and certain paraproteins. The mixing test is useful to distinguish between deficiency vs inhibitor, and mixing test results usually suggest subsequent test ordering. The most common mixing test protocol mixes patient plasma with normal plasma in a 1:1 ratio, followed by repeat assay of the PT or PTT immediately after mixing and repeated 1–2 hours later. Sample mixes that correct to normal at both intervals suggest that the original abnormal result was due to factor deficiency, while sample mixes that fail to correct to normal at one or both intervals suggest the presence of an inhibitory substance¹. If heparin is suspected as the inhibitor, screening tests for the presence of heparin can be done with the thrombin time assay and reptilase assay (discussed below) or by direct assay of heparin or low-molecular-weight heparin using anti-factor X_a methods (see Chapter 10, Monitoring of Anticoagulant Therapy).

The mixing test is most useful for evaluating prolonged PTT results. Almost all prolonged PT samples result from factor deficiency, and the mixing test is less useful in this circumstance.

The Thrombin Time Assay

The thrombin time (TT) measures the conversion of fibrinogen to fibrin. It is performed by the addition of purified thrombin to patient plasma; the resulting clotting time is a function of fibrinogen concentration and activity. The TT is a screening test for the presence of heparin in a plasma sample. Other causes for a prolonged thrombin time include quantitative deficiency of fibrinogen, qualitative abnormality of fibrinogen (dysfibrinogen), elevated levels of fibrin degradation products (FDP), the presence of certain paraproteins, and markedly increased levels of fibrinogen ($> 5 \text{ g/L}$)(1).

If heparin is suspected as the cause of a prolonged TT, confirmation of heparin presence can be made using heparin assays (see

anti-X_a assays in Chapter 10, Monitoring of Anticoagulant Therapy) or by using the reptilase clotting time that also measures the conversion of fibrinogen to fibrin. The reptilase clotting time is not affected by heparin. Therefore, a plasma sample with a prolonged TT, but a normal reptilase clotting time indicates the presence of heparin.

For hypofibrinogenemia to prolong a TT, the fibrinogen value will usually be $\leq 0.7\text{--}1\text{ g/L}$. If the thrombin concentration in the TT assay is more than 3 U/mL, the assay will be less sensitive in detecting abnormalities.

Fibrinogen Assays

Fibrinogen is a heterodimeric molecule, with each half of the molecule composed of three polypeptide chains - A α , B β and γ . It is acted upon by thrombin to produce fibrin monomers that polymerize to form fibrin strands, and ultimately a fibrin clot. Circulating fibrinogen molecules are structurally heterogeneous, and not all fibrinogen molecules are capable of participating in clot formation. Therefore antigenic assays and clot-based assays may return different results depending upon the composition of fibrinogen molecules in a specific patient sample³. Only clottable fibrinogen is of interest for the purpose of hemostasis screening.

Fibrinogen assays available on current automated coagulation analyzers include the Clauss and PT-derived methods. The Clauss method is a modified thrombin time. To initiate clotting, excess thrombin is added to patient plasma that has been diluted with buffer. The clotting time is proportional to the fibrinogen concentration. The dilution with buffer decreases the effects of interfering substances (e.g., heparinoids, FDPs) on the Clauss reaction, relative to the thrombin time assay. However, most manufacturers only claim to inhibit interference from heparin up to a concentration of 1–2 U/mL. Higher concentrations may result in falsely low fibrinogen measurements. FDPs generally result in decreased fibrinogen estimates by the Clauss method, as well, though the degree of interference varies by manufacturer. Some manufacturers include FDP inhibitors in their reagents^{3,4}. Since Clauss assays have excess thrombin in the reagents, carryover of thrombin to subsequent tests is a potential problem that should be addressed during instrument validation.

In the PT-derived assays, the fibrinogen concentration is proportional to the total change in optical signal observed during the PT assay.

PT-derived assays have the advantage that a fibrinogen level can be derived directly from the PT assay without additional time or expense. In addition, FDPs generated by thrombolysis do not interfere with these assays^{3,4}. However, these assays are not recommended for routine use in the clinical laboratory since they have lower precision than the Clauss assays, and suffer from decreased accuracy at low or high fibrinogen concentrations, or when turbid plasma is tested⁴.

In most circumstances, the PT, PTT and platelet assays are sufficient for screening bleeding patients. However, in cases where fibrinogen levels may drop precipitously (e.g., disseminated intravascular coagulation [DIC] in obstetric patients), a fibrinogen assay is an essential screening test, since the PT and PTT are relatively insensitive to low levels of fibrinogen (see Chapter 11, Coagulation Testing and Transfusion Medicine).

D-dimer Assays

D-dimer is formed through the proteolytic action of plasmin on polymerized fibrin that has been cross-linked by factor XIII (Figure 5.4). The presence of D-dimer in the circulation is evidence that clot has formed and has been cleaved by plasmin. As such, D-dimer is an effective screening assay for two conditions: DIC and venous thromboembolism (VTE). In DIC, excessive thrombin is formed resulting in clot formation and activation of the fibrinolytic system that, in turn produces D-dimer (see Chapter 8, Acquired Coagulation Disorders and TTP). In VTE, clot forms in the deep venous system of the pelvis and/or proximal, lower extremities secondary to pre-disposing conditions (Virchows triad). Clots may then embolize to the lungs. Plasmin cleaves cross-linked fibrin polymers in the clot releasing D-dimer into the circulation.

D-Dimer assays are immunoassays with different antibody specificities for the heterogeneous D-dimer fragments produced by the action of plasmin on cross-linked clot (Figure 5.4). Therefore, D-dimer levels produced by different assays are generally not interchangeable. The sensitivity of D-dimer assays is judged relative to reference ELISA assays performed in microtitre wells. The manual agglutination assays are the least sensitive tests, while the automated ELISA and immunoturbidimetric assays are the most sensitive, rapid assays. Any of the available assays is suitable for screening for DIC. Many patients, particularly hospitalized patients, have low levels of plasma

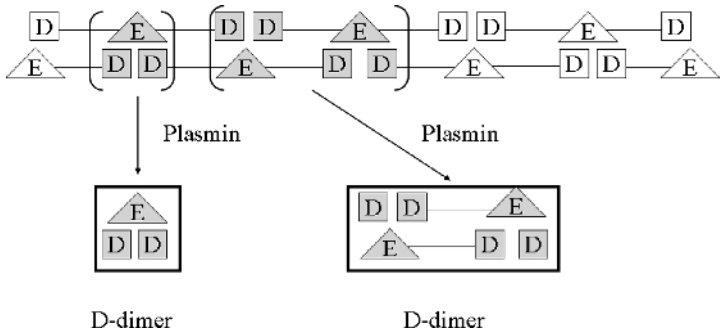


Figure 5.4. Proteolytic action of plasmin on cross-linked fibrin polymer producing heterogenous D-dimer molecules. Abbreviations: D = D domain of fibrinogen, E = E domain of fibrinogen.

D-dimer that exceed the reference interval, but they do not have clinical evidence of DIC. Therefore, assay-specific cutoffs should be established to maximize sensitivity and specificity for the diagnosis of DIC⁵.

Sensitive D-dimer assays have been demonstrated to have excellent negative predictive value for the diagnosis of VTE, when combined with a pre-test probability assessment⁶. Manual agglutination assays are insufficiently sensitive to be used to rule out VTE.

The Bleeding Time (BT) Test

The BT test was developed to provide information on platelet function and the likelihood of bleeding with surgery or invasive procedures. However, an extensive literature on the BT has evolved indicating that the test has minimal clinical utility, including a position paper by the College of American Pathologists and the American Society of Clinical Pathologists¹. A clinical outcomes study published in 2001 found that when the BT test was discontinued at a tertiary medical center, there were no negative impacts, including no increase in procedural or surgical bleeding⁷. The authors of that study concluded that appropriate evaluation for patients suspected of having platelet dysfunction should focus on the patient and family history of bleeding, as well as specific tests for platelet dysfunction, including laboratory evaluation of von Willebrand disease and platelet aggregation studies. Figure 5.5 summarizes an algorithm used to evaluate patients for platelet dysfunction.

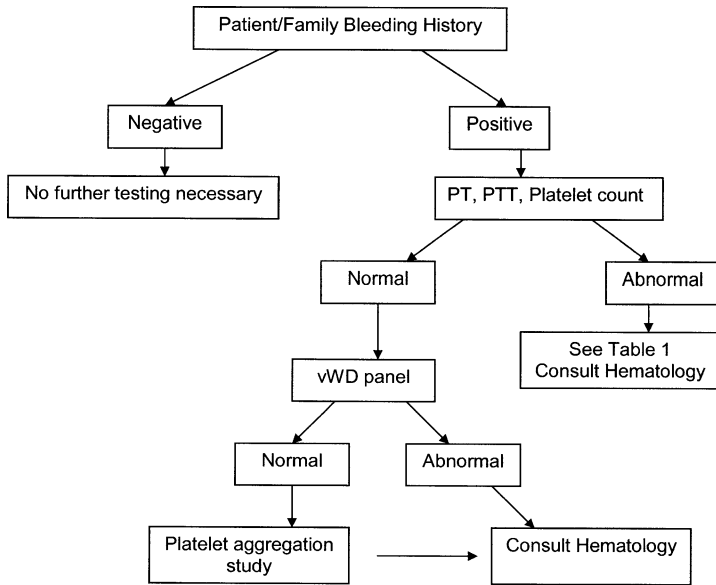


Figure 5.5. An algorithm to evaluate patients for possible platelet dysfunction. The patient and family history are important in deciding whether to pursue laboratory evaluation. If the history is positive, the screening tests of hemostasis (PT, PTT, platelet count) are used for further evaluation. The vWD panel consists of factor VIII activity, von Willebrand factor antigen, and ristocetin cofactor activity. (Abbreviations: vWD = von Willebrand's disease) Modified from Lehman CM, Blaylock RC, Alexander DP, et al. Discontinuation of the bleeding time test without detectable adverse clinical impact. *Clin Chem* 2001; 47: 1204–11.

Other laboratory tests to evaluate platelet function are discussed in Chapter 6, Testing for Inherited Bleeding Disorders, and Chapter 7, Testing for Acquired Platelet Disorders.

Interpretation of Hemostasis Screening Tests in Patients with Bleeding Disorders

Table 5.1 summarizes the differential diagnosis and recommended subsequent laboratory evaluation for patients with a positive bleeding history using the three hemostasis screening tests – PT, PTT, and platelet count.

Table 5.1. Interpretation of Hemostasis Screening Tests in Patients with Bleeding Disorders, and Recommended Subsequent Evaluation

| PT | PTT | Platelet Count | Differential Diagnosis | Recommended Tests |
|-----------|------------|-----------------------|---|--|
| ↑ | - | - | <i>Common</i> Acquired F VII deficiency (Liver disease, malnutrition, warfarin therapy) | Liver function tests Drug history |
| | | | <i>Rare</i> Some cases of DIC Factor VII inhibitor Dysfibrinogenemia | D-dimer Mixing test and F VII assay Functional/antigenic fibrinogen levels |
| | | | Inherited F VII deficiency Certain F X variants | F VII level F X level |
| - | ↑ | - | <i>Common</i> Heparin | Mixing test, TT assay, heparin assay |
| | | | Deficiency or inhibitor of Factors VIII, IX, XI | Mixing test, F VIII, IX, XI levels. If F VIII level is low, vWD panel |

(Continued)

Table 5.1. (Continued)

| | | | | |
|---|---|---|--|--|
| | <i>Rare</i> | | | Mixing test, LA panel, platelet aggregation F X assay |
| | Lupus anticoagulant with platelet dysfunction Certain F X variants | | | |
| | <i>Common</i> | | | |
| ↑ | Vitamin K deficiency Liver disease, warfarin, heparin | - | | Liver function tests Drug history |
| | <i>Rare</i> | | | |
| | DIC | | | D-dimer |
| | Dysfibrinogenemia | | | Functional/antigenic fibrinogen levels |
| | LA with hypoproteinemia Primary fibrinolysis | | | Mixing test, LA panel, prothrombin level |
| | Deficiency or inhibitor to factors X, V, prothrombin, or fibrinogen | | | Fibrinogen level, FDP levels (exclude DIC with D-dimer) |
| | <i>Common</i> | | | Mixing test, factor levels |
| | DIC | | | D-dimer |
| ↑ | Liver disease | ↓ | | Liver function tests |

(Continued)

Table 5.1. (Continued)

| PT | PTT | Platelet Count | Differential Diagnosis | Recommended Tests |
|----|-----|----------------|--|---|
| - | - | ↓ | <p><i>Rare</i> Heparin-induced thrombocytopenia</p> <p><i>Common</i> Decreased platelet production Increased platelet production Splenomegaly Hemodilution</p> <p><i>Rare</i> Certain inherited platelet disorders (Bernard-Soulier syndrome, Wiskott-Aldrich syndrome) Myeloproliferative disorders</p> <p><i>Common</i> vWD Acquired platelet dysfunction (uremia)</p> | <p>Drug history, Heparin-PF4 ELISA</p> <p>Bone marrow</p> <p>Physical exam History</p> <p>Blood smear evaluation</p> <p>Platelet aggregation study</p> <p>Bone marrow Cytogenetics, PCR tests (BCR-ABL, JAK-2, etc)</p> <p>vWD panel, multimeric analysis Renal function tests (platelet aggregation study)</p> |
| - | - | ↑ | | |
| - | - | - | | |

(Continued)

As mentioned above, these are screening tests, and normal test results do not exclude a bleeding disorder¹.

The category that may require the most extensive testing is the last one, in which all screening tests are normal. Although most patients will have common disorders such as mild von Willebrand disease or platelet dysfunction, some patients will require extensive testing before a diagnosis is made.

Key Points

- Screening tests of hemostasis (platelet count, PT, PTT) are useful in suggesting diagnostic possibilities in bleeding patients.
- The presence of normal values for platelet count, PT, and PTT assays does not exclude the possibility of a bleeding disorder. In fact, the most common bleeding disorders (von Willebrand's disease, platelet dysfunction) will usually present with this coagulation profile.
- Isolated, prolonged PT values suggest inherited or acquired (liver disease, malnutrition, warfarin therapy, DIC) factor VII deficiency.
- Isolated, prolonged PTT values in bleeding patients suggest either heparin therapy, factor deficiency or inhibitor to factors VIII, IX, or XI.
- Mixing studies should be performed to evaluate prolonged PTT values. Failure to correct to the normal reference interval with mixing suggests the presence of heparin, a lupus anticoagulant, or a specific factor antibody.
- Prolongations of both the PT and PTT suggest either inherited deficiency of a common pathway factor (uncommon), acquired multiple factor deficiency (DIC, vitamin K deficiency), or heparin or warfarin therapy.
- PT-derived fibrinogen methods are not recommended for routine use in the coagulation laboratory.
- D-dimer is a sensitive test for the diagnosis of DIC. However, many patients with low levels of plasma D-dimer do not have DIC, so assay-specific cutoffs must be established.
- Sensitive D-dimer assays, in combination with a pre-test probability assessment, have excellent negative predictive value for ruling out VTE. Manual agglutination assays cannot be used to rule out VTE.

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6

Testing for Inherited Bleeding Disorders

George M. Rodgers

Testing for Inherited Bleeding Disorders

Inherited bleeding disorders can be classified into three categories – those due to deficient platelet number or function, those due to deficient coagulation protein activity, and those due to vascular abnormality. The coagulation laboratory has a key role in evaluating patients in the first two categories, while disorders of vascular function are usually diagnosed by physical examination and certain genetic tests. Table 6.1 is a comprehensive listing of inherited platelet and coagulation disorders associated with bleeding, categorized by manner of genetic transmission. This chapter will focus on laboratory aspects of qualitative platelet disorders, von Willebrand's disease, and the coagulation disorders associated with bleeding. Unlike certain inherited thrombotic disorders that are associated with highly-conserved point mutations (e.g., factor V Leiden), the inherited bleeding disorders are polygenic and are best diagnosed with functional coagulation assays.

Qualitative Platelet Disorders

In these disorders, platelet dysfunction is associated with a normal platelet count. Normal platelet function results from several platelet activities. Platelet adhesion to subendothelium is the initial response to vascular injury, and is mediated by the plasma adhesion protein, von Willebrand's factor (vWF) and its platelet receptor, glycoprotein (GP) 1b. Platelet-platelet interaction (aggregation) is mediated by plasma fibrinogen and its platelet receptor, GP IIb-IIIa. Following platelet activation, secreted ADP amplifies platelet aggregation to

Table 6.1. Summary of Inherited Platelet and Coagulation Disorders Associated with Bleeding

| Disorders associated with: | Thrombocytopenia | Platelet dysfunction (Qualitative platelet disorders) | Coagulation disorders |
|----------------------------|--|--|---|
| | <i>Autosomal recessive</i> | <i>Autosomal recessive</i> | <i>X-linked</i> |
| | Bernard-Soulier syndrome | Bernard-Soulier syndrome | Factor VIII deficiency (hemophilia A) |
| | Congenital amegakaryocytic thrombocytopenia | Glanzmann's thrombasthenia, Storage pool disorder | Factor IX deficiency (hemophilia B) |
| | Gray platelet syndrome | Gray platelet syndrome | <i>Autosomal recessive</i> |
| | Thrombocytopenia with absent radius | Cyclooxygenase deficiency, Thromboxane synthase deficiency | Factor XI deficiency, Prothrombin deficiency, Factor V deficiency |
| | <i>Autosomal dominant</i> | Gray platelet syndrome | Factor VII deficiency |
| | May-Hegglin anomaly, Quebec platelet disorder | Scott's syndrome | Factor X deficiency, Factor XIII deficiency |
| | Epstein's syndrome, Fechtner's syndrome | | Afibrinogenemia, Hypofibrinogenemia |
| | Sebastian's syndrome | | α_2 Antiplasmin deficiency |
| | Montreal platelet syndrome, Hereditary macrothrombocytopenia | | Plasminogen activator inhibitor deficiency Combined deficiency of factors V/VIII |
| | <i>X-linked</i> | | <i>Autosomal dominant</i> |
| | Wiskott-Aldrich syndrome | | von Willebrand's disease |
| | X-linked thrombocytopenia GATA-1 mutation | | Dysfibrinogenemia |

generate the platelet thrombus¹. These hemostatic events are summarized in Figure 6.1. The platelet plug is consolidated (reinforced) by fibrin generated from the coagulation mechanism. All of these platelet functions (adhesion, aggregation, secretion) can be measured in the laboratory, and deficient platelet GP function can be inferred or

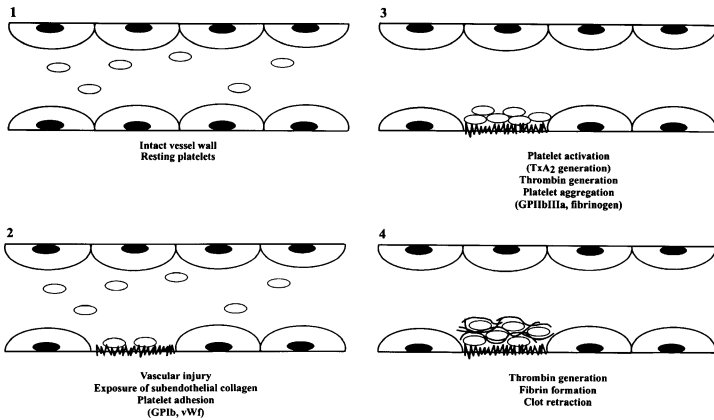


Figure 6.1. Sequence of hemostatic events following vascular injury. 1. In the resting state, platelets are non-activated and the endothelium is intact. 2. Following vascular injury, subendothelial components are exposed to induce platelet adhesion, mediated by vWF. 3. Collagen exposure activates platelets leading to platelet secretion, thromboxane A₂ generation and formation of thrombin. These events lead to platelet aggregation. 4. Thrombin generation results in cross-linked fibrin that reinforces the platelet plug. From Rodgers GM, Diagnosis of bleeding disorders. Kjeldsberg CR, et al., eds. In Practical Diagnosis of Hematologic Disorders. ASCP Press. Chicago, 2006, Fourth ed., p. 306, with permission.

quantitated. Screening tests for platelet function such as the bleeding time are not recommended and the utility of other screening tests such as the PFA-100 is unproven (see Chapter 7, Testing for Acquired Platelet Disorders and Chapter 3, Instrumentation for the Coagulation Laboratory).

Platelet Aggregation

The traditional assay for platelet function is platelet aggregation using platelet-rich plasma. Since countless prescription and over-the-counter medications and dietary supplements may impair platelet function, patients' medication lists must be reviewed by the coagulation laboratory prior to testing. Drugs containing aspirin must be discontinued 10–14 days prior to phlebotomy, while most other drugs should be stopped 4–7 days before phlebotomy. Citrated blood is obtained from the patient and a control subject (who also is taking no interfering medications), and platelet-rich plasma (PRP) is prepared by slow-speed

centrifugation. The blood sample should not be refrigerated. The PRP specimens are mixed with patient platelet-poor plasma to obtain a final platelet count $\sim 250,000/\mu\text{L}$.

Platelet aggregation is based on spectrophotometric monitoring of the turbid PRP sample after addition of various agonists (platelet activating stimuli). Agonist-induced platelet aggregation results in decreased turbidity and increased light transmittance. Figure 6.2 illustrates normal platelet aggregation patterns with PRP. The agonists used in platelet aggregation studies commonly include ADP (low-dose $0.5\text{--}2\mu\text{g/mL}$; high-dose $10\text{--}20\mu\text{g/mL}$), dilutions of a collagen suspension, arachidonic acid, and ristocetin (low-dose $0.5\mu\text{g/mL}$; high-dose $1\text{--}2\mu\text{g/mL}$). Some laboratories also use epinephrine as an agonist. This panel of agonists will test various platelet functions, including adhesion, aggregation, secretion and activation².

Zhou and Schmaier have published platelet aggregation procedures to standardize performance and interpretation of platelet function testing³. They ask patients to avoid all medications for 10 days prior to assay, to avoid coffee on the day of testing, and to fast for at least 4 hours before phlebotomy.

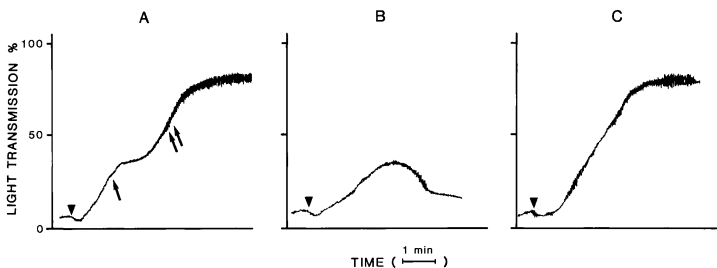


Figure 6.2. Normal platelet aggregation patterns. A. The primary and secondary waves of the platelet aggregation response. The primary wave (single arrow) results from agonist-induced platelet aggregation whereas the secondary wave (two arrows) results from platelet secretion of endogenous ADP and recruitment of additional activated platelets. B. Low-dose agonist-induced primary wave aggregation. The amount of stimulus is insufficient to fully activate the platelet and induce secretion of stored ADP. C. High-dose agonist-induced platelet aggregation in which the primary and secondary aggregation waves have merged into a single aggregation wave. Triangles indicate addition of the agonist. From Rodgers, GM. Platelet physiology and laboratory evaluation of platelet function. *Clin Obstet Gynec* 1999; 42: 349–59, with permission.

These authors recommend that each aggregation run last for 5–10 minutes, to allow observation of both primary and secondary aggregation waves, as well as the phenomenon of platelet disaggregation which may occur in some patients with platelet dysfunction.

Interpretation of platelet aggregation patterns can be done in two ways: calculation of the extent (percent) of aggregation, or determination of the initial rate of aggregation per minute. Zhou and Schmaier have published normal values for platelet aggregation results; most agonists result in maximal extent (percent) of aggregation of 70–80%³.

Figure 6.3 summarizes platelet aggregation profile results in patients with Glanzmann's thrombasthenia, Bernard-Soulier syndrome, and storage pool/secretion disorders (thrombopathies). Glanzmann's thrombasthenia platelets have deficient GPIIb-IIIa receptors, so that aggregation dependent on this receptor is decreased or absent. Consequently, aggregation induced by ADP, collagen, arachidonic acid (all agonists *except* ristocetin) is abnormal. In contrast, patients with Bernard-Soulier syndrome have platelets deficient in GPIb receptors, so that aggregation dependent on this receptor is decreased or absent. Since ristocetin is the only agonist dependent on normal GPIb function, Bernard-Soulier

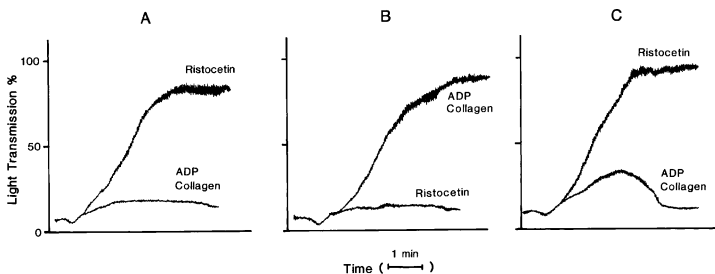


Figure 6.3. Platelet aggregation profiles in vWD and the inherited qualitative platelet disorders. Typical responses to ADP, collagen, and ristocetin are shown. Three patterns of aggregation are shown: A. Glanzmann's thrombasthenia. B. Bernard-Soulier syndrome and vWD. Both diseases give similar aggregation results with decreased or absent response to ristocetin. The two diseases are distinguished by Bernard-Soulier patients having giant platelet morphology with thrombocytopenia and vWD patients having normal platelet morphology and decreased ristocetin cofactor activity. C. Thrombopathies (secretion defects and abnormalities in prostaglandin synthesis) From Rodgers GM. Platelet physiology and laboratory evaluation of platelet function. Clin Obstet Gynec 1999; 42: 349–59, with permission.

syndrome platelets have normal aggregation responses to all agonists but ristocetin².

Platelets from patients with storage pool disorder or secretion defects will exhibit normal primary aggregation responses to all agonists, but since the defect is in platelet activation and/or secretion, secondary aggregation is defective to all agonists except ristocetin².

Platelet aggregation results can be reported quantitatively (percentage change of transmittance) or qualitatively (normal or abnormal), with a comment as to the type of abnormality and possible diagnosis. It is important to remember that numerous common medications can induce marked platelet dysfunction, and this potential confounding issue should be considered when interpreting an abnormal platelet aggregation study.

Table 6.2 summarizes the platelet aggregation profiles seen with the inherited qualitative platelet disorders. Although von Willebrand's disease (vWD) is best diagnosed with other tests, platelet aggregation can be used to diagnose vWD. Deficiency of vWF results in a profile where all agonist responses are normal, except for ristocetin, a pattern similar to that seen in Bernard-Soulier syndrome. In Bernard-Soulier syndrome, the defect is in the vWF receptor (GP1b); in vWD, the defect is absent vWF. The two disorders can be distinguished by assay of vWF activity with the ristocetin cofactor assay; levels are normal in Bernard Soulier syndrome and decreased in vWD (Table 6.2).

The key clinical distinction in evaluating patients for qualitative platelet disorders is to differentiate bleeding from a platelet disorder vs. vWD. Both disorders have similar clinical features, but are treated differently – vWD patients should receive a vWF replacement product, while patients with platelet dysfunction should receive platelet transfusions for significant bleeding.

Whole blood aggregation methods are also available that are less laborious than the PRP method. Certain whole blood assays also report platelet secretion data in addition to aggregation data, permitting direct identification of platelet dysfunction due to secretion defects.

von Willebrand's Disease

vWD is the most common inherited bleeding disorder, affecting ~ 1% of the population. The disorder is characterized by deficiency or qualitative abnormality of vWF, the plasma adhesion protein necessary for primary hemostasis. vWF circulates in the blood in a complex with factor

Table 6.2. Platelet Aggregation Profiles in the Inherited Qualitative Platelet Disorders and von Willebrand's Disease

| Disorders | <i>Aggregation response</i> | | | | Comments |
|----------------------------|-----------------------------|---------|-----------|----------|---|
| | ADP | Primary | Secondary | Collagen | |
| Bernard-Soulier syndrome | N | N | N | A | Giant platelets and thrombocytopenia are present. vWF activity assay is normal. |
| Glanzmann's thrombasthenia | A | A | A | N | Normal platelet morphology |
| Storage pool disorder | N | A | A | N | Platelet morphology by light microscopy is usually normal. Electron microscopy is abnormal. |
| Secretion defect | N | A | A | N | Platelet morphology by light and electron microscopy is normal. |
| vWD | N | N | N | A | Platelet morphology is normal. vWF activity assay is decreased. |

N = normal, A = abnormal, vWD = von Willebrand's disease

Source: Rodgers, GM. Qualitative platelet disorders and von Willebrand's disease. In Kjeldsberg C, ed. Practical Diagnosis of Hematologic Disorders. Chicago, IL: ASCP Press, 2006; p. 331.

VIII, the plasma protein deficient in hemophilia A. *In vivo*, vascular injury results in exposure of subendothelial components that promote vWF-mediated platelet adhesion. This event is dependent primarily

on the highest molecular weight component (multimers) of vWF. *In vitro* assays of platelet function using ristocetin mimic *in vivo* platelet adhesion; such assays (ristocetin-induced platelet aggregation, ristocetin cofactor activity) measure vWF function².

Assays for vWF

Several assays for vWF are available to assess patients with bleeding disorders for vWD. Most commonly, a vWD “panel” is obtained that measures antigenic and functional properties of the factor VIII – vWF complex. Panel components include vWF antigen, a quantitative assay of vWF, as measured by ELISA, Laurell immunoelectrophoresis, or immunoturbidimetric assay; vWF activity, as measured by a test called ristocetin cofactor activity; and factor VIII activity, as measured by coagulation or chromogenic substrate assay. Of these three panel components, ristocetin cofactor activity is frequently the most abnormal, but assaying all 3 components is recommended. If an abnormal vWD panel result is obtained, the patient’s disorder is further classified by multimeric analysis, in which the patient’s vWF is analyzed by immunoblot and characterized as type 1 (quantitative mild to moderate deficiency), type 2 [qualitative abnormality in which high-molecular-weight multimers are absent (type 2B), or high- and intermediate-molecular-weight multimers are absent (type 2A)], or type 3 (absence of vWF). This classification is clinically important since patients with type 1 vWD, but not types 2 or 3 vWD may respond to desmopressin, a non-transfusion therapy that can increase plasma vWF levels. Another assay that may be useful in diagnosing patients with vWD when the standard panel results are normal is ristocetin-induced platelet aggregation². vWF activity may also be measured by a collagen binding assay.

vWF Immunoassay

Historically, Laurell rocket immunoelectrophoresis has been the standard method to quantitate vWF antigen. This assay uses a precipitating polyclonal antibody to vWF in an electroimmunodiffusion technique, but this method is very labor-intensive. Current methods are based on ELISA techniques, which can be automated. Values less than 50% of normal are usually diagnostic of vWD. vWF is an acute-phase response protein, and can be increased by stress, trauma,

infection, inflammation, pregnancy, and female hormones. Consequently, a normal value does not exclude vWD, and repeat testing may be needed to establish the diagnosis².

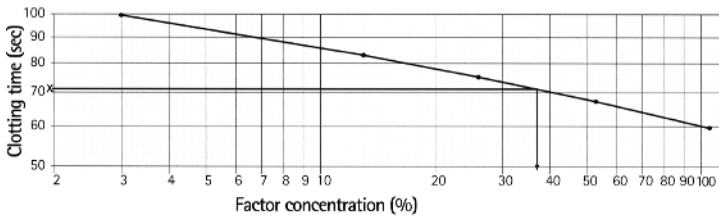
Ristocetin Cofactor Activity (vWF Activity) Assay

This is a functional assay of vWF that traditionally measures the ability of patient plasma to aggregate normal platelets in the presence of ristocetin, compared to normal plasma. This is a laborious technique that requires normal platelets, and is being replaced by ELISA methods in which a monoclonal antibody against a functional vWF epitope is used. Values less than 50% of normal are usually diagnostic of vWD. As with the vWF antigen assay, acute-phase events, pregnancy, and female hormone use may confound the diagnosis of vWD².

Factor VIII Activity Assay

This assay is useful in evaluating patients for factor VIII deficiency due to hemophilia A or vWD, and in monitoring response to therapy. The ability of dilutions of patient plasma deficient in factor VIII to correct the prolonged partial thromboplastin time (PTT) of commercial factor VIII-deficient plasma is compared to normal plasma. For example, if a 1:40 dilution of normal plasma shortens the clotting time of deficient plasma to the same extent as a 1:10 dilution of patient plasma, the patient plasma has 25% of normal factor VIII activity⁴. Figure 6.4 depicts a typical standard curve for this assay.

Dilutions of reference plasma (1:10, 1:20, 1:40, 1:80, 1:160, 1:320) and patient plasma (1:10, 1:20) are made in buffer (Owren's or imidazole). One part of diluted plasma is added to one part of factor VIII-deficient plasma and a PTT is performed. The 1:10 dilution of reference plasma is arbitrarily defined as 100% activity, the 1:20 dilution as 50%, etc. A line of best fit is obtained (Figure 6.4) when clotting times are plotted against per cent factor VIII activity. The factor VIII activities that give the same clotting times as the 1:10 and 1:20 patient plasma dilutions are extrapolated from the graph. The 1:10 dilution value is the patient's actual factor VIII level; the 1:20 dilution value must be multiplied by two, and the mean of the two values is reported. Values less than 50% of normal suggest factor VIII deficiency due to hemophilia A or vWD. As with vWF, factor VIII is an acute-phase response protein, and normal factor VIII levels may



Factor Assay Standard Curve Data

| Factor Concentration (%) | Normal plasma dilution | Clotting time (sec) |
|--------------------------|------------------------|---------------------|
| 105% | 1:10 | 59.1 |
| 53% | 1:20 | 67.2 |
| 26% | 1:40 | 75.2 |
| 13% | 1:80 | 82.7 |
| 3% | 1:320 | 97.4 |

Figure 6.4. A typical factor assay standard curve. Dilutions of normal pooled plasma are made (1:10–1:320), and clotting times are measured. Results of clotting times vs. factor concentration are plotted on log-log paper, and the best-fit line is drawn. Patient plasma is diluted 1:10 and 1:20, and clotting times are measured. As indicated in the figure, the 1:10 dilution of patient plasma had a clotting time of 71 sec (“x” on the Y-axis). Interpolation from the standard curve yields a factor value of 37%. These curves can be prepared manually, or fully-automated coagulation analyzers can be used to prepare dilutions, add reagents, measure clotting times, plot standard curves, and calculate patient results. From Rodgers GM. Inherited coagulation disorders. In Kjeldsberg CR, ed. Practical Diagnosis of Hematologic Disorders. ASCP Press. Chicago, 2006, Fourth ed., p. 351, with permission.

not exclude mild deficiency. A difficult phlebotomy (activated plasma sample) may also artifactually increase the factor VIII level⁴.

vWF multimeric Analysis

This is an immunoblot that is useful to distinguish quantitative vs. qualitative types of vWD. Patient and normal plasma are electrophoresed using dilute agarose (0.7–1%). The separated plasma proteins are transferred to nitrocellulose paper, incubated with an antibody to vWF, followed by immunodetection using the avidin-biotin-peroxidase technique. The resulting immunoblot (Figure 6.5) illustrates the full range of multimers in normal plasma and in plasma from patients with type 1 vWD, as well as absence of high-and/or intermediate-molecular-weight-multimers in

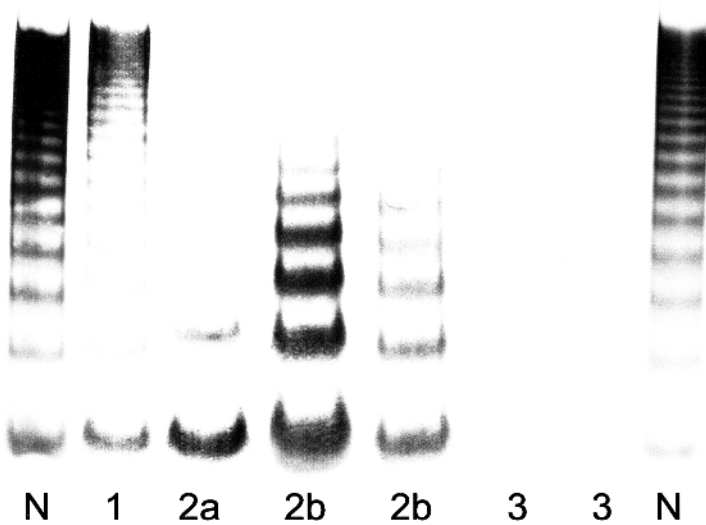


Figure 6.5. Multimeric analysis of vWF. Plasma was obtained from a normal subject (N), and from patients with various types of vWD (1,2A, 2B, 3). Plasma was electrophoresed in an agarose gel, then vWF was identified using an immunoperoxidase method. The dark bands at the top of the gel (N) represent the high-molecular-weight multimers most important in platelet adhesion. Note the generalized decrease in band intensity in type 1, the loss of intermediate and high-molecular-weight multimers in type 2A, the loss of only high-molecular-weight multimers in type 2B, and the virtual absence of all multimers in type 3. From Rodgers GM, Qualitative platelet disorders and von Willebrand's disease. In Kjeldsberg CR, ed. In Practical Diagnosis of Hematologic Disorders. ASCP Press. Chicago, 2006, Fourth ed., p. 339, with permission.

patients with type 2 vWD, or complete absence of multimers as in patients with type 3 vWD².

Appropriate Test Ordering for Platelet-type Bleeding Disorders

Patients with mucocutaneous bleeding, a normal platelet count, and a family history of excessive bleeding are appropriate candidates for testing of platelet-type bleeding disorders which include vWD and the qualitative platelet disorders. Given the limitations of screening tests such as the bleeding time, it would be reasonable to initially evaluate patients for vWD since this disorder is much more common than the

disorders of platelet function. An exception to this recommendation would be patients with abnormal platelet morphology on blood smear examination (giant platelets, gray platelets, etc.) that suggests a qualitative disorder for which platelet aggregation studies would be the appropriate initial test.

If the vWD panel is normal, platelet aggregation testing using a complete panel of agonists (collagen, ADP, arachidonic acid, ristocetin, all in various concentrations) is appropriate. It is important to ensure that all antiplatelet medications are discontinued prior to testing. Some patients with vWD will have normal panel results, but may be detected with ristocetin-induced platelet aggregation. If patients have normal results for both the vWD panel and platelet aggregation test, and the likelihood of the platelet-type bleeding disorder is thought to be high, repeat vWD testing is indicated, since factor VIII and vWF are acute-phase reactants, and many patients with mild vWD will have normal studies. For women who are on hormone therapy or pregnant, and in whom normal vWD panel results will not exclude the diagnosis, testing of symptomatic family members may be useful².

One genetic modifier of plasma vWF concentrations is the patient's ABO blood group; individuals with type O have ~ 25% lower vWF concentrations than do those with a non-O blood type⁵. Correct interpretation of borderline-low vWF panel results should take into consideration the patient's blood type.

The Platelet Function Analyzer (PFA)-100®

The PFA-100® device is now available as a screening test for disorders of platelet function. An international consensus panel has recently evaluated the literature on this device in terms of its clinical utility⁶. These authors concluded that this assay should be considered optional in the evaluation of platelet function, based on a paucity of outcome data that has been published with the PFA-100®⁶.

Inherited Coagulation Disorders

The qualitative platelet disorders and vWD described above result in defective primary hemostasis – formation of an ineffective platelet thrombus. The coagulation mechanism (secondary hemostasis) is necessary to generate fibrin to reinforce the platelet plug. The inherited coagulation disorders have in common deficient fibrin formation that

frequently leads to delayed bleeding. Assays are based on the factor VIII activity method described above in which the ability of patient plasma to shorten prolonged clotting times (PT-or PTT-based) of commercially-deficient plasma is compared to normal plasma. For example, the factor IX activity assay uses factor IX-deficient plasma in a PTT-based assay. The factor VII activity assay uses factor VII-deficient plasma in a prothrombin time (PT)-based assay. Factors VIII, IX, XI, XII, prekallikrein, and high-molecular weight kininogen are assayed using PTT-based methods, while factors VII, X, V, and prothrombin are assayed using PT-based methods. Diagnosis of a specific coagulation-type bleeding disorder depends on specific coagulation factor assay results. Table 6.1 illustrates how abnormal PT and/or PTT results lead to recommended specific factor assays. Table 6.3 summarizes the effects on PT and PTT assays of coagulation factor deficiencies, classified further as to whether or not excessive clinical bleeding is present.

Table 6.3. Results of PT and PTT Assays in Coagulation Factor Deficiency Disorders

| Factor deficiency | PT | PTT | Excessive bleeding? |
|---------------------------------|-----------|------------|----------------------------|
| XII | NI | ↑ | No |
| Prekallikrein | NI | ↑ | No |
| High-molecular-weight kininogen | NI | ↑ | No |
| XI | NI | ↑ | Yes |
| IX | NI | ↑ | Yes |
| VIII | NI | ↑ | Yes |
| VII | ↑ | NI | Yes |
| X | ↑ | ↑ | Yes |
| V | ↑ | ↑ | Yes |
| Prothrombin | ↑ | ↑ | Yes |
| Fibrinogen | ↑ | ↑ | Yes |
| XIII | NI | NI | Yes |

PT and PTT test results in patients with coagulation factor deficiency disorders are shown and classified as to whether excessive bleeding is seen with the disorder. Patients with mild bleeding disorders may have normal PT and PTT results, and identification of these patients may depend on PT and PTT reagent sensitivities. Although deficiency of factor XII, prekallikrein, and high-molecular weight kininogen is associated with increased PTT values, there is no excessive bleeding seen in these disorders.

Abbreviations: NI = normal; ↑ = increased; PT = prothrombin time; PTT = partial thromboplastin time.

Table 6.4 summarizes pediatric reference intervals for coagulation tests^{7,8}. Note that there is age-dependence for many analytes, including the PT, prothrombin, factors VII, VIII, IX, and X, alpha₂-antiplasmin, antithrombin and protein C.

Factor XIII (Clot Stability) Assay

The assay principle for factor XIII activity differs from other factor assays described above. The screening assay for factor XIII activity is performed by recalcifying patient and control plasma to produce a

Table 6.4. Summary of Pediatric Reference Intervals

| Age | 7-9 | 10-11 | 12-13 | 14-15 | 16-17 | Adult |
|--|-----------|-----------|-----------|-----------|-----------|-----------|
| N | 245 | 164 | 164 | 164 | 150 | 120 |
| PT (sec) | 13.0-15.4 | 13.0-15.6 | 13.0-15.2 | 12.8-15.4 | 12.6-15.7 | 12.3-14.4 |
| PTT (sec) | 27-38 | 27-38 | 27-39 | 26-36 | 26-35 | 26-38 |
| Factor II, % | 78-125 | 78-120 | 72-123 | 75-135 | 77-130 | 86-150 |
| Factor V, % | 69-132 | 66-136 | 66-135 | 61-129 | 65-131 | 62-140 |
| Factor VII, % | 67-145 | 71-163 | 78-160 | 74-180 | 63-163 | 80-181 |
| Factor VIII, % | 76-199 | 80-209 | 72-198 | 69-237 | 63-221 | 56-191 |
| Factor IX, % | 70-133 | 72-149 | 73-152 | 80-161 | 86-176 | 78-184 |
| Factor X, % | 74-130 | 70-134 | 69-133 | 63-146 | 74-146 | 81-157 |
| Factor XI, % | 70-138 | 66-137 | 68-138 | 57-129 | 65-159 | 56-153 |
| RCF, % | 52-176 | 60-195 | 50-184 | 50-203 | 49-204 | 51-215 |
| vWF Ag, % | 62-180 | 63-189 | 60-189 | 57-199 | 50-205 | 52-214 |
| Fibrinogen, % | 198-413 | 197-410 | 215-378 | 204-392 | 208-438 | 211-441 |
| Alpha ₂ - antiplasmin, % | 88-147 | 90-144 | 87-142 | 83-136 | 77-134 | 82-133 |
| AT, % | 90-135 | 90-134 | 90-132 | 90-131 | 87-131 | 76-128 |
| Plasminogen, % | 76-116 | 74-117 | 66-114 | 71-124 | 75-132 | 71-144 |
| Protein C, % | 70-142 | 68-143 | 66-162 | 69-170 | 70-171 | 83-168 |
| Protein S, % | 66-140 | 69-139 | 72-139 | 68-145 | 77-167 | 66-143 |
| Male | | | | | | |
| Protein S, % | 62-151 | 65-142 | 70-140 | 55-145 | 51-147 | 57-131 |
| Female | | | | | | |

PT and PTT results are expressed in seconds. All other analytes are expressed as percent of normal.

Abbreviations: PT = prothrombin time, PTT = partial thromboplastin time, RCF = ristocetin cofactor activity, vWF Ag = von Willebrand factor antigen, AT = antithrombin

Source: Flanders MM, et al. Pediatric reference intervals for seven common coagulation assays. Clin Chem 2005; 51: 1738-1742 and Flanders MM, et al. Pediatric reference intervals for uncommon bleeding and thrombotic disorders. J Pediatr 2006; 149: 275-277.

fibrin clot. Control and patient clots are suspended in solutions of strong denaturing agents (urea, trichloroacetic or monochloroacetic acid) for 24 hours. Clots that have been appropriately crosslinked by factor XIII_a can withstand such conditions and no clot lysis is seen. Clots formed in the absence of factor XIII dissolve in the presence of denaturing agents⁴. Quantitative factor XIII assays can be performed to confirm the abnormal screening assay results. The quantitative assays are not routinely performed in coagulation laboratories; they measure transglutaminase activity.

Fibrinogen Activity Assay

This assay utilizes a thrombin time-based assay. Serial dilutions of a fibrinogen reference standard are prepared and a calibration curve is done. The clotting time is inversely proportional to the amount of fibrinogen in the sample. A typical reference interval for fibrinogen in adults is 150–350 mg/dL⁹. Low fibrinogen levels are observed in inherited afibrinogenemia or hypofibrinogenemia, dysfibrinogenemia, primary fibrinolysis, and disseminated intravascular coagulation. Diagnosis of dysfibrinogenemia requires demonstration of a normal fibrinogen antigen level with a decreased functional level.

Alpha₂-antiplasmin Assay

This assay measures functional activity of an enzyme associated with an uncommon bleeding disorder. This disorder is usually considered in patients with coagulation-type bleeding in whom other factor deficiencies have been excluded. Plasmin is added to the patient's plasma and residual plasmin activity is measured by a fluorescent or chromogenic substrate assay. Alpha₂-antiplasmin levels can also be measured using immunologic assays⁴.

Assays for Factor VIII Antibodies (Inhibitor Assays)

Up to 30% of patients with inherited factor VIII deficiency will develop antibodies to factor VIII. Identification of these antibodies is important since they are associated with increased bleeding, and will usually require specific treatment. Patients may acquire antibodies to factor VIII in the absence of inherited hemophilia in association with connective tissue or malignant diseases, or in certain non-malignant settings such as pregnancy.

The principle of these assays is based on the mixing study (inhibitor screen) that was discussed in Chapter 5, Hemostasis Screening Assays. Normal plasma is incubated with patient plasma suspected of having a factor VIII antibody, and residual factor VIII activity of the mixed sample is measured. A screening assay for factor VIII antibodies is less labor intensive than the formal Bethesda assay that quantitates the factor VIII antibody level.

In the screening assay, one part of patient plasma is mixed with an equal part of normal pooled plasma that contains 100% factor VIII activity. After a 2 hour incubation at 37°C, the mixture is diluted in saline (1:5) and assayed for factor VIII. If this assay result is 35% or higher, no antibody is present, or if present, is too weak to be clinically important^{4,9}.

Patient samples with factor VIII results < 30–35% should be further evaluated with a Bethesda assay to quantitate the factor VIII antibody level. Serial dilutions of patient plasma in buffer (1:2, 1:4, 1:8, etc.) are incubated with equal volumes of normal plasma for 2 hours at 37°C. The residual factor VIII activity of the incubation mixtures is determined. The inhibitor level (in Bethesda units) is the reciprocal of the dilution of the patient plasma that results in 50% inhibition of factor VIII activity. An antibody level < 5 Bethesda units is labeled as a low-level inhibitor, while a level ≥ 5 Bethesda units is considered a high-level inhibitor⁴.

Laboratories that offer Bethesda assays should also provide similar assays against porcine factor VIII. These assays are similar to the above-mentioned method except that porcine factor VIII is substituted for human plasma. The anti-porcine factor VIII level provides useful clinical information in that if the patient has high antibody levels towards human factor VIII, but low levels against porcine factor VIII, porcine factor VIII may be a treatment option.

Figure 6.6 illustrates a factor VIII Bethesda assay calibration curve that converts residual factor VIII activity measured by incubating patient and reference plasma to Bethesda inhibitor units.

Key Points

- The gold standard assays for evaluating platelet function are platelet aggregation methods using either whole blood or platelet-rich plasma. Discontinuing patients' anti-platelet medications prior to testing is essential to appropriate interpretation of platelet aggregation results.

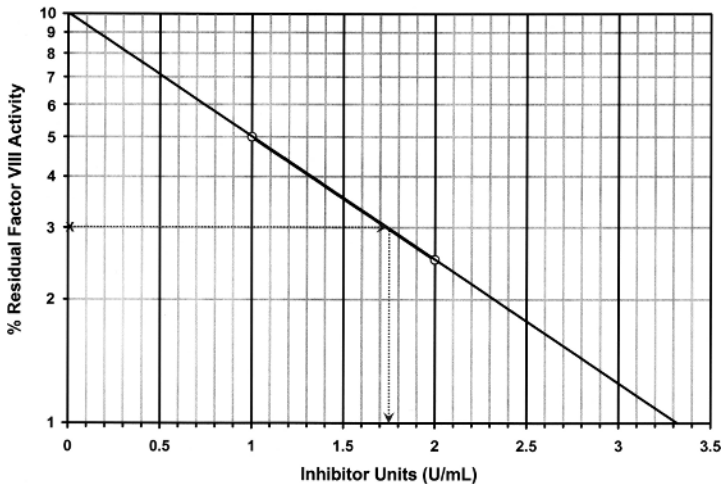


Figure 6.6. A factor VIII Bethesda assay calibration curve. In this example, the residual factor VIII activity measured after incubating patient and reference plasma for 2 hours was 30% (small “x” on the Y-axis). Interpolation from the graph indicates that a factor VIII antibody is present at a level of 1.75 Bethesda units. If the patient sample was diluted prior to incubation with reference plasma, 1.75 would be multiplied by the dilution factor to obtain the final result.

- The most common inherited bleeding disorder is von Willebrand’s disease (vWD); diagnosis is best achieved with the vWD panel (factor VIII activity, von Willebrand factor antigen, ristocetin cofactor activity), with multimeric analysis used to classify the vWD type.
- With the exception of vWD, factor VIII and factor IX deficiencies, other inherited bleeding disorders are very uncommon. It may be most appropriate for reference laboratories to perform these esoteric tests.
- Pediatric reference intervals are now available and should be used to correctly classify younger patients with possible bleeding disorders.

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7

Testing for Acquired Platelet Disorders

Christopher M. Lehman

Testing for Acquired Platelet Disorders

Acquired platelet disorders can be broadly categorized into disorders of platelet number (Table 7.1) and disorders of platelet function (Table 7.2), though there can be overlap (e.g., myeloproliferative disorders). The hematology laboratory traditionally assesses platelet number using automated hematology analyzers, with targeted confirmation by manual counting. Platelet antibody assays may be performed by the immunology and/or coagulation laboratories, or by the transfusion service. Platelet function testing has traditionally been performed by special coagulation laboratories using platelet-rich plasma or whole blood platelet aggregation. However, new platelet function analyzers are being marketed as potential substitutes for traditional platelet aggregometry (see Chapter 3, Instrumentation for the Coagulation Laboratory).

Platelet Counting Methods

Reference Method

Until relatively recently, the gold standard for platelet counting was the phase contrast manual count. Patient whole blood is diluted and loaded onto a hemocytometer and platelets are counted in a defined volume of sample. The raw counts are converted into a platelet concentration by a calculation that takes into account the dilution factor and the number and volume of the squares counted¹. While this method is still utilized in the laboratory as a backup for automated methods, it suffers from poor precision and accuracy due to the small sample size that is evaluated. This created difficulties for manufacturers of automated hematology analyzers, since the reference method was less precise and accurate

Table 7.1. Etiologies of Acquired Disorders of Platelet Number

| |
|--|
| I. Causes of Thrombocytopenia |
| Decreased Production |
| Marrow Aplasia |
| Marrow Infiltration |
| Myelodysplasia |
| Megaloblastic Anemia |
| Drug Effects |
| Increased Peripheral Destruction |
| Non-immunologic |
| Disseminated Intravascular Coagulation |
| Heparin-induced Thrombocytopenia Type I |
| Artificial Cardiac Valves |
| Extra Corporeal Membrane Oxygenation |
| Atrio-ventricular Fistulae |
| Left Ventricular Assist Devices |
| Hypersplenism |
| Liver Failure |
| Eclampsia/Preeclampsia |
| Infection |
| Thrombotic Thrombocytopenic Purpura |
| Hemolytic Uremic Syndrome |
| HELLP Syndrome |
| Immunologic |
| Autoimmune Thrombocytopenic Purpura |
| Sepsis |
| Neonatal Alloimmune Thrombocytopenia |
| Post-transfusion Purpura |
| Drug-Induced Thrombocytopenia |
| Heparin Induced Thrombocytopenia Type II |
| Systemic Lupus Erythematosus |
| Lymphoproliferative disorders |
| Platelet Transfusion Refractoriness |
| Hemodilution |
| Pseudothrombocytopenia |
| Anticoagulant-mediated antibodies |
| II. Causes of Thrombocytosis |
| Reactive |
| Infection |
| Chronic Inflammation |
| Iron Deficiency |
| Myeloproliferative Disorders |

Table 7.2. Etiologies of Acquired Disorders of Platelet Function

I. Causes of Acquired Platelet Dysfunction

Physiologic

Uremia

Myeloproliferative Disorders

Hyperfibrinolysis

Von Willebrand Disease (indirect cause of platelet dysfunction)

Iatrogenic

Cardiopulmonary Bypass

Drug-induced

Non-steroidal Anti-inflammatory Drugs

Aspirin

Unfractionated and Low Molecular Weight Heparins

Thienopyridines

Clopidogrel

Platelet Glycoprotein IIb/IIIa Inhibitors

Other Drugs

than the methods that they were attempting to validate. This problem was solved in 2001 when the International Council for Standardization in Haematology adopted a cytofluorometric-based platelet counting standard that labels platelets with two platelet-specific antibodies: anti-CD 61 and anti-CD 41². The ratio of platelet to red blood cell (RBC) events counted by cytofluorometry is multiplied by the RBC count produced by an automated hematology analyzer to derive the platelet concentration (counts/ μL).

Automated Platelet Counters

Automated hematology analyzers use electromagnetic fields to count blood cells, including platelets. The oldest method, the impedance method, uses static electrical fields, and volume-dependent changes in conductivity produced by blood cells as they pass through an aperture to identify and size platelets. Different field frequencies can be overlaid to optimize an analyzer's ability to differentiate cell types. The impedance method is susceptible to interference from non-cellular (e.g., cryoglobulins) and cellular particulate debris (e.g., RBC and white blood cell fragments), as well as microcytic red blood cells and RBC ghosts that have a volume distribution that overlaps that of platelets. Overlap of the platelet distribution with microcytic RBCs generally limits the

ability of impedance counters to detect large platelets. Sophisticated computer algorithms have been designed to minimize interferences with the impedance platelet count, including mathematical extrapolation to include counts of platelets larger than the actual measurement range³.

A second approach to platelet counting uses light or fluorescence scatter detected at different angles to identify, count and size platelets³. Information obtained from optical detection methods provides better differentiation of platelets from microcytic RBCs allowing direct counting of large platelets⁴. Therefore, optical counting has been shown to be more accurate than impedance counting when specimen selection is weighted towards conditions that produce microcytosis (e.g., immune-mediated hemolysis), red cell fragmentation (e.g., thrombotic thrombocytopenic purpura) and large platelets (e.g., autoimmune thrombocytopenic purpura)⁵. However, comparisons involving random specimen selection, or thrombocytopenic specimens collected from oncology patients demonstrate excellent correlation of both optical and impedance methods with counts produced by the international cytofluorometric reference method^{5,6}. Of particular importance, current impedance and optical methods are sufficiently accurate to guide transfusion therapy at platelet count triggers of 15–20,000/ μL ^{6,7}. Transfusion triggers of 5–10,000/ μL are probably beyond the capability of all current platelet-counting methods, with the possible exception of an immunologic assay^{6,8}.

Pre-Analytic Considerations

Pre-analytic factors must be considered when evaluating a low platelet count. Specimen clotting and platelet clumping due to insufficient mixing or platelet activation of the sample during phlebotomy must be ruled out by microscopy on any patient presenting with apparent thrombocytopenia by automated analysis. Hematology analyzers flag platelet clumps with variable sensitivity and specificity. A new specimen should be collected in the case of clotting or platelet clumping. Use of protocols to disaggregate platelet clumps in an attempt to produce an accurate platelet count should be used cautiously, and only after appropriate validation⁹. Pseudothrombocytopenia – *in vitro* platelet clumping due to anticoagulant-dependent (primarily EDTA), platelet autoantibodies – can also be ruled out by microscopy. Interference due to pseudothrombocytopenia can usually be overcome by collecting specimens to be analyzed for platelet counts in a sodium citrate or ACD

tube. Counts from citrate or ACD tubes should be corrected for the dilutional effect of the liquid anticoagulant (10% dilution for sodium citrate and 20% for ACD).

Platelet Antibody Testing

Human platelets express a variety of platelet-specific antigens, as well as class I human leukocyte antigens (HLA) that are targets for alloimmunization. In addition, neo-antigens may be formed as a result of interactions between native platelet antigens and therapeutic drugs (e.g., heparin). Antibodies may be formed against these antigenic structures, as a result of infusion of therapeutic drugs, or exposure to non-self, platelet-specific antigens through pregnancy, transfusion of platelet-containing blood components or hematopoietic allo-transplantation (active immunization), or immunoglobulin administration (passive immunization). Finally, autoantibodies directed against glycoproteins common to all normal platelets may be produced after an acute infection, as a result of an autoimmune disease, or without a clear precipitating cause (idiopathic).

Immune-Mediated Thrombocytopenia

Immune thrombocytopenia results when antibody bound to a patient's platelets (allo- or auto-antibody) interacts with the reticulo-endothelial system and platelets are sequestered and destroyed (Table 7.3). Alloimmune thrombocytopenia may occur as a result of active transfer by the placenta of platelet-specific (and possibly HLA) IgG antibodies into the fetal circulation and subsequent clearance of fetal platelets (neonatal alloimmune thrombocytopenia or NAIT); or as a result of binding of platelet-specific, alloantibody (produced secondary to prior pregnancy or platelet transfusion), to transfused platelets resulting in destruction of both transfused and native, bystander platelets (post-transfusion purpura or PTP). Alloimmune thrombocytopenia may also occur as a result of platelet-specific antibody production by passenger lymphocytes carried in a solid organ transplant, or after hematopoietic allo-transplantation when the recipient immune system makes antibodies against the donor's platelet antigens. On the other hand, with the exception of rare cases of HLA-induced NAIT¹⁰, production of HLA antibodies doesn't produce thrombocytopenia. Instead, HLA antibodies

Table 7.3. Platelet Antibody-mediated Thrombocytopenic Disorders

| |
|---|
| I. Alloimmune Etiologies |
| Neonatal Alloimmune Thrombocytopenia |
| Post-transfusion Purpura |
| Drug-induced Thrombocytopenia |
| Heparin-induced Thrombocytopenia |
| Infusion of Platelet Alloantibodies (e.g., immunoglobulin products) |
| Alloantibodies Produced by Transplanted Lymphocytes |
| II. Autoimmune Etiologies |
| Autoimmune Thrombocytopenic Purpura |
| Autoimmune Diseases (e.g., systemic lupus erythematosus) |

pre-dispose platelet transfusion-dependent patients to platelet transfusion refractoriness – poor increments in platelet count after transfusion of non-HLA matched platelet products.

Autoimmune thrombocytopenic purpura (AITP) results when patients produce antibodies against their own platelets resulting in rapid clearance of both the patient's and transfused platelets due to the fact that the antibodies are directed against platelet glycoproteins common to all normal platelets (GPIIb/IIIa and/or GPIb). A patient's own platelets may also be destroyed as a result of drug-induced, immune thrombocytopenia. This may result from an immune response against neoantigens formed as a result of drug-platelet interaction, or as a result of the production of non-drug dependent autoantibodies.

HLA Antibody Tests

Circulating HLA antibodies are routinely detected and identified using indirect methods including antibody-mediated cytotoxicity assays, cytofluorometric assays using manufactured microbeads coupled to purified HLA glycoproteins, or ELISA using immobilized, purified HLA glycoproteins bound to microtiter wells.

Platelet Antibody Tests

Testing for platelet antibodies has undergone significant evolution over the last five decades¹¹. Phase I tests were designed to detect the effect of serum/plasma platelet antibodies on control platelets after incubation. Test endpoints included platelet aggregation, platelet

activation, platelet lysis, and release of platelet granule contents. These tests were primarily used to detect autoantibodies, but had poor sensitivity and specificity, and survive today as the serotonin release assay used for diagnosis of heparin induced antibodies, and aggregation methods for detecting the effects of cardiopulmonary bypass, platelet inhibitory drugs, and heparin dependent antibodies. Phase II assays measure platelet-associated immunoglobulins that associate with platelets both non-specifically and through specific binding to antigens. Total platelet associated immunoglobulin assays measure intracellular IgG released by platelet lysis, in addition to surface associated immunoglobulins. Since non-specific platelet binding of immunoglobulins is common in thrombocytopenic disorders other than ITP, these assays proved to be relatively non-specific, though very sensitive for ITP.

The development of phase III assays provided the opportunity to improve specificity for platelet autoantibody detection, as well as a platform for the identification of alloantibodies through the use of specific platelet glycoprotein targets. In the simplest form of these assays, the ability to type platelet glycoprotein antigens allows for screening of patients' serum or plasma against antigen-defined, donor platelets with subsequent detection of bound antibody through immunofluorescence detected by cytofluorometry. This format can be modified to detect drug-dependent antibodies by incubating the platelets with the test drug prior to incubation with patient serum/plasma.

In the more sophisticated assays, antigenically defined, immunopurified or solubilized platelet glycoproteins are captured on a solid phase support (e.g., plastic wells, beads) for testing. A patient's serum/plasma or an eluate prepared from a patient's own platelets (autoantibody) or from sensitized donor platelets (auto- or alloantibody) is incubated with the glycoproteins, and binding of antibody is detected using labeled, anti-human antibody. Detection can be accomplished through radioactivity, standard ELISA methods, immunofluorescence or RBC agglutination. In a variation of this format, auto- or alloantibody is bound to patient or donor platelets, respectively, before platelet solubilization. The antibody-glycoprotein complex is then captured and the bound antibody is detected as already described. The advantage of this test format is that the glycoprotein epitopes are in their native state when antibody is bound, and not modified by the purification process.

Testing for Heparin-Dependent Antibodies

The administration of heparin to patients can result in the formation of heparin-dependent antibodies that are directed against a neoantigen produced by the association of heparin with platelet factor 4 (PF4). The production of heparin-dependent antibodies can result in heparin-induced thrombocytopenia (HIT) - a clinical syndrome characterized by thrombocytopenia and a risk of venous or arterial thrombosis of 30–75%¹². The College of American Pathologists recommends that reference laboratories validate more than one assay for the detection of heparin dependent antibodies to maximize sensitivity and specificity¹³. An immunologic assay and a functional assay are routinely employed in the evaluation of patient serum for heparin dependent antibodies.

The PF4 ELISA tests are immunoassays that employ PF4 complexed with either heparin or a polyanionic compound to bind heparin dependent antibodies¹⁴. Positive reactions can be confirmed by inhibiting antibody binding by adding excess heparin to the reaction. The PF4 ELISA assay is a sensitive assay ($\geq 90\%$), but less specific ($\leq 90\%$) than the serotonin release assay (SRA) discussed below. The particle gel immunoassay (PaGIA), uses PF4-heparin complexes bound to colored particles to bind heparin-dependent antibodies that results in agglutination of the antibody bound particles. The particles are centrifuged through gel columns such that agglutinated particles remain at the top of the gel, while free particles migrate to the bottom of the column. These particles can also be adapted for use in a cytofluorometry-based assay. The PaGIA exhibits slightly lower sensitivity than the ELISA but is closer to the SRA in specificity¹⁴.

Due to less complicated test formats, the PF4 immunoassays are frequently used as screening tests for patients suspected of a diagnosis of HIT. PF4 immunoassays that detect only IgG antibodies demonstrate better specificity, without loss of sensitivity¹³, though none are commercially available at this time. A strong positive result has greater predictive value for a diagnosis of HIT than a weak positive, and the British Committee for Standards in Haematology recommends reporting the optical density value of a test as well as the cutoff value for a positive result¹⁵.

The SRA is a functional assay that identifies heparin dependent antibodies through induction of $\geq 20\%$ serotonin release from donor platelets in the presence of low concentrations of heparin (0.1–0.3 U/mL), and $\geq 50\%$ inhibition of that release in the presence of high concentrations of heparin (100 U/mL). SRA test sensitivity

reportedly ranges from 90 to 98% depending upon the expertise of the laboratory, and specificity is generally $\geq 95\%$ except when the decrease in platelet count occurs 5 or more days after heparin exposure¹³. Due to the high specificity of this test, it is considered the “gold standard” method. However, the requirement for fresh platelets susceptible to heparin dependent antibodies, the need to wash the platelets to maximize sensitivity, and the use of radioisotopes to detect serotonin release makes this assay impractical for most laboratories. Consequently, it is generally used as a confirmatory test¹³.

One laboratory has noted a high rate of false-negative results on the initial specimens collected from patients suspected of having HIT¹⁴. In a retrospective review, one third of negative results by immunologic assay turned positive on subsequent specimens collected from patients with continuing suspicion of HIT. A second report documents three of eleven patients with HIT who were negative by immunologic and functional assay on the initial specimen, but seroconverted over the following seven days¹⁶. Causes for false negative results have not been determined, but may include low titers at the time of testing, and interference from PF4-heparin complexes circulating at the time of specimen collection¹⁶. Therefore, in cases where clinical suspicion for HIT is high, retesting of additional specimens collected after a negative result may be indicated.

Clinical Utility of Platelet Antibody Tests

Platelet antibody testing is indicated for the detection of alloantibodies in cases of NAIT, post-transfusion purpura (PTP), and for heparin-dependent antibodies in heparin-induced thrombocytopenia (HIT). Even though the platelet count recovers after withdrawal of heparin, confirmation of HIT is helpful in guiding decisions about further anticoagulation therapy. Testing for other drug-induced platelet antibodies is usually not necessary, since thrombocytopenia generally resolves after withdrawal of the implicated drug, and non-cross-reacting drugs can generally be substituted for the original drug. Testing for autoantibodies in suspected cases of autoimmune thrombocytopenia is more controversial. Phase II tests for platelet associated immunoglobulins are too non-specific to be useful. Phase III tests that identify binding of autoantibody to platelet-specific glycoproteins appear to have improved positive predictive value, but poor negative predictive value¹⁷. If this relationship is born out in larger prospective trials, phase III assays would be useful for ruling in AITP, but not for ruling it out.

Platelet Function Testing

Platelet function testing by standard platelet aggregation tests is the gold standard for identifying platelet dysfunction, however, availability is limited to specialty laboratories. Consequently, new, less manual test devices are being evaluated for routine use (see Chapter 3, Instrumentation for the Coagulation Laboratory). Acquired platelet dysfunction may be produced by disease states or by administration of therapeutic drugs or interventions (Table 7.2).

Myeloproliferative Disorders

Myeloproliferative disorders (MPD) may be complicated by platelet dysfunction. Platelet function testing is not indicated in these disorders, however, since it is not predictive of clinical outcome¹⁸.

Uremia

Renal failure predisposes patients to bleeding secondary to anemia due to decreased displacement of circulating platelets towards vessel walls by red blood cells, and due to uremia. Uremia leads to the overproduction and accumulation of nitric oxide in the circulation that directly inhibits platelet function¹⁹. Uremic patients are at increased risk for spontaneous bleeding as well as excessive bleeding as a complication of invasive procedures – most commonly renal biopsy. Dialysis or treatment with desmopressin acetate (DDAVP) significantly reduces the bleeding risk. Unfortunately, the level of uremia cannot predict the risk of bleeding. The bleeding time test has traditionally been used to assess bleeding risk, but the predictive value of the test is poor²⁰. Preliminary evaluations of two other methods have been reported. In one study, the PFA-100® failed to detect uremia-induced platelet dysfunction after the hematocrit of the samples had been normalized²¹. In a second study, thromboelastography demonstrated positive and negative predictive values of only 71 and 74%, respectively, for post-biopsy, renal hemorrhage²².

Antiplatelet Drugs

Interest in monitoring antiplatelet drugs has focused on aspirin, clopidogrel, and the platelet GP IIb/IIIa receptor inhibitors. Clinician interest in measuring the platelet effects of aspirin are two-fold: to determine

the potential for a slightly increased risk of bleeding during and after invasive procedures, and to determine the degree of aspirin-induced platelet inhibition in patients taking prophylactic doses to prevent cardio- and cerebrovascular events. The increased number of patients taking aspirin for its anti-thrombotic effects means more patients present for invasive procedures with aspirin on-board. In general, there is little need for determination of platelet function in these patients. For elective procedures, aspirin should be discontinued, and the procedure should be delayed 3–5 days allowing regeneration of roughly one half of the platelet pool²³. The exception to this rule would be patients undergoing emergency coronary artery bypass grafting (CABG) who have not received other pre-operative anticoagulants or anti-platelet agents, and do not have either thrombocytopenia or platelet dysfunction. Platelet transfusions are indicated when surgery must be performed emergently, and bleeding is unexpectedly brisk. Aprotinin may be indicated for CABG patients who have received aspirin and have other risks for bleeding, as noted above²³.

Resistance to the platelet inhibitory effects of aspirin, as documented by laboratory testing, has been clearly demonstrated. There is preliminary evidence that this “aspirin resistance” can result in treatment failure and increased thrombotic events²⁴. Platelet resistance as predicted by the PFA-100[®] and the VerifyNow[™] platelet function analyzers has been reported to correlate with treatment failure. The International Society on Thrombosis and Haemostasis recommends that testing for aspirin resistance be limited to research protocols only, since appropriate therapy for aspirin resistant individuals has not been defined, and current assays have not been sufficiently validated²⁴.

The current situation for monitoring the effects of clopidogrel, a platelet ADP receptor blocker, and the GP IIb/IIIa inhibitors is similar to aspirin. Due to a risk for excessive post-operative bleeding, the Society of Thoracic Surgeons recommends discontinuation of clopidogrel 5–7 days before CABG, and discontinuation of GP IIb/IIIa inhibitors either 4–6 hours before surgery (short-acting inhibitors), or 12–24 hours before surgery (long-acting inhibitors). Platelet transfusion has been demonstrated to decrease the incidence of post-operative bleeding when GP IIb/IIIa inhibitors are still on-board during surgery²³. Preliminary evidence suggests that resistance to the actions of clopidogrel, as measured by a cone and plate analysis²⁵, and resistance to GP IIb/IIIa inhibitors, as measured by the VerifyNow[™] ²⁶, occurs in patients and may predict therapeutic failure. Of note, the PFA-100[®]

has not been found to be useful for monitoring the platelet effects of either clopidogrel²⁷ or GP IIb/IIIa inhibitors²⁶.

Cardiopulmonary Bypass

Cardiopulmonary bypass (CPB) procedures result in a demonstrable, transient platelet function defect that has been correlated with post-operative bleeding. Proposed causes of this platelet dysfunction include the platelet effects of high-dose heparin administered before and during the procedure, activation of platelets upon exposure to the materials that make up the CPB circuit, hypothermia, and platelet effects of protamine administration post-procedure²⁸. Thromboelastography (TEG[®], ROTEM[®]), the Sonoclot[®], the PFA-100[®], the HemoSTATUS[™], the Plateletworks[®] system, and the Hemostasis Analysis System have all been reported to detect platelet dysfunction secondary to CPB and to correlate with excessive bleeding²⁹⁻³¹, though data supporting the predictive value of the HemoSTATUS[™] and the PFA-100[®] are inconsistent³¹.

TEG[®] and the PFA-100[®] have been incorporated into transfusion protocols for cardiac surgery requiring CPB with resulting decreases in both transfused blood components as well as blood loss compared with standard transfusion practice^{32,33}. However, it is difficult to assess the role of rapid assessment of hemostasis through near-patient testing, in general, and the role of platelet function assessment, specifically, in the improved outcomes. Cardiac surgery transfusion practice can be highly variable across surgeons and institutions³⁴, and implementation of a strict protocol may contribute to a decrease in transfused components. A recent clinical trial comparing two algorithm-driven transfusion protocols for cardiac surgery, one including TEG[®] and PFA-100[®] monitoring, and the other utilizing routine laboratory coagulation testing, failed to demonstrate either a difference in the number of components transfused or a difference in median blood loss. Both algorithm-driven protocols demonstrated decreased blood component transfusion when compared with a historical control group, but no difference in median blood loss³⁵.

Summary and Key Points

Excessive bleeding (and less commonly thrombosis) may result from acquired platelet disorders that include thrombocytopenia and/or platelet dysfunction. There have been significant improvements in platelet

counting and antibody detection methods in recent years. There has also been a concurrent development of automated platelet function analyzers that are designed for routine use in the laboratory or at the patient's bedside. The clinical role for routine platelet function testing, however, has not yet been defined.

- An immunologic-based reference method has been endorsed for use by manufacturers and reference laboratories for the validation of automated platelet counters.
- Optical- and impedance-based platelet analyses perform comparably in the general patient population. Optical-based platelet analysis is superior to impedance-based analysis when known interferences are encountered in specific patient populations.
- Phase II platelet associated immunoglobulin assays are too non-specific to be useful for the diagnosis of autoimmune thrombocytopenia. Phase III assays are more specific, but need further clinical validation to assess diagnostic utility.
- Tests for heparin-dependent antibodies are indicated for the diagnostic workup of HIT, but may be falsely negative on the initial specimen drawn when the diagnosis is first considered.
- Clinical indications for acquired platelet dysfunction testing by automated analyzers have not been sufficiently defined, nor have outcomes been sufficiently validated to warrant routine use.

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8

Acquired Coagulation Disorders and TTP

George M. Rodgers

Acquired Coagulation Disorders and TTP

The most common causes of acquired clotting factor deficiencies associated with bleeding are decreased or abnormal synthesis of clotting factors caused by liver disease or disseminated intravascular coagulation (DIC). The latter is seen in many severe illnesses, including metastatic cancer and infectious diseases. Vitamin K deficiency is another common cause of bleeding especially in hospitalized patients. Uncommon causes of acquired coagulation bleeding disorders include antibodies to coagulation factors and abnormal fibrinolysis.¹

Thrombotic thrombocytopenic purpura (TTP) is encountered in clinical practice primarily as an acquired, thrombotic, autoimmune disorder directed against the ADAMTS-13 protease that is responsible for the processing of ultra-high-molecular-weight multimers of von Willebrand factor (UHMWM-vWF) to normal vWF.²

Liver Disease and Vitamin K Deficiency

The liver is the major site of clotting factor synthesis, and bleeding can occur in patients with severe hepatitis or cirrhosis. In these disorders, coagulation protein synthesis is reduced (all clotting factors but factor VIII and von Willebrand factor). In addition to altered coagulation protein levels, other hemostatic defects exist with liver disease, including decreased clearance of activated clotting factors and increased levels of degradation products of fibrinogen and fibrin. Fibrin degradation products inhibit hemostasis by interfering with both platelet function and fibrin formation. Fibrinolysis also may be enhanced in liver disease¹.

Significant liver disease, with associated portal hypertension and splenomegaly, can result in mild to moderate thrombocytopenia. Splenomegaly results in splenic sequestration of platelets. Hepatoma and cirrhosis also are associated with synthesis of qualitatively abnormal fibrinogen (dysfibrinogen) (1).

When vitamin K is deficient, the vitamin K-dependent clotting factors do not bind calcium, and although they are synthesized in normal amounts, are inactive. Vitamin K deficiency can be seen in patients with hepatic or biliary disease, malabsorption, malnutrition, warfarin therapy, or antibiotic treatment¹.

Vitamin K deficiency is a clinical diagnosis. Hospitalized patients who have screening coagulation test results of an isolated, prolonged prothrombin time (PT) or prolongation of both the PT and partial thromboplastin time (PTT) values, and that have clinical associations as outlined above will likely have vitamin K deficiency; no further laboratory tests are usually necessary. For patients in whom the diagnosis is uncertain, a D-dimer test will rapidly evaluate DIC (see Chapter 5, Hemostasis Screening Assays). The response to oral or parenteral vitamin K is also diagnostic of vitamin K deficiency with improvement or normalization of the PT and PTT within 24 hours. If laboratory diagnosis of vitamin K deficiency is important, factor assays demonstrating low levels of the vitamin K-dependent coagulation proteins (prothrombin, VII, IX, X) with normal levels of the other coagulation proteins (e.g., factors V and VIII) confirm vitamin K deficiency.

DIC

The basis for DIC is excessive thrombin formation, which may result in thrombi in the microcirculation, and/or depletion of platelets and fibrinogen, leading to a bleeding tendency. Common causes of DIC include sepsis, burn or traumatic injuries, obstetrical complications, and cancer. The hallmark of all of these disorders is unregulated thrombin activity in blood¹.

Figure 8.1 illustrates the consequences of thrombin activity in blood: thrombin cleavage of fibrinopeptides A and B from fibrinogen generates fibrin monomers; polymerization of fibrin monomers form fibrin polymers; factor XIII_a cross-linking activity generates insoluble fibrin polymers; and lastly, the fibrinolytic mechanism produces fibrin degradation products (FDPs), including D-dimer, a clinically-useful

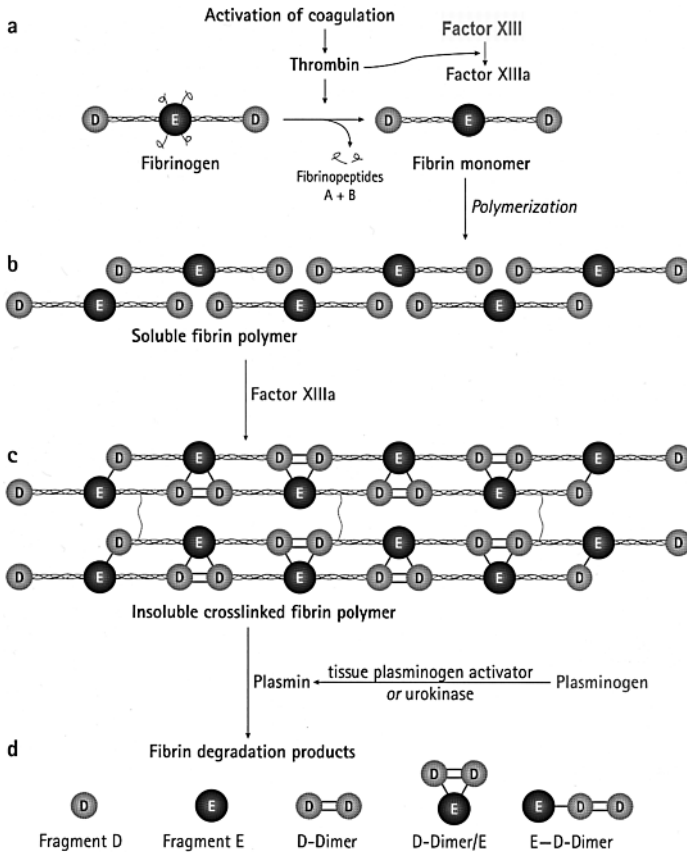


Figure 8.1. Generation of cross-linked fibrin by thrombin and factor XIIIa following activation of coagulation, and consequences of plasmin degradation of cross-linked fibrin. (a) Following activation of coagulation, thrombin activates the transglutaminase, factor XIII to factor XIIIa, and cleaves fibrinopeptides A and B from fibrinogen to generate fibrin monomers. (b) Fibrin monomers align longitudinally and with adjacent monomers to form fibrin polymers. (c) Factor XIIIa cross-links the D-domains of fibrin monomers to form rigid fibrin polymers. (d) Plasminogen activators (TPA or UK) convert plasminogen to plasmin; plasmin degrades cross-linked fibrin to fibrin degradation products, including fragment D, fragment E, and D-dimer, in addition to other products. From Rodgers GM, Acquired coagulation disorders. In Kjeldsberg CR, ed. Practical Diagnosis of Hematologic Disorders. ASCP Press. Chicago, 2006, Fourth ed., p. 359, with permission.

FDP whose presence in large amounts confirms the diagnosis of DIC (see Chapter 5, Hemostasis Screening Assays).

Antibodies to Coagulation Factors

Rarely, patients with previously-normal hemostasis who develop prolonged PT and/or PTT assays and bleeding will not have either vitamin K deficiency or DIC. These patients will probably have acquired antibodies against a clotting factor, usually factor VIII or factor V¹. The appropriate evaluation is to first perform mixing studies (see Chapter 5, Hemostasis Screening Assays) to determine if an inhibitor (antibody) is present. Next, specific factor assays confirm the affected factor. Thus, for a patient with previously-normal PT and PTT values who develops bleeding with an isolated, prolonged PTT, failure to correct the prolonged PTT with a mixing study, followed by assay of factors VIII, IX, XI would diagnose this patient's acquired bleeding disorder. Antibodies to factor VIII are the most commonly seen.

For patients who have acquired prolonged PT and PTT values, performing mixing studies with both the PT and PTT assays is indicated, followed by assay of common pathway coagulation proteins—fibrinogen, prothrombin, factor V, and factor X. Of these clotting factors, antibodies to factor V are most commonly seen.

Abnormal Fibrinolysis

Abnormal fibrinolysis is an uncommon bleeding disorder in which excessive plasmin is generated. Plasmin is appropriately generated following initiation of coagulation and thrombin formation (secondary fibrinolysis). For example, patients with DIC have secondary fibrinolysis which is appropriate. In contrast, in abnormal or primary fibrinolysis, there is no laboratory evidence for thrombin formation (negative D-dimer). Examples of clinical conditions associated with abnormal fibrinolysis include inherited α_2 -antiplasmin deficiency, tumors of the genito-urinary tract, gynecologic tumors, acute promyelocytic leukemia, liver disease, and fibrinolytic drugs¹.

The diagnosis of abnormal fibrinolysis is suggested by a low fibrinogen level, elevated FDP levels, a negative D-dimer assay, and an underlying disorder as outlined above. The presence of an elevated D-dimer level indicates that the abnormal fibrinolysis is secondarily due to excessive thrombin formation, probably DIC.

Thrombotic Thrombocytopenic Purpura

Thrombotic thrombocytopenic purpura usually presents as an acquired thrombocytopenic disorder in adults. The pathologic basis for TTP is autoimmune; patients acquire an auto-antibody to the protease responsible for processing ultra-high-molecular-weight multimers of von Willebrand factor to normal vWF. This protease is known as the vWF-cleaving protease or ADAMTS-13². Failure to process UHMWM-vWF leads to disseminated platelet thrombosis and the clinical manifestations of TTP—microangiopathic hemolytic anemia, thrombocytopenia, neurologic abnormalities, fever, and renal dysfunction. TTP may rarely be inherited (Upshaw-Schulman syndrome); in these cases, the disorder presents in childhood in patients with mutations in the ADAMTS-13 gene².

TTP is primarily a clinical diagnosis because the fulminant nature of the disease requires emergent therapy. However, since the recognition of the basis of the disease, assays have been developed to more rapidly provide a diagnosis. Previous methods were based on demonstrating the presence of UHMWM-vWF in TTP patient plasma; this method required an immunoblot technique and took 1–2 days turn-around time. More recent methods quantitate ADAMTS-13 activity using a fluorogenic substrate (FRETs-vWF73) that provides assay results in a few hours. Results are expressed in terms of ADAMTS-13 activity in normal, pooled plasma; most patients with classic TTP will have ADAMTS-13 levels < 5–10% of normal³.

Patients with hemolytic-uremic syndrome (HUS) generally will have normal ADAMTS-13 levels; therefore, ADAMTS-13 activity assays provide a means to distinguish TTP from HUS (4).

Summary and Key Points

- Acquired coagulation disorders are most commonly caused by vitamin K deficiency, liver disease, and DIC. The clinical situation, routine hemostasis tests (PT, PTT, platelet count) and a D-dimer assay will usually suggest the correct diagnosis.
- Uncommon acquired coagulation disorders include antibodies to clotting factors (usually factors VIII or V) and are suggested by results of the routine hemostasis tests (PT, PTT) and mixing tests.

- Abnormal fibrinolysis is suggested by the appropriate clinical situation, a low fibrinogen level, elevated FDP levels, and a normal D-dimer assay.
- The diagnosis of TTP remains primarily clinical; however, availability of assays for the ADAMTS-13 protease (FRETs-vWF73 assay) offers the possibility of a rapid laboratory confirmation of the diagnosis.

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9

Testing for Inherited and Acquired Thrombotic Disorders

George M. Rodgers

Laboratory Testing of Inherited and Acquired Thrombotic Disorders

Virchow's triad, the major risk factors that predispose to thrombosis, include vascular injury, stasis, and hypercoagulability. Arterial and venous thrombosis have different pathogenic mechanisms. For example, arterial thrombosis primarily involves vascular injury (atherosclerosis) and platelet deposition. Venous thrombosis primarily involves stasis of blood flow and hypercoagulability, which is defined as altered blood composition associated with a thrombotic tendency. Fibrin deposition is the normal consequence of venous thrombosis.

The laboratory evaluation of inherited thrombosis is based on identifying abnormalities in regulatory proteins of coagulation; also known as natural anticoagulants, as well as mutations of other coagulation proteins. Over the past 10–15 years, major inherited etiologies of thrombosis including the factor V Leiden and prothrombin gene mutations have been identified. Acquired etiologies of thrombosis, including cancer, antiphospholipid antibodies, and hyperlipidemia are more common than inherited etiologies. This chapter will summarize the pathologic basis for thrombosis as related to laboratory testing for inherited and acquired etiologies. Methods for laboratory evaluation of thrombosis will be presented, and the utility of laboratory thrombosis testing will be discussed.

Pathophysiology of Thrombosis

Vascular endothelium plays a pivotal role in maintaining thromboresistance of the blood. The coagulation mechanism is modulated by several

endothelial cell regulatory mechanisms¹. Figure 9.1 summarizes the antithrombotic properties of endothelium that can be evaluated in the laboratory. A key regulatory mechanism is the protein C pathway, which consists of two vitamin K–dependent plasma proteins—protein C and protein S. Protein C is converted to an active form, activated protein C (APC); this activation is mediated by an endothelial cell receptor, thrombomodulin. Thrombomodulin forms a complex with thrombin; this complex activates protein C generating APC, which in turn inactivates factors Va and VIIIa. Protein S binds to the endothelial cell surface, providing a receptor for APC¹. A factor V gene mutation called factor V Leiden results in thrombosis due to the inability of APC to degrade the abnormal factor V_a molecule. This phenomenon is termed APC resistance and can be measured by the laboratory².

Antithrombin is a natural anticoagulant that irreversibly binds to and inactivates activated clotting factors such as factor X_a and thrombin. This inactivation is catalyzed by heparin-like glycosaminoglycans on the endothelial cell surface (see Figure 9.1) or by therapeutic heparin¹.

The fibrinolytic mechanism components include plasminogen, tissue-plasminogen activator (TPA), plasmin, α_2 – antiplasmin, and plasminogen activator inhibitor (PAI-1). Fibrinolysis is initiated when vascular thrombosis triggers endothelial cell secretion of TPA. In the presence of TPA, plasminogen is converted to plasmin that degrades fibrin clots. Plasmin activity is regulated by α_2 – antiplasmin, while TPA activity is regulated by PAI-1. Either deficiency or excess of these components may occur leading to hypofibrinolysis and a thrombotic risk or hyperfibrinolysis and a bleeding risk¹.

The Inherited Thrombotic Disorders

Table 9.1 summarizes the inherited thrombotic disorders and describes their prevalence, inheritance patterns, and clinical features. Abnormalities of the protein C pathway (protein C, protein S, factor V Leiden, thrombomodulin) constitute most cases of inherited thrombosis². Most inherited disorders are transmitted in an autosomal dominant manner, and venous thromboembolism is the usual clinical feature. The importance of inherited fibrinolytic disorders (TPA deficiency or excess PAI-1 activity) is uncertain.

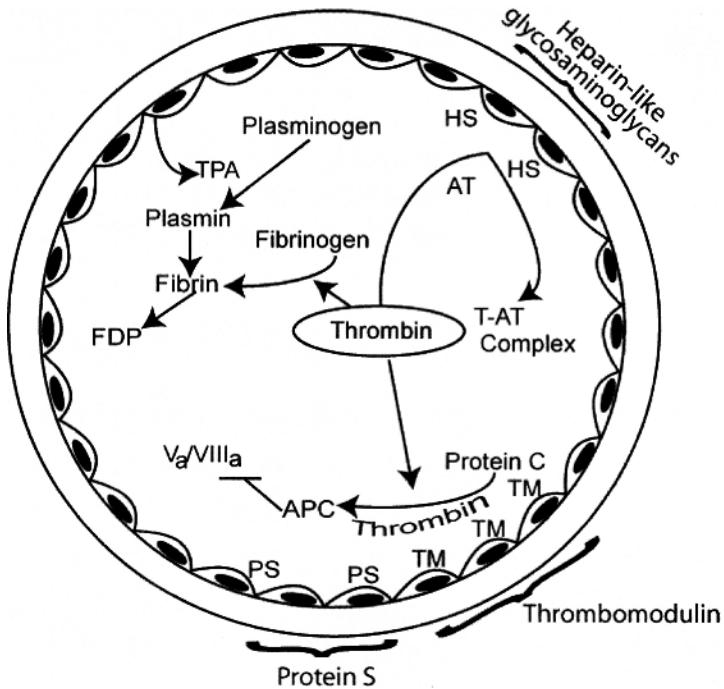


Figure 9.1. Regulation of thrombin activity by vascular endothelium. The effects of major antithrombotic properties of the blood vessel wall are shown. Regulation of thrombin activity is important because thrombin is a major factor in thrombosis (platelet activation, fibrin formation). Abnormalities in regulation of thrombin activity may lead to hypercoagulability and an increased risk of thrombosis. Major antithrombotic properties include: heparin-like glycosaminoglycans (heparan sulfate, HS) on the luminal surface that catalyze antithrombin (AT) inhibition of thrombin, generating inactive thrombin-antithrombin (T-AT) complexes; thrombomodulin (TM), an endothelial cell receptor for thrombin. The thrombin-TM complex converts protein C to APC; protein S functions as a cofactor binding protein for APC, permitting inactivation of factors $V_a/VIII_a$, resulting in inactivation of coagulation; and endothelial cell secretion of tissue-plasminogen activator (TPA) that initiates fibrinolysis. Most of the coagulation components shown in this figure can be assayed by the laboratory to identify thrombosis risk. From Rodgers GM. Hemostatic properties of normal and perturbed vascular cells. FASEB J 1988; 2: 116–23, with permission.

Table 9.1. Summary of the Inherited Thrombotic Disorders

| Classification and Disorders | Inheritance | Estimated Prevalence* | Clinical Features |
|---|--------------------|------------------------------|--|
| <i>Deficiency or qualitative abnormalities of inhibitors to activated coagulation factors</i> | | | |
| AT deficiency | AD | 1% | Venous thromboembolism (usual and unusual sites), heparin resistance |
| TM deficiency | AD | 1%–5% | Venous thrombosis |
| Protein C deficiency | AD | 1%–5% | Venous thromboembolism |
| Protein S deficiency | AD | 1%–5% | Venous and arterial thromboembolism |
| APC resistance due to Factor V Leiden | AD | 20%–50% | Venous thromboembolism |
| <i>Abnormality of coagulation zymogen or cofactor</i> | | | |
| Prothrombin mutation | AD | 5%–10% | Venous thromboembolism |
| Elevated factor VIII | Unknown | 20%–25% | Venous thromboembolism |
| Elevated factor IX | Unknown | ~10% | Venous thromboembolism |
| Elevated factor XI | Unknown | ~10% | Venous thromboembolism |
| <i>Impaired clot lysis</i> | | | |
| Dysfibrinogenemia | AD | 1%–2% | Venous thrombosis > arterial thrombosis |
| Plasminogen deficiency | AD,AR | 1%–2% | Venous thromboembolism |

(Continued)

Table 9.1. (Continued)

| | | | |
|-------------------------|----|---|--|
| TPA deficiency | AD | ? | Venous thromboembolism |
| Excess PAI-1 activity | AD | ? | Venous thromboembolism and arterial thrombosis |
| <i>Metabolic defect</i> | | | |
| Homocysteinemia | AR | 1 in 300,000 live births; 10–25% of patients with recurrent thrombosis | Arterial and venous thrombosis (homozygous patients); premature development of coronary and cerebral arterial thrombotic disease (heterozygous patients) |

Abbreviations: AT = antithrombin; APC = activated protein C; TPA = tissue plasminogen activator; PAI-1 = plasminogen activator inhibitor-1; TM = thrombomodulin; AD = autosomal dominant; AR = autosomal recessive; ? = uncertain prevalence of abnormal fibrinolysis.

* Prevalence data are estimated by pooling information from studies in which large groups of patients with thrombosis were screened for these disorders. Results are expressed in terms of a percentage that each disorder might constitute of the total patient population with inherited thrombosis. (Assays for TM mutations are not widely available.)

Source: Robetorye RS, Rodgers GM. Update on selected inherited venous thrombotic disorders. *Am J Hematol* 2001; 68: 256–268.

Another common inherited thrombotic disorder is the prothrombin gene mutation. This mutation is associated with elevated plasma prothrombin levels, which may explain the predisposition to thrombosis³.

Homocysteinemia is a metabolic disorder associated with thrombosis. Although pediatric patients present clinically with homozygous homocysteinemia, (homocystinuria), adult patients heterozygous for homocysteinemia have primarily premature arterial disease (myocardial infarction, stroke, peripheral vascular disease). Heterozygous homocysteinemia may account for a significant number of patients with arterial vascular disease in the absence of traditional risk factors (e.g., smoking, hypertension, hyperlipidemia). Between 1% to 2% of the general population have heterozygous homocysteinemia. Homocysteinemia is also associated with venous thromboembolism⁴.

A recently described inherited risk factor for thrombosis is elevated levels of factor VIII activity. Although factor VIII is an acute-phase response protein, as many as 10% to 20% of patients with recurrent venous thrombosis have elevated factor VIII levels as their only risk factor².

Increased levels of other coagulation factors, including fibrinogen, factor IX, and factor XI have also been associated with thrombosis. However, routine laboratory testing of these factors is not recommended by the College of American Pathologists Consensus Conference on Thrombophilia (see Table 9.2).

General Principles of Thrombosis Testing

1. Laboratory evaluation should be postponed until 2 to 3 months after the acute thrombotic event when the patient is clinically well and has not received anticoagulant therapy for 2 weeks. Thrombosis induces an acute-phase response that makes interpretation of coagulation-based tests difficult. Reliable data for assays such as antithrombin, protein C, and protein S activities are best obtained in the absence of anticoagulants. If anticoagulants cannot be discontinued in the affected patient, then surrogate testing of symptomatic family members who are not receiving anticoagulants can be done. However, if DNA-based assays (for the factor V Leiden or the prothrombin gene mutations) are performed, these results will not be affected by acute-phase changes of thrombosis or anticoagulant

Table 9.2 Summary of the College of American Pathologists' recommendations on laboratory testing for inherited thrombosis

| Thrombotic Disorder | Who should be tested? | Test method(s) | Comments |
|----------------------------|---|---|---|
| Factor V Leiden | First VTE at age <50 years Recurrent VTE First unprovoked VTE First VTE, unusual site First VTE, positive family history First VTE related to pregnancy or hormonal therapy Unexplained 2nd or 3rd trimester pregnancy loss As above | APC resistance assay using factor-V deficient plasma or DNA-based assay | Patients with relatives who are known to have FVL should be tested directly with DNA-based assays. Patients with positive APC resistance assays should have confirmatory DNA tests. |
| Prothrombin gene mutation | As above | DNA-based assay | Prothrombin activity assays should not be used |
| Homocysteinemia | Arterial vascular disease Controversial for VTE | HPLC or immunoassays | Genotyping for MTHFR mutations is not recommended. Fasting may not be necessary. Proper sample processing is necessary. Testing in VTE patients may be appropriate to identify and treat affected patients with vitamins. |

(Continued)

Table 9.2 (Continued)

| Thrombotic Disorder | Who should be tested? | Test method(s) | Comments |
|----------------------------|--|---|--|
| Protein C deficiency | Infants with neonatal purpura fulminans VTE patient from a family with known PC deficiency Asymptomatic female from a known PC-deficient family prior to hormonal therapy Patient with VTE from a family with known PS deficiency | Chromogenic substrate assays are preferred Functional assays are useful Immunologic assays are discouraged Functional assay or Immunoassay for free PS Total PS antigen assays not recommended | Avoid testing during acute thrombosis or anticoagulant therapy. Exclude causes of acquired PC deficiency. Consider age-dependent reference ranges. Abnormal functional assay results should be confirmed with an immunoassay for free PS. Exclude acquired causes of PS deficiency. Avoid testing during acute thrombosis, anticoagulant therapy, and pregnancy. Consider age- and gender-dependent reference ranges. |

(Continued)

Table 9.2 (Continued)

| | | | |
|-----------------------------------|--|---|---|
| Antithrombin deficiency | Patient with VTE from a family with known AT deficiency Asymptomatic female from a known AT-deficient family prior to hormonal therapy Controversial | Chromogenic substrate assays are preferred AT antigen assays not recommended Factor VIII activity assay | Exclude acquired causes of AT deficiency. Avoid testing during acute thrombosis or anticoagulant therapy. Test 6 months after thrombosis. Avoid anticoagulant therapy. |
| Elevated factor VIII levels | Controversial | Factor VIII activity assay | Test 6 months after thrombosis. Avoid anticoagulant therapy. |
| Dysfibrinogenemia | Not recommended | | |
| Heparin cofactor II | Not recommended | | |
| Factor XIII polymorphisms | Not recommended | | |
| Plasminogen activator inhibitor-1 | Not recommended | | |
| Plasminogen deficiency | Test in non-DVT patients with ligneous conjunctivitis | | |

FVL = factor V Leiden, VTE = venous thromboembolism, APC = activated protein C, HPLC = high performance liquid chromatography, MTHFR =methylentetrahydrofolate reductase, PC = protein C, PS = protein S, AT = antithrombin. From the College of American Pathologists' Consensus Conference on Thrombophilia. Arch Pathol Lab Med 2001; 126:1277-1433, with permission.

- therapy. Similarly, homocysteine testing will not be affected by acute-phase changes or anticoagulant therapy. If factor VIII activity testing is to be done, it should be deferred until 6 months after the thrombotic event².
2. The probability of obtaining positive thrombosis testing results is increased if the patient population being investigated is restricted to young patients (< 50 years of age) with recurrent thrombosis or patients with a single event and a positive family history for thrombosis².
 3. Functional coagulation assays are recommended over immunologic assays for evaluation of antithrombin, protein C, or protein S deficiencies. Functional assays detect both quantitative deficiency and qualitative abnormality of the protein. On the other hand, functional assays are affected by anticoagulant therapy and interpretation of abnormal functional assay results must take into account whether the patient is receiving anticoagulants².
 4. Assay for common etiologies first (factor V Leiden/APC resistance, prothrombin gene mutation, homocysteinemia)².
 5. Heterozygous homocysteinemia should be considered as a cause for thrombosis in middle-aged patients with premature vascular disease as well as a cause of venous thrombosis².

Laboratory Testing Strategy for Inherited Thrombosis

There are two testing strategies for inherited thrombosis—arterial and venous etiologies. Most cases of arterial thrombosis are not inherited, but acquired, including disorders such as diabetes, hyperlipidemia, and other causes of atherosclerosis, plus other etiologies such as vasculitis, myeloproliferative disorders, etc. Inherited etiologies for arterial thrombosis include elevated levels of PAI-1, homocysteinemia, and some patients with protein C or S deficiencies.

In contrast to the limited etiologies for arterial thrombosis, inherited venous thrombosis testing is more detailed. Figure 9.2 depicts one suggested strategy for testing in the venous thrombosis setting⁵. Assay of the most common etiologies is done initially (factor V Leiden/APC resistance, prothrombin gene mutation, homocysteinemia). If these tests are normal, then evaluations of uncommon causes of venous thrombosis are done (antithrombin, protein C, protein S). If the patient is to be evaluated for elevated factor VIII levels, this should be deferred for at least 6 months.

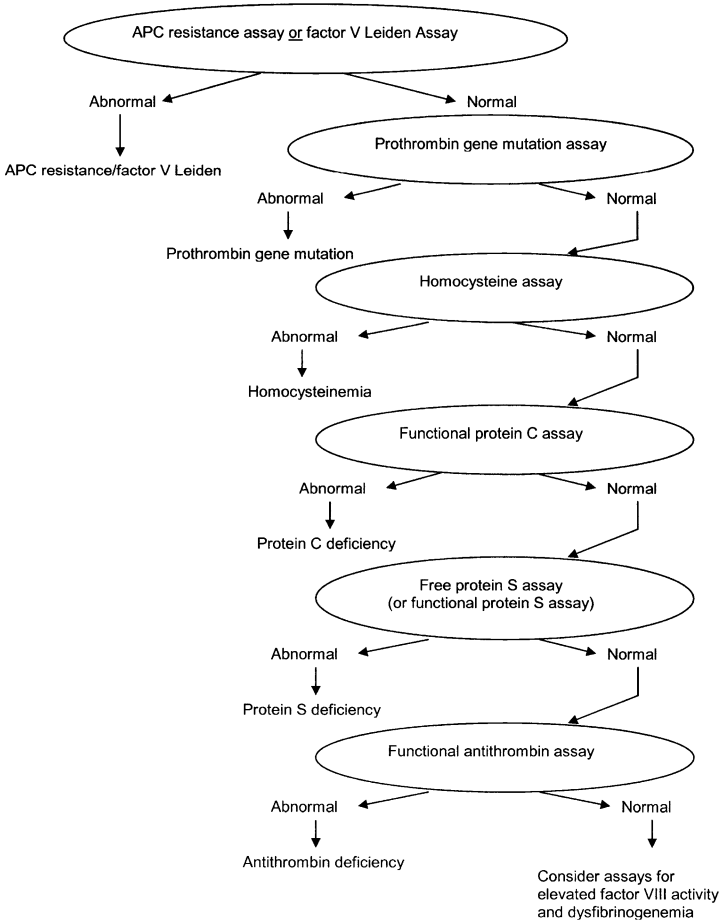


Figure 9.2. An algorithm approach for the evaluation of patients with inherited venous thrombosis. It is assumed that patients have been appropriately selected and that certain assays (antithrombin, protein C, protein S) will not be performed until anticoagulation has been completed. The CAP Consensus Panel recommends confirmation of positive results for APC resistance assays with the factor V Leiden DNA test. Testing for elevated factor VIII levels is controversial, but if this is to be done, testing should be deferred until 6 months after the thrombotic event when anticoagulation has been discontinued. From Rodgers GM, Chandler WL. Laboratory and clinical aspects of inherited thrombotic disorders. *Am J Hematol* 1992; 41: 113–22, with permission.

Utility of Inherited Thrombosis Testing

The two major reasons to evaluate patients for inherited thrombosis are: 1) to screen family members for whom an inherited diagnosis may change management, and 2) to identify an inherited thrombotic disorder in the patient that may affect their management (e.g., intensity or duration of anticoagulation) (6). There is justification for the first reason to test, because for example, if a patient has female siblings or children who might also inherit a thrombotic disorder, this would potentially affect their treatment with oral contraceptives, or change counseling regarding pregnancy. In contrast, there is limited justification to test patients for the second reason (a thrombosis test result affects patient management). Several clinical trials in the past few years have produced results suggesting that positive test results for inherited thrombotic disorders do not predict recurrence of thrombosis; therefore, a positive result would not change patient treatment. Also, there is no data that patients with an inherited thrombotic disorder require a different intensity of anticoagulation than thrombosis patients without an inherited disorder⁶. One conclusion from this data is that testing should primarily be done if there are potentially affected female family members. It should also be noted that comprehensive testing can be expensive – between \$1,000–\$2,000 for a single patient evaluation.

Laboratory Tests for Inherited Thrombosis

This section will briefly describe the principles of interpretation of common assays used to evaluate etiologies of inherited thrombosis. Most assays are available as commercial kits with standard methodologies that can be automated.

Activated Protein C Resistance Assay

APC resistance due to the factor V Leiden mutation is a very common cause of inherited venous thrombosis, especially in Caucasian populations. Test options include a clotting-based assay or a DNA-based test^{7,8}.

APC is an anticoagulant that will prolong the clotting time of normal plasma, but patients with the factor V Leiden (or certain other) mutations will exhibit clotting times that are less prolonged (i.e., these plasmas exhibit APC resistance). In the APC resistance assay, two determinations are made, one with and one without the addition of APC.

A ratio is obtained [e.g., Partial Thromboplastin Time with APC/Partial Thromboplastin Time without APC] and compared with that of a normal population. The ideal APC resistance assay uses either prothrombin time (PT) or partial thromboplastin time (PTT)-based assays in which patient plasma is diluted in factor V-deficient plasma. Using factor V-deficient plasma makes the test useful in patients who are receiving heparin or warfarin therapy, or who have lupus anticoagulants. If a normalized assay is done (the APC ratio of the patient is divided by the APC ratio of a control pooled plasma sample from the same test run), this method can distinguish normal vs. heterozygotes vs. homozygotes with the factor V Leiden mutation. If the laboratory has test samples from large numbers of patients receiving heparin or warfarin, and the local assay method does not use factor V-deficient plasma dilutions, then uninterpretable results may occur. In this situation, it may be preferable to perform factor V Leiden DNA testing.

Factor V Leiden DNA Test

With the discovery of the factor V Leiden mutation and the prothrombin gene mutation, each due to highly conserved point mutations, the utility of molecular diagnostic testing for thrombosis was enhanced.

For diagnosis of factor V Leiden, the original method used the restriction enzyme *Mnl* to digest a 267-bp amplified fragment of patient DNA. Digestion of normal patient DNA results in three fragments. A variety of molecular methods are available (summarized in the CAP Consensus Conference report)⁸, including fluorescent detection of real-time PCR products, as well as non-PCR based methods.

Although DNA-based assays will be more expensive than the APC resistance assays, there will be no interferences, and the result should unequivocally be either normal, or heterozygous or homozygous for the factor V Leiden mutation. Negative test results for the factor V Leiden mutation will not exclude APC resistance due to other genetic defects (~5% of patients with APC resistance).

Prothrombin Gene Mutation Assay

As with the factor V Leiden mutation, the prothrombin gene mutation is mostly seen in Caucasian populations. Although many patients with this mutation will have elevated levels of prothrombin activity, it is recommended that patients be tested for the specific DNA abnormality³.

Multiple molecular diagnostic methods are available for assay, including restriction endonuclease digestion, automated fluorescence detection, or real-time fluorescence detection. Results are reported as normal, heterozygous, or homozygous for the prothrombin gene mutation³.

The typical prothrombin gene mutation is G20210A. A variant prothrombin gene mutation has been reported in African-American patients – C20209T. This latter mutation may be under-recognized because standard PCR assays for the G20210A mutation may not detect the variant mutation. Laboratories that evaluate large numbers of African-American patients for thrombosis testing should consider assays for the C20209T mutation.

Homocysteine Assays

Unlike the other coagulation analytes discussed in this chapter that are linked to thrombosis, homocysteine is an amino acid whose levels can be elevated in inherited or acquired circumstances. It is usually assayed in the chemistry laboratory. Elevated levels of homocysteine may be associated with arterial or venous thrombosis, but consensus opinion suggests focusing on evaluation of patients with arterial thrombosis (Table 9.2)⁴.

There are also molecular diagnostic tests available to identify mutations in the methylene tetrahydrofolate reductase (MTHFR) gene; however, consensus opinion is that testing for the MTHFR mutation should not be performed⁴.

Earlier data suggested obtaining fasting samples for homocysteine measurement; this is no longer recommended. Either plasma or serum samples can be used, but collected blood samples should be placed on ice and red cells promptly separated. Quantification in the past was done by high-performance liquid chromatography, but the availability of fluorescence-based immunoassays has expanded the use of homocysteine measurement in smaller laboratories.

Protein C Assay

Unlike the highly conserved point mutations of factor V Leiden or the prothrombin gene mutations, deficiencies of protein C, protein S, and antithrombin are caused by numerous mutations. Consequently, molecular diagnostic tests are not clinically useful in evaluating patients for these disorders.

A variety of antigenic and functional assays are available to measure protein C levels⁹. Antigenic assays include ELISA, electroimmunoassay (Laurell rocket technique), and radioimmunoassay. The antigenic assays measure protein C levels (normal and des-carboxy forms), so these methods are not affected by oral anticoagulant therapy. However, antigenic assays will not measure protein C function and will therefore not detect qualitative protein C abnormalities. Functional protein C assays are preferred since they will detect both quantitative and qualitative protein C deficiency. Functional assay methods include clotting and chromogenic techniques. A common functional assay method uses Protac[®], a snake venom activator of protein C. The APC generated is then measured in a PTT-based assay (clotting method) or in an amidolytic assay (chromogenic substrate method). The CAP Consensus Study recommends use of the latter chromogenic assay, primarily because therapeutic heparin levels do not affect assay results. Anticoagulant therapy and elevated factor VIII levels will affect the clot-based methods⁹. However, the Russell's viper venom-based clotting method containing heparin neutralizer eliminates the effects of both elevated factor VIII levels and therapeutic heparin.

One important aspect of interpreting protein C levels is that there is age-dependence for protein C levels; younger patients may not be correctly classified unless pediatric reference intervals for protein C levels are considered. Table 6.4 summarizes pediatric reference intervals for coagulation analytes, including protein C.

Numerous acquired variables impact protein C levels (Table 9.3), and interpretation of abnormal results should be made with caution. It is useful for the laboratory to provide a comment on factors that may result in low protein C levels, to assist clinicians in correct interpretation of the test. Ideally, evaluation of protein C deficiency should be done at a time distant from the acute thrombotic event when anticoagulation has been completed for at least two weeks.

Protein S Assay

As with protein C assays, antigenic and functional methods are available to measure protein S levels¹⁰. Total protein S antigen assays can be performed using ELISA methods, Laurell rocket technique, radioimmunoassay, etc. Functional assays can be either based on PT or PTT clotting assays. The "gold standard" test for protein S has been identified as free protein S antigen levels¹⁰. Unlike protein C measurement,

Table 9.3 Acquired Conditions Associated with Protein C, Protein S, and Antithrombin Deficiency

| Analyte | Causes of Deficiency |
|--------------|--|
| Protein C | DIC |
| | Acute thrombosis |
| | Vitamin K deficiency, including oral anticoagulant therapy |
| | Newborn infants, children |
| | Liver disease |
| | Post-operative state |
| Protein S | DIC |
| | Acute thrombosis |
| | Inflammatory illness of any cause |
| | Vitamin K deficiency, including oral anticoagulant therapy |
| | Newborn infants, children |
| | Liver disease |
| | Pregnancy |
| | Nephrotic syndrome |
| Antithrombin | DIC |
| | Acute thrombosis |
| | Liver disease |
| | Oral contraceptives |
| | Nephrotic syndrome |
| | Pregnancy |
| | Heparin (therapeutic levels) |

Conditions, drugs, or diseases listed in this table may result in acquired deficiency of protein C or S, or antithrombin. These causes should be considered before evaluating patients for inherited deficiency.

Abbreviations: DIC=disseminated intravascular coagulation

Source: Kjeldsberg C, et al. Practical Diagnosis of Hematologic Disorders ASCP Press, Chicago, 2006, p. 387, with permission.

functional protein S assays are less useful because factor V Leiden (APC resistance) interferes with functional protein S assays results. For example, spuriously low protein S levels may be seen in patients with factor V Leiden when tested with functional assays for protein S.

Free protein S levels can be measured by monoclonal antibody-ELISA. Functional protein S assays are based on PT or PTT assays. Diluted patient plasma is added to protein S-deficient plasma in the presence of APC and factor Va for the PTT format assay. Positive

results in the functional assay should be confirmed with another assay method, e.g., a free protein S assay.

Numerous acquired conditions may result in protein S deficiency (Table 9.3), and these must be considered in the interpretation of a low protein S test result. It is useful to comment on these acquired etiologies when reporting the test results to assist clinician interpretation. Age-dependent reference ranges are necessary to correctly classify patients (Table 6.4). Additionally, males have higher mean plasma levels of protein S, so gender should also be taken into account when reporting protein S levels.

Antithrombin Assay

Functional and antigenic assays are available to measure antithrombin levels¹¹; functional methods are preferred since many antithrombin-deficient patients have qualitatively-abnormal molecules that would be missed if antigenic assays were used. Functional assays for antithrombin measure heparin cofactor activity using a chromogenic substrate method to assay thrombin or factor Xa inhibition. Antigenic methods include Laurell immunoelectrophoresis, radial immunodiffusion, and microlatex particle immunoassay¹¹.

Plasma antithrombin levels can be markedly decreased by therapeutic heparin, and long-term warfarin treatment may increase antithrombin levels. Therefore, patients should be tested off of anticoagulant therapy. Other acquired conditions that can affect antithrombin results are listed in Table 9.3. Pediatric reference ranges should be used to correctly classify laboratory results of children (see Chapter 6, Testing for Inherited Bleeding Disorders, Table 6.4).

Laboratory Testing for Other Inherited Thrombotic Disorders

Assays exist to measure numerous other analytes that have been linked to thrombosis, for example, dysfibrinogenemias, heparin cofactor II, and fibrinolytic components (plasminogen, tissue plasminogen activator, and plasminogen activator inhibitor-1). However, the association between these parameters and thrombosis is either weak or unproven, and the CAP Consensus Conference has recommended that they not be tested for (Table 9.2).

Assay for elevated factor VIII levels is controversial. Factor VIII is an acute-phase response protein, so elevated levels would be seen

during the acute event. Also, the assay for factor VIII is clot-based, so heparin therapy would affect the measurement. A summary of the CAP recommendations on thrombophilia testing is given in Table 9.2.

The Acquired Thrombotic Disorders

Table 9.4 summarizes the acquired thrombotic disorders. These include autoimmune disorders such as vasculitis and antiphospholipid antibodies, hematologic disorders such as the myeloproliferative disorders, paroxysmal nocturnal hemoglobinuria, thrombotic thrombocytopenic purpura, and heparin-associated thrombocytopenia with thrombosis, metabolic disorders such as obesity, diabetes and hyperlipidemia, and miscellaneous disorders such as pregnancy, nephrotic syndrome, hormone therapy, and cancer. Laboratory evaluation of thrombotic thrombocytopenic purpura and heparin-associated thrombocytopenia are discussed in Chapters 7, Testing for Acquired Platelet Disorders and 8, Acquired Coagulation Disorders and TTP. D-dimer assays which are frequently elevated in cancer patients with thrombosis

Table 9.4 Acquired Thrombotic Disorders

| |
|--|
| Autoimmune disease |
| Vasculitis |
| Antiphospholipid antibodies |
| Inflammatory bowel disease |
| Hematologic diseases |
| Myeloproliferative disorders (polycythemia vera, essential thrombocytosis) |
| Plasma cell disorders (Waldenstrom's macroglobulinemia) |
| Heparin – associated thrombocytopenia with thrombosis |
| Paroxysmal nocturnal hemoglobinuria |
| Metabolic disorders |
| Diabetes |
| Hyperlipidemia |
| Obesity |
| Miscellaneous disorders |
| Trauma |
| Post-surgery |
| Cancer |
| Hormone therapy |
| Nephrotic syndrome |

are discussed in Chapter 5, Hemostasis Screening Assays. Antiphospholipid antibodies are frequently assayed in the coagulation laboratory and will be discussed in this chapter.

Testing for antiphospholipid antibodies (anticardiolipin antibodies, lupus anticoagulant) is appropriate for thrombosis patients (arterial and venous), especially if they have idiopathic clots without a family history, or if there is autoimmune disease. Testing for antiphospholipid antibodies is also appropriate for evaluating patients with recurrent miscarriage. Criteria for diagnosis of antiphospholipid antibody syndrome require demonstration of antibody persistence for 12 weeks (Table 9.5).

Anticardiolipin Antibody Assay

A standardized ELISA is recommended using high-sensitivity microtiter plates; these plates permit greater antigenic density and improve detection of antibodies¹². The buffer system is also important. Reporting of antibody titer is critical in interpretation of an anticardiolipin antibody ELISA result. Moderate or high titer antibody levels are required for diagnosis, and the laboratory should provide clinicians with an interpretive comment that allows correct classification of patients. Positive results for I_gG antibodies are considered the most important, although positive results for I_gM antibodies that persist may also be clinically important. The utility of I_gA anticardiolipin antibody testing is uncertain.

Laboratory evaluation of β_2 -glycoprotein-1 antibodies is being considered in the routine investigation of antiphospholipid antibodies. The most recent consensus conference on laboratory criteria used to diagnose antiphospholipid antibody syndrome recommends testing for antibodies to β_2 -glycoprotein-1. Table 9.5 summarizes the recent criteria for diagnosing patients with antiphospholipid antibody syndrome.

Lupus Anticoagulant (LA) Assay

The LA is an antiphospholipid antibody that affects phospholipid-based coagulation assays; it is clinically associated with thrombosis and miscarriage. The LA antibody actually recognizes a protein in a phospholipid-protein complex; the protein is usually β_2 -glycoprotein-1 or prothrombin. Assays to detect the LA are coagulation-based,

Table 9.5 Revised Clinical and Laboratory Criteria for the Antiphospholipid Antibody Syndrome

Clinical criteria

| | |
|--|---|
| Vascular thrombosis – or pregnancy morbidity - | arterial or venous or small vessel one or more unexplained deaths of a morphologically-normal fetus at or beyond 10 weeks gestation or one or more premature births of a morphologically – normal neonate before the 34th week of gestation because of eclampsia/preeclampsia or placental insufficiency or three or more unexplained consecutive spontaneous abortions before the 10th week of gestation (with exclusion of anatomic, hormonal or genetic causes) |
|--|---|

Laboratory criteria

| | |
|--|---|
| anticardiolipin antibody: or lupus anticoagulant: or anti- β_2 -glycoprotein-1 antibody: | (I _g G/I _g M), in moderate or high-titer, present on at least 2 occasions 12 weeks apart, measured by standardized ELISA present on at least 2 occasions 12 weeks apart measured by ISTH criteria (Table 9.6). (I _g G/I _g M), present in titer > 99th percentile on at least 2 occasions 12 weeks apart, measured by a standardized ELISA |
|--|---|

Definite APS is present if at least one clinical and one laboratory criteria are present.

Source: Miyakis S, et al. International consensus statement on an update of the classification criteria for definite antiphospholipid syndrome (APS). *J Thromb Haemost.* 2006; 4: 295–306.

and therefore can be impacted by numerous variables that affect clot-based tests, including sample collection, processing, and the presence of anticoagulants. A good quality phlebotomy (non-activated sample) is necessary. Platelet-poor plasma (platelet count < 10,000/ μ l) is critically important; if the sample platelet count is higher, many LAs will

Table 9.6 ISTH Criteria for Laboratory Diagnosis of the LA

-
- Prolongation of at least one phospholipid-dependent coagulation test with platelet – poor plasma (PTT, dilute PT, DRVVT, kaolin clotting time, etc.)
 - Failure to correct the prolonged clotting time when patient and normal plasma are mixed
 - Correction of the originally-prolonged clotting time by addition of excess phospholipid
 - Exclusion of other inhibitors to coagulation (heparin, factor VIII antibodies)
-

Abbreviations: ISTH = International Society of Thrombosis and Haemostasis; LA = lupus anticoagulant.

Source: Brandt JT, *et al.* Criteria for the diagnosis of lupus anticoagulants: An update. *Thromb Haemost* 1995; 74: 1185–1190.

not be detected since platelet phospholipid in a frozen-thawed sample will neutralize antibody activity^{12,13}.

Guidelines have been published recommending laboratory criteria for diagnosis of the LA. It is suggested that more than one test system (phospholipid-dependent coagulation assay) be used to optimize identification of the LA. Test options include the PTT, dilute PT, dilute Russell's Viper venom time (DRVVT), kaolin clotting time, Taipan venom test and Textarin time. International Society of Thrombosis and Haemostasis (ISTH) guidelines for laboratory detection of the LA are summarized in Table 9.6.

If the screening test demonstrates a prolonged clotting time, a mixing study is performed with normal plasma. If the mixing study result fails to correct, then an inhibitor is suspected. When excess phospholipid or hexagonal-phase phospholipids are added to the screening test sample, the prolonged clotting time shortens or corrects, validating that the originally-prolonged clotting time was due to antibodies to phospholipid. Lastly, other inhibitors to coagulation should be excluded clinically or by the laboratory (heparin, factor VIII antibodies).

“Integrated” coagulation reagents have been developed that permit testing for the LA using the ISTH criteria. For example, a DRVVT kit is available that uses reagents for screening and confirmation of the LA. Similarly, the Staclot LA[®] test also uses a methodology to screen and confirm the LA. Some of the integrated LA kits contain a heparin neutralizer that permits performing the test on patients receiving heparin therapy.

Utility of Laboratory Thrombosis Testing

The availability of hypercoagulability or thrombosis panels and identification of new thrombophilia disorders have led to increased test ordering of these assays. There are two reasons to test patients – to screen other family members who may be similarly affected, and to possibly change the management of the patient being tested. The current literature is controversial on this subject, with some investigators insistent that widespread testing is useful, while others state that the information obtained is of limited usefulness. Based on the literature (summarized in the review article by Rondina et al.), several conclusions can be drawn:

1. Testing for the inherited disorders (factor V Leiden, prothrombin gene mutation, protein C and S deficiencies, antithrombin deficiency) is of limited utility in guiding treatment for the affected patients. Large clinical trials have demonstrated that positive results for these tests do not predict recurrent thrombosis, and therefore, do not assist clinicians in determining duration of anticoagulation.
2. Testing for the above inherited disorders may be useful if the patient being tested has female siblings or children for whom a diagnosis might change management (e.g., hormonal therapy).
3. Testing for homocysteinemia is controversial. While patients who have this diagnosis made can be treated with vitamins, at this time, there is no data that shows a clear-cut clinical benefit from suppressing homocysteine levels.
4. Testing for antiphospholipid antibodies (anticardiolipin antibodies, lupus anticoagulant) is helpful in patient management, not because it changes the intensity of anticoagulation, but rather because patients should be anticoagulated for the duration that antiphospholipid antibodies persist.

Most clinicians are not aware of these limitations of thrombosis testing. Laboratory directors should educate their clinicians so that they are aware of when thrombosis testing can be helpful (family screening), what tests to order (functional assays for the natural anticoagulants), and when to order them (not acutely; not while the patient is anticoagulated), etc.

Key Points

- The most common inherited thrombotic disorders are activated protein C resistance (factor V Leiden), the prothrombin gene mutation, and homocysteinemia.
- Antithrombin deficiency, and deficiencies of proteins C and S are very uncommon. Optimal assays for these analytes include functional coagulation methods or free protein S antigen assay.
- If assays for antithrombin, protein C, and protein S are to be done, postpone testing until the patient has completed anticoagulation and the patient is clinically well.
- Be aware that inherited thrombosis testing is most useful for family screening. There is little, if any, data that such results alter management of the patient with thrombosis.
- Pediatric reference intervals are now available for inherited thrombosis analytes (antithrombin, protein C, and protein S deficiencies).
- Appropriate testing for antiphospholipid antibodies includes standardized assays for anticardiolipin antibodies, β_2 -glycoprotein-1 antibodies, and the lupus anticoagulant.

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10

Monitoring Anticoagulant Therapy

Sterling T. Bennett

Monitoring Anticoagulant Therapy

Anticoagulant medications are widely used in the prevention and treatment of thromboembolism. For many years, the antithrombotic therapeutic armamentarium was limited essentially to heparin, warfarin, and aspirin, but over the past several years the number of medications has increased significantly, with a consequent demand for better understanding of the laboratory aspects of therapeutic monitoring.

Laboratory monitoring falls into the two broad categories of general monitoring and specific monitoring. General monitoring is directed toward the assessment of bleeding or other untoward effects of therapy. These tests include hematocrit, hemoglobin, platelet count, occult blood, and so forth. The need for general monitoring is common to all anticoagulants and will not be discussed in detail in this chapter.

Specific monitoring is directed toward the assessment of the specific anticoagulant effects of a given medication and is the focus of this chapter. The basics of anticoagulant therapies and the laboratory tests used in their monitoring will be discussed. A summary of anticoagulant monitoring may be found in Table 10.1.

Warfarin

Warfarin is an oral anticoagulant medication, widely used for the prevention of thromboembolic events, including deep venous thrombosis, pulmonary embolism, myocardial infarction, and stroke^{1,2}.

Mechanism of Action

Warfarin is one of the coumarins or vitamin K antagonists. It does not have direct anticoagulant properties, but exerts its effect by inhibiting

Table 10.1. Summary of Anticoagulant monitoring

| Anticoagulant | Mode of Action | Plasma Half-life | Monitoring Test(s) | Therapeutic Range | Reversal | Notes |
|---|--|-------------------------|------------------------------|--|---|---|
| Warfarin (vitamin K antagonists) | Interference with post-translational γ -carboxylation of vitamin K-dependent clotting factors | 20–60 hours | INR | 2.0–3.0, moderate intensity 2.5–3.5, high intensity | Vitamin K, fresh frozen plasma, recombinant factor VIIa | |
| Heparin, unfractionated, standard-dose IV or high-dose SC | Potentiation of AT inactivation of factors IIa and Xa | 0.5–2.5 hours | PTT Anti-Xa heparin assay | Method-specific 0.3–0.7 IU/mL | Protamine sulfate | The PTT therapeutic range endpoints are the PTT responses corresponding to the anti-Xa heparin therapeutic range. |
| Heparin, unfractionated, low-dose SC | Potentiation of AT inactivation of factors IIa and Xa | | None | N/A | None | |
| Heparin, unfractionated, high-dose IV | Potentiation of AT inactivation of factors IIa and Xa | 0.5–2.5 hours | ACT | Procedure-specific | Protamine sulfate | Target ACT is determined by the nature of the procedure, |

(Continued)

Table 10.1. (Continued)

| | | | | | | |
|--------------|--|-------------|--|--|-------------------------------|--|
| Heparin, LMW | Potential of AT inactivation of factors IIa and Xa | 3–4 hours | Anti-Xa LMW heparin assay | 0.6–1.1 IU/mL, twice daily dosing 1.0–2.0 IU/mL, once daily dosing (for enoxaparin) | Partial reversal by protamine | e.g., cardiac catheterization, cardiopulmonary bypass, ECMO, etc. |
| Fondaparinux | Potential of AT inactivation of factor Xa | 17–21 hours | None routinely. Anti-Xa fondaparinux assay for renal impairment. | 0.3–0.6 mg/L | None | Monitoring of LMW heparin concentration is not routinely required, but is indicated in renal impairment, morbid obesity or low body weight, prolonged therapy, or young age. |
| | | | | | | Monitoring of fondaparinux concentration is not routinely required, but is indicated in renal impairment. |

(Continued)

Table 10.1. (Continued)

| Anticoagulant | Mode of Action | Plasma Half-life | Monitoring Test(s) | Therapeutic Range | Reversal | Notes |
|----------------------|--|-------------------------|---|---|-----------------|--|
| Argatroban | Inhibition of thrombin by reversibly binding to its active catalytic site | 24 minutes | PTT | 1.5–3.0 times baseline, not to exceed 100 seconds | None | Therapeutic targets may not be reliable for all PTT assays due to the highly variable dose-response. |
| Bivalirudin | Inhibition of thrombin by reversibly binding to its active catalytic site and substrate-binding site | 25 minutes | ACT (PCI procedures) None routinely; ACT for renal impairment | >300 seconds Procedure-specific | None | Dosage is weight-based. Monitoring is not routinely required, but patients with renal impairment should be monitored with the ACT with a therapeutic range determined by clinical protocol. |

(Continued)

Table 10.1. (Continued)

| | | | | | | |
|--|--|---------------|------|------------------------------|------|--|
| Lepirudin | Inhibition of thrombin by irreversibly binding to its active catalytic site and substrate-binding site | 40–45 minutes | PTT | 1.5–2.5 times reference mean | None | Therapeutic targets may not be reliable for all PTT assays due to the highly variable dose-response. |
| Drotrecogin alfa (activated protein C) | Degradation of factors Va and VIIIa | 30 minutes | None | None | None | |

This table identifies tests for monitoring the anticoagulant effect only. Other testing required to monitor anticoagulation therapy in general, including hemoglobin or hematocrit, platelet count, occult blood, creatinine, etc., are not specifically listed.

Abbreviations: ACT = activated clotting time; PTT = activated partial thromboplastin time; AT = antithrombin; INR = International Normalized Ratio; LMW = low molecular weight; IV = intravenous; SC = subcutaneous.

the vitamin K pathway. Clotting factors II, VII, IX, and X, and the antithrombotic proteins C and S are synthesized in the liver as inactive proteins. Vitamin K-H₂, reduced vitamin K, is an essential cofactor in the post-translational γ -carboxylation of 10–12 glutamate residues on these proteins, creating calcium-binding sites required for activity. In this process, vitamin K-H₂ is oxidized to vitamin K epoxide. In a two-step process catalyzed by the enzymes vitamin K epoxide reductase and vitamin K reductase, vitamin K-H₂ is regenerated and becomes ready to participate in another γ -carboxylation. The vitamin K cycle and its inhibition by warfarin are illustrated in Figure 10.1.

Warfarin inhibits both enzymes, decreasing the availability of vitamin K-H₂ and decreasing the number of calcium-binding sites on newly synthesized clotting factors. Active clotting factors are not affected by warfarin, but as they are metabolized, hypofunctional proteins replace them. Thus the onset of warfarin's effect is related to the half-life of the vitamin-K-dependent clotting factors, shown in Table 10.2. A partial effect is achieved quickly with the short half-life of factor VII, but the full onset of anticoagulation takes several days because of factor II's long half-life.

Rationale for Laboratory Monitoring

The therapeutic effectiveness of warfarin is well established, but therapeutic monitoring is required for several reasons¹. First, warfarin has a relatively narrow therapeutic window. Under-anticoagulation greatly reduces warfarin's therapeutic efficacy, while over-anticoagulation greatly increases the risk of bleeding. Severe bleeding episodes may be fatal or lead to severe morbidity. Second, the dose-response between individuals is highly variable and may be quite variable in the same

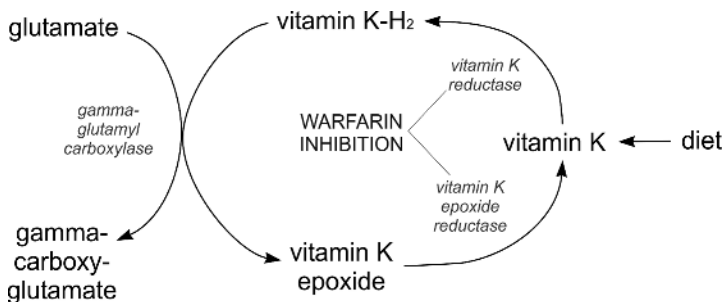


Figure 10.1. Vitamin K cycle and warfarin inhibition of key enzymes.

Table 10.2. Half-Lives of Vitamin-K-Dependent Proteins

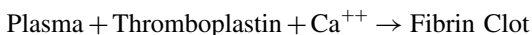
| Protein | Half-life, h |
|----------------|---------------------|
| Factor II | 60 |
| Factor VII | 4–6 |
| Factor IX | 24 |
| Factor X | 48–72 |
| Protein C | 8 |
| Protein S | 30 |

Abbreviations: h = hours.

individual over time. Third, warfarin's effect may be either potentiated or inhibited by a large number of medications. For example, potentiators include acetaminophen, erythromycin, miconazole, propranolol, and cimetidine, and inhibitors include barbiturates, prednisone, carbamazepine, nafcillin, and cholestyramine, to name a few. Fourth, warfarin's effect is influenced by fluctuations in dietary vitamin K intake. Dietary vitamin K is obtained principally from green leafy vegetables, olive oil, soybean oil, cottonseed oil, and canola oil and to a lesser extent from butter, margarine, liver, milk, ground beef, coffee, and pears. Multivitamins and herbal remedies are additional sources of vitamin K. The half-life of vitamin K is only about 1½ days, so continual intake is required, and changes in vitamin K intake affect the anticoagulant activity of warfarin within days. Fifth, coexisting diseases or illnesses may affect the absorption and metabolism of warfarin and vitamin K and the synthesis of clotting factors. Sixth, as with many medications, patient compliance is an issue with warfarin and this affects the stability of the anticoagulant effect. All these factors combine to make laboratory monitoring vital for the safe management of warfarin therapy.

PT and INR Basics

The prothrombin time (PT) test is the assay most commonly used to monitor warfarin therapy. The PT measures the time required for the following reaction:



Thromboplastins are reagent preparations rich in tissue factor and phospholipids. Thromboplastins stimulate clot formation via the

extrinsic and common pathways, involving factors VII, X, V, and II, and fibrinogen. Many different thromboplastins are commercially available.

A serious limitation of the PT is that results vary considerably with different analyzers and thromboplastins and even different lots of the same thromboplastin. Consequently, PT results are inconsistent between laboratories and even within laboratories over time, making the PT inadequate for establishing therapeutic ranges and monitoring warfarin therapy, particularly when more than one laboratory is involved.

The International Normalized Ratio (INR) was developed to standardize PT values via the mathematical transformation

$$INR = \left(\frac{PT}{MNPT} \right)^{ISI},$$

where PT is the prothrombin time, MNPT is the mean normal PT, and ISI is the International Sensitivity Index of the thromboplastin. The objective of the INR is to translate any given PT value into the PT ratio (i.e., PT/MNPT) that would have been obtained had the PT been performed by the *reference method*, which consists of PT testing using the tilt-tube method with an international reference thromboplastin preparation of the World Health Organization (WHO)³.

Let's further examine the elements of the INR transformation. The MNPT is the geometric mean of PT determinations from a set of reference subjects (i.e., normal subjects) using the laboratory's analyzer and thromboplastin combination, hereafter referred to as the *working method*. Refer to Chapter 4, Validation of Coagulation Assays, Instruments and Reagents for additional information on the selection of reference subjects. Given n PT values, the geometric mean is calculated by the formula

$$\text{Geometric MNPT} = (PT_1 \cdot PT_2 \cdot PT_3 \cdot \dots \cdot PT_n)^{1/n}.$$

In practice, the arithmetic mean is frequently used, calculated by the formula

$$\text{Arithmetic MNPT} = \frac{PT_1 + PT_2 + PT_3 + \dots + PT_n}{n}.$$

Although, technically speaking, using the arithmetic mean deviates from the INR schema, substituting the arithmetic MNPT for the geometric MNPT has no practical significance⁴. Dividing a PT value by the MNPT gives the PT ratio (PTR). Prior to the widespread adoption of the INR

system, the PTR was used in an effort to standardize PT results between working methods, but the lab-to-lab variation was too great.

The ISI represents the relative responsiveness or sensitivity of a working method compared to the reference method. For a given reduction in active clotting factor concentrations, the PT may be more prolonged or less prolonged, depending on the working method. Those methods that generate prolonged clotting times with relatively small reductions in clotting factor activities are considered responsive or sensitive. The sensitivity of a working method depends largely on the characteristics of the thromboplastin, but different models of analyzers with the same thromboplastin will usually exhibit some difference in sensitivity.

With warfarin therapy, the concentrations of active vitamin-K-dependent factors decrease, while the concentrations of inactive clotting factors increase. Inactive clotting factors are simply proteins prevented from undergoing post-translational γ -carboxylation of glutamate residues because of the decreased availability of vitamin K-H₂. They go by the unwieldy moniker of PIVKAs (*Proteins Induced by Vitamin K Antagonists*). Thromboplastins have differential sensitivity to concentrations of active clotting factors and PIVKAs, so comparison of methods is best made when all are at a steady state which may take up to 6 weeks.

The ISI of a working method is determined by split-sample PT testing against a reference method. According to the WHO protocol, fresh plasma samples from 20 non-warfarinized normal subjects and 60 stably warfarinized subjects are tested by both the working method and reference method. The ISI is the slope of the orthogonal regression line of the reference method log-PT on the working method log-PT, as illustrated in Figure 10.2. Thus, a working method whose sensitivity is similar to that of the reference method will have an ISI close to 1.0, but a less sensitive working method will have a higher ISI. The College of American Pathologists recommends that laboratories use thromboplastins with an ISI ≤ 1.70 and the Clinical and Laboratory Standards Institute (CLSI) recommends an ISI ≤ 1.50 ^{5,6}. The trend over the past decade has been for laboratories to adopt sensitive thromboplastins⁷.

In practice, most laboratories do not have the expertise or resources to determine their methods' ISI values according to the WHO protocol and rely instead on thromboplastin manufacturers to provide ISI values. Manufacturers typically assign ISI values to classes or models of analyzers, but not to specific analyzers.

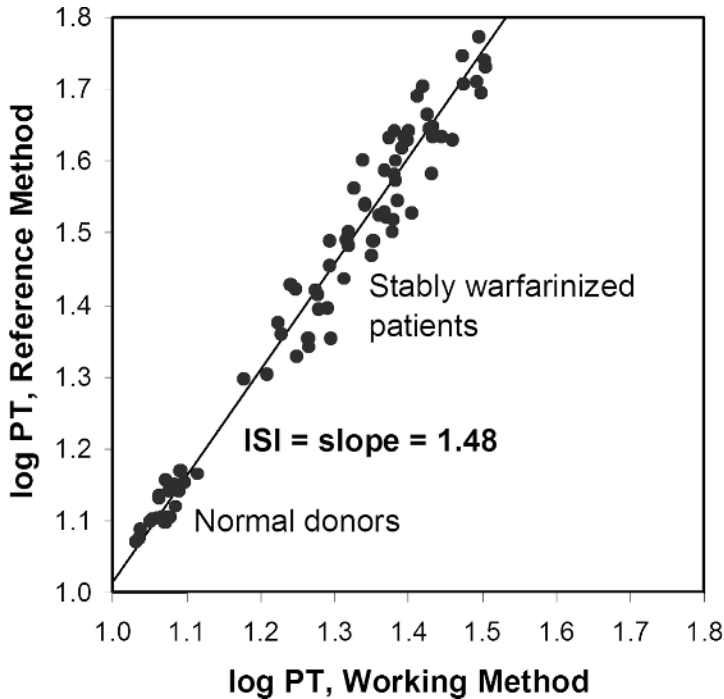


Figure 10.2. Representation of ISI assignment. The ISI is the slope of the line from the orthogonal regression of the log-PT values from the reference method on the log-PT values from the working method. Refer to the Method Comparison section of Chapter 4, Validation of Coagulation Assays, Instruments and Reagents for a discussion of orthogonal regression.

INR and Calibration

Calibration procedures are routinely used in many areas of the clinical laboratory, particularly in chemistry, immunochemistry, and hematology assays. Calibration is fundamentally the process of determining the mathematical relationship between some measurement such as absorbance or impedance and some value of interest such as an analyte concentration or cell count. Calibration equations are then used to translate measurements on patients' specimens into the values of interest.

The INR system is, in essence, a calibration of the PT where the measurement is the PT by the working method and the value of interest

is the PTR that would have been obtained with the reference method. Table 10.3 compares the INR with a typical calibration. While there is much in common, some important differences are worth noting. In a typical calibration, calibrators with assigned target values traceable to a reference or definitive method are used on each analyzer to establish the relationship between the measured responses and the target values, yielding analyzer-specific calibration equations. Additional standards with assigned target values are then used to verify the calibration and determine or verify the range of accuracy. In the INR schema, calibrators and other standards as such do not exist. The thromboplastin reagents themselves are “calibrated” against reference thromboplastins,

Table 10.3. Comparison of a Typical Calibration and the INR

| Component | Typical Calibration | INR |
|--------------------------|---|--|
| Measured response | Absorbance, impedance, luminescence, etc. | Clotting time (i.e., PT) |
| Calculated value | Concentration | INR |
| Standards | Calibrators | Thromboplastins |
| Target values | Standards traceable to definitive or reference method | Thromboplastins indexed to WHO reference thromboplastins |
| Calibration parameters | Slope and intercept, or coefficients of non-linear calibration equations | ISI and MNPT |
| Calibration “location” | Analyzer | Analyzer (MNPT) and manufacturer (ISI) |
| Range of accuracy | Determined in laboratory or determined by manufacturer and verified in laboratory | Difficult or impossible to determine in most labs |
| Calibration verification | Verified in laboratory | Difficult or impossible to do in most labs |
| Limitations | Matrix effects | Analyzer effects |

Abbreviations: INR = International Normalized Ratio; PT = prothrombin time; WHO = World Health Organization; ISI = International Sensitivity Index; MNPT = mean normal prothrombin time.

resulting in the assignment of a set of ISI values for classes or models of analyzers. Thus the calibration occurs in part on each analyzer with the determination of the MNPT and in part at the manufacturer's location with the assignment of the ISI. The absence of calibrators and standards in this schema means that the INR calibration is not analyzer-specific and that laboratories cannot readily verify the accuracy of their INRs.

It should come as little surprise that a host of publications have reported inadequacies with the INR system. Interested readers may find references of these reports in other publications^{1,8}. Problems include the incorrect assignment of class- or model-specific ISI values by manufacturers and individual analyzer effects that invalidate the class- or model-based ISI values. Nevertheless, the INR system appears to have decreased the overall lab-to-lab variation in PT values for monitoring warfarin therapy and has allowed the adoption of standardized therapeutic ranges throughout the world.

Plasma Calibrators

Due to the limitations of the calibration provided by the INR schema, described above, the past several years have seen a growing interest in analyzer-specific INR calibration in a typical manner using certified plasma calibrators. Calibrators with assigned INR values are now commercially available in many countries, though not yet in the United States, and procedures for using INR calibrators are described in CLSI document H-54⁸.

Several reports have been published of improvements in accuracy and interlaboratory variation through the use of INR calibrators⁹⁻¹⁴. However, a number of fundamental issues still need to be addressed, the most significant of which is the assignment of target INR values to calibrators. Currently, different reference thromboplastins, analytical methods, and methods of producing factor-deficient plasma may produce significantly different target INR values¹⁰. The ability to reliably determine the "true" INR of calibrators remains elusive. Additional unresolved questions involve the required precision of the calibration procedure, number of calibrators to be used, stability of calibration curves, determination of a full calibration curve versus an ISI-only calibration, and matrix effects of calibrators versus fresh patients' specimens^{10,15}.

INR calibration using plasma calibrators is appealing from both a theoretical perspective and the documented deficiencies of the INR

schema, and results of studies of calibration look promising, but as of this writing it appears prudent to maintain some level of caution due to the unresolved issues.

Evaluation of Discrepant INRs

Warfarin patients generally receive INR testing over an extended period of time and often from more than one laboratory. It is not uncommon, in the author's experience, for the laboratory director to receive inquiries about INR values that appear discrepant, either between two laboratories or from one laboratory at different times. Causes of discrepant INRs are listed in Table 10.4. Some discrepancies are caused by the laboratory, some are caused by patients, and some reflect inherent limitations of the INR schema. Not all discrepancies can be resolved, but all reports of possible discrepancies should be taken seriously and evaluated to assure that correctable issues are identified and addressed.

When evaluating an INR discrepancy, the laboratory director should encourage the patient's physician to increase the thoroughness

Table 10.4. Causes of Discrepant INR Values

| Testing | Patient |
|---|---|
| Different citrate concentrations | Changes in diet (including herbal remedies and vitamin supplements) |
| Improper specimen handling | |
| Misidentified specimen | Changes in warfarin dosage |
| Incorrect ISI assignment | Changes in patient compliance |
| Incorrect MNPT determination | Changes in medications |
| Incorrect INR equation | Variation in specimen collection time |
| Incorrect calculations | relative to warfarin ingestion |
| Incorrect entry of ISI or MNPT in computer system | Biological variation in clotting factor levels |
| Reporting errors | Genetic determinants of warfarin metabolism |
| Analytical imprecision | |
| Analyzer problems | |
| Reagent problems | |
| Lupus anticoagulant | |
| Analyzer-specific effects not corrected by INR schema | |

Abbreviations: INR = International Normalized Ratio; ISI = International Sensitivity Index; MNPT = mean normal prothrombin time.

of the dietary and medication history. One vignette will serve as illustration. A patient whose INRs had been stable for many months had a markedly elevated INR. On questioning, the patient denied any changes in diet, medications, or compliance. Finally the physician asked the patient to bring her medications to the office. The medications included warfarin tablets marked with a “5”. The physician had expected the patient to be on a dose of 2 mg/day. The patient said she had run out of her latest prescription, fished in the medicine cupboard and found some old tablets that appeared to her aged eyes to be marked with a “2” and started taking them until it was time for another appointment with her physician. Unfortunately, the number on the tablets was actually a 5, so the patient had been taking a much larger dose than necessary.

Therapeutic Monitoring

The recommended test for monitoring warfarin therapy is the INR. Use of any other test should be discouraged. Common therapeutic targets are shown in Table 10.1, although it is important to remember that these ranges may not apply to all patients¹. Guidelines for warfarin dosing, frequency of testing, and treatment of over-dosage are beyond the scope of this chapter, but the interested reader is referred to an excellent review by Ansell and colleagues¹.

A few additional points about warfarin therapeutic monitoring may be useful. First, the PT assay is sensitive to changes in factor VII levels. Because factor VII has a short half-life, the PT/INR may begin to be prolonged within hours of a warfarin dose. Some non-compliant patients have admitted to taking warfarin only the day prior to testing to make it appear to the physician that the patient is compliant with the prescribed therapy. In cases where patient compliance is suspect but the INR indicates a warfarin effect, specific factor assays showing decreased factor VII levels with normal or nearly normal factor II levels suggest that warfarin has been taken only recently. During long-term therapy at steady state, factor II and factor VII levels should be roughly equally decreased.

Second, heparin in therapeutic doses does not significantly affect the PT/INR with most working methods, since most thromboplastin reagents contain heparin neutralizers. So for patients on both heparin and warfarin, the INR reflects only the warfarin effect. The situation is different for direct thrombin inhibitors (DTIs), discussed later in this chapter. DTIs affect clot-based assays that involve thrombin, including

the PT. In patients being treated with DTIs and warfarin, the INR reflects both a warfarin effect and DTI effect. During the conversion from a DTI to warfarin, the INR therapeutic target should be increased or the dose of DTI should be decreased^{16,17}.

Third, patients with significantly supratherapeutic INRs are frequently treated with oral vitamin K to partially reverse the warfarin effect. Many physicians believe the reversal will occur within a few hours, but significant lowering of the INR may take 24 hours or longer due to the relatively long half-life of warfarin².

Unfractionated Heparin

Unfractionated heparin is an IV or subcutaneously administered anticoagulant, widely used for the treatment and prevention of thromboembolic events, including deep venous thrombosis, pulmonary embolism, stroke, myocardial infarction, unstable angina, and some cases of disseminated intravascular coagulation (DIC), and for anticoagulation during cardiopulmonary bypass, percutaneous coronary intervention, and extracorporeal membrane oxygenation (ECMO) procedures¹⁸.

Mechanism of Action

UFH is a heterogeneous group of anionic mucopolysaccharides, called glycosaminoglycans, with anticoagulant properties. The molecular weight ranges from 3,000 to 30,000 daltons, averaging 15,000–18,000 daltons, around 45–50 saccharides. It carries the name *heparin* because it was originally extracted from liver¹⁸. Pharmaceutical preparations are usually derived from porcine intestinal mucosa or bovine lung. The term “heparin” generally refers to unfractionated heparin, so named because of its heterogeneity in size and function. In this chapter, the abbreviation UFH will be used for unfractionated heparin and LMWH for low-molecular weight heparin (discussed later).

Heparin, both UFH and LMWH, exerts its anticoagulant effect via several mechanisms, but the most important by far is the potentiation of the serine protease inhibitor antithrombin (AT), formerly called antithrombin III. By itself, AT is a slow inhibitor of thrombin. Heparin binds to AT through a specific pentasaccharide sequence, inducing a conformational change that increases the rate of binding to thrombin by 1000-fold. This relationship is illustrated in Figure 10.3. Heparin-AT also inhibits factor Xa, factor IXa, factor VIIa-tissue factor

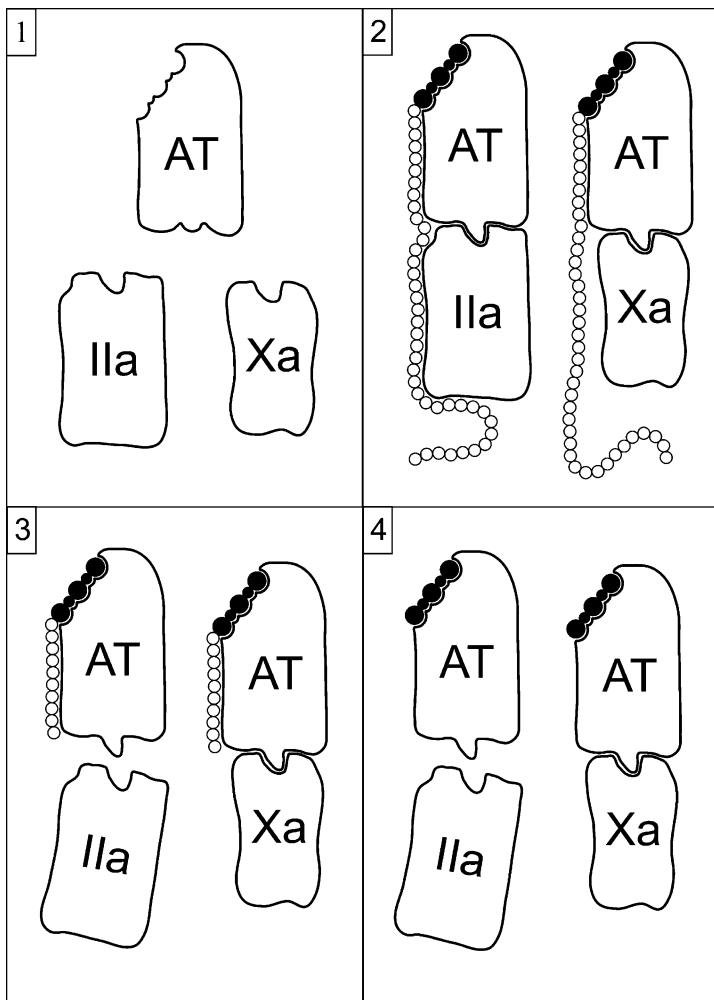


Figure 10.3. Schematic representation of the potentiation of AT by heparin and its derivatives. Panel 1 illustrates the lack of activity of AT in its native state against factor IIa (thrombin) or factor Xa. Panel 2 shows the interaction and activity of UFH and AT. The binding of UFH to AT occurs through a specific high-affinity pentasaccharide sequence (shaded), inducing a conformational change in AT that markedly enhances its activity. The long saccharide chain of UFH is needed for anti-IIa but not anti-Xa activity. Panel 3 shows the interaction and activity of LMWH and AT. The binding of LMWH to AT occurs via the high-affinity pentasaccharide. AT-LMWH complexes in which

complex, factor XIa, factor XIIa, and plasma kallikrein. It indirectly inhibits thrombin-induced activation of platelets and factors V and VIII. Thrombin is 10-fold more sensitive to heparin-AT inhibition than is factor Xa, and factor Xa is more sensitive than the other coagulation factors. AT binds covalently to the active serine centers of coagulation factors, then heparin dissociates and can be reutilized¹⁹.

In pharmaceutical preparations, only one-third of UFH molecules possess the high-affinity pentasaccharide required for AT potentiation. The remaining two-thirds have minimal anticoagulant activity at therapeutic levels. Inhibition of thrombin by heparin-AT requires heparin molecules of 18 or more saccharides. Complexes with small heparin molecules inhibit factor Xa and other factors, but not thrombin.

Several other minor mechanisms contribute to the anticoagulant effect of UFH. At high concentrations, UFH molecules of 24 or more saccharides, with or without the high-affinity pentasaccharide, bind to heparin cofactor II and catalyze the inactivation of thrombin. The non-specific binding of heparin to platelets inhibits platelet function (usually) and contributes to the hemorrhagic effects of heparin therapy. Platelet binding is more pronounced with large heparin molecules than with small heparin molecules. Heparin induces the secretion of tissue factor pathway inhibitor (TFPI) by endothelial cells, which reduces the procoagulant activity of the factor VIIa-tissue factor complex. Finally, heparin binds to von Willebrand factor (vWF), inhibiting platelet adhesion mediated by vWF¹⁸.

Rationale for Laboratory Monitoring

Heparin is a highly effective anticoagulant when present in an adequate concentration. Laboratory monitoring is required for a number of reasons. First, UFH has a relatively narrow therapeutic window. When used for treatment, inadequate concentrations fail to produce the intended therapeutic response with increased risk of recurrence or extension of thromboembolism, while excessive concentrations yield

Figure 10.3. (Continued) the LWMH molecules have fewer than 18 saccharides lose anti-IIa activity but retain anti-Xa activity. LMWH molecules of 18 or more saccharides retain anti-IIa activity. Panel 4 shows the interaction and activity of fondaparinux and AT. Fondaparinux is a synthetic version of the high-affinity pentasaccharide. Fondaparinux-AT complexes do not have anti-IIa activity.

an increased risk of hemorrhagic complications. When used for anticoagulation for invasive procedures, failure to achieve an adequate level of anticoagulation may lead to procedure failure, clotting in extracorporeal circuits, or embolic events. Second, the dose-response is highly unpredictable between patients and even in the same patient over time. Non-specific binding of heparin to endothelial cells, macrophages, platelets, and a variety of plasma proteins makes the dose-response highly variable. Even with weight-based loading doses and infusion rates, therapeutic levels of anticoagulation are difficult to achieve and maintain. Third, the rate of heparin resistance is high. Heparin resistance, defined as a requirement for $>35,000$ U UFH per 24 hours to achieve a therapeutic level, occurs in up to 25% of patients treated with UFH. Heparin resistance is classically caused by AT deficiency, particularly with AT levels $<25\%$, but is also associated with increases of factor VIII, fibrinogen, platelet factor 4, and heparin-binding proteins. Many of these proteins are acute-phase reactants whose concentrations rise during the acute episodes that create the need for heparin therapy.

Heparin-induced Thrombocytopenia

One complication of UFH deserves special mention in the context of laboratory monitoring. Heparin-induced thrombocytopenia (HIT) is a serious, potentially life-threatening complication that requires routine monitoring of platelet counts during UFH therapy. HIT is defined as an antibody-mediated decrease in platelet count of 50% or more from baseline. HIT is associated with a 20–75% risk of venous or arterial thromboembolic events including deep venous thrombosis (DVT), pulmonary embolism (PE), myocardial infarction (MI), stroke, occlusion of extremities or digits, and so forth. It should be noted that HIT is *not* the non-immune-mediated, transient, modest drop in platelet counts commonly observed with heparin therapy, although this phenomenon has also been called HIT or Type I HIT.

During UFH therapy, some individuals develop antibodies to the heparin-platelet factor 4 (PF4) complex. Although the mechanism of thrombus formation in HIT has not been fully elucidated, research suggests that hypercoagulability involves the following: 1) platelet activation with release of procoagulant microparticles, due to heparin-PF4-IgG immune complexes' occupancy and crosslinking of platelet Fc receptors; 2) tissue factor expression on endothelial cells activated

by the antibodies' recognition of PF4 bound to endothelial heparin sulfate; and 3) tissue factor expression by monocytes activated by HIT antibodies²⁰.

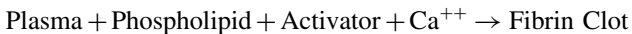
HIT is typically recognized by platelet count monitoring, but sometimes thrombotic complications are the first clue. For individuals who have not previously received heparin, HIT occurs on day 5 or later. For those who have previously received heparin, particularly within the prior 100 days, HIT may occur within 24 hours ("rapid-onset HIT"). In a small number of HIT patients, the onset of thrombocytopenia begins several days after heparin has been stopped ("delayed-onset HIT"). The risk of HIT varies among patient groups. Postoperative patients have a higher risk than medical patients who have a higher risk than obstetric patients. Cardiac surgical patients are more likely to develop HIT than orthopedic surgical patients, but when orthopedic surgical patients develop the antibody, they are more likely to have a thrombotic event.

Two types of assays are used to detect the presence of HIT antibodies (i.e., anti-heparin-PF4 antibodies). Activation assays are based on the ability of serum or plasma HIT antibodies to activate platelets in the presence of heparin. One or more measures of platelet activation are used, particularly serotonin release and platelet aggregation. These tests are non-standardized and vary widely in their accuracy. Variability between donor platelets and variability between heparin preparations are confounding factors, and alterations in the concentrations of platelets, heparin, and patient serum may produce different results. Nevertheless, in experienced hands the activation assays are more accurate than antigenic assays, with sensitivity as high as 98%. Antigenic assays are based on the detection of antibodies that bind to heparin-PF4 complexes or to polyvinylsulfonate-heparin complexes. Antigenic assays are more standardized, technically easier, and have more rapid turnaround time than activation assays. Antigenic assays detect clinically insignificant antibodies more frequently than do activation assays, and antigenic assays may miss antibodies formed against proteins other than PF4, but the overall sensitivity of antigenic assays is above 90%. Both types of assays have specificities above 95% when the platelet count falls within 4 days of initiation of UFH therapy. When the platelet count falls later, the specificity is 80–97% for activation assays and 50–93% for antigenic assays.

For more information about HIT, the interested reader is referred to an excellent review by Warkentin and Greinacher²⁰.

PTT Basics

The activated partial thromboplastin time (PTT) test is the assay most commonly used to monitor heparin therapy. The PTT measures the time required for the following reaction:



Clot formation occurs via the intrinsic and common pathways, involving high molecular weight kininogen, prekallikrein, factors XII, XI, IX, VIII, X, V, and II, and fibrinogen. Many different phospholipid and activator reagents are available commercially²¹.

The PTT is prolonged by IV heparin and has been used successfully in the management of heparin therapy for many decades. However, as with PT results for monitoring warfarin therapy, PTT results vary widely with different analyzers and reagents and are inconsistent between laboratories. PTT values may vary considerably even with different reagents or different lots of reagent from the same manufacturer. Unlike with the PT assay, there is no INR equivalent with the PTT for standardizing results and establishing universal therapeutic ranges. Each laboratory must establish therapeutic range limits for its working method and must verify or adjust the limits whenever a reagent change is made. See the discussion below for determination of therapeutic ranges.

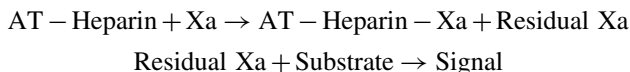
The popularity of the PTT for monitoring UFH therapy is based on several factors. First, there is a logical appeal in a test that provides a physiologic measurement of the anticoagulant effect rather than simply providing a heparin concentration. Second, the PTT is widely available with relatively short turnaround time, has good reproducibility, and is inexpensive. Third, clinicians have decades of experience with the PTT and have achieved a comfort level with its use for UFH monitoring. Fourth, until recently there have not been good alternatives. These characteristics have combined to provide a general level of clinical satisfaction with the PTT assay.

The PTT also possesses some limitations and drawbacks for monitoring UFH therapy^{18,22}. First, the PTT is a non-standard assay, as described above, so therapeutic ranges must be established in each laboratory. Different types of heparin may produce different PTT response curves. Second, the PTT is useful only for monitoring therapeutic IV doses or high-dose subcutaneous UFH therapy. The PTT is not useful for standard-dose subcutaneous therapy, due to limited PTT response, or for the high-dose IV UFH required for cardiac catheterization or other procedures, due to an excessive PTT response. Third,

the PTT is affected by variables other than UFH, including increased concentrations of factor VIII and fibrinogen, decreased concentrations of AT or intrinsic and common pathway proteins, lupus anticoagulant, and thrombolytic agents. When lupus anticoagulants are present, the PTT becomes unpredictable for assessing UFH response and should not be used. The anti-Xa assay, discussed below, is the test of choice for patients with lupus anticoagulants. Fourth, specimens have relatively short stability, particularly when heparin is present. See Chapter 2, Collection of Coagulation Specimens for additional discussion of specimens. However, the advantages of the PTT are perceived as outweighing the disadvantages for most patients, so the PTT assay continues to enjoy widespread use for monitoring IV UFH therapy.

Anti-Xa Heparin Assay Basics

The heparin anti-Xa assay, or factor Xa inhibition test, is increasingly used to supplement or replace the PTT for monitoring heparin therapy. The principle of the anti-Xa assay is the inhibition of factor Xa by AT-heparin complexes, as illustrated in the following reactions²²:



In these reactions, the AT-heparin complex is provided by the test plasma and factor Xa is provided as a defined quantity of reagent. The AT-heparin complexes inhibit factor Xa essentially in a 1:1 ratio. Residual factor Xa is available to react with a substrate to form a signal, usually a chromogen, whose strength is inversely proportional to the heparin concentration. The signal is compared against a standard curve to yield a quantitative factor Xa inhibitory effect in IU/mL. The standard curve is dependent on the heparin preparation used. Different curves need to be established for UFH and LMWH.

The anti-Xa assay has several favorable characteristics. First, the anti-Xa assay is simple to perform and automatable on many coagulation analyzers. Second, citrated specimens used for other coagulation tests may be used without additional special handling. Third, the test is not affected by coagulation factor concentrations, lupus anticoagulants, or other biological variables. Fourth, the anti-Xa assay is sensitive to the anticoagulant effect of LMWH and fondaparinux, discussed below, making it more versatile than the PTT for anticoagulant monitoring.

Fifth, the anti-Xa assay can theoretically be standardized to provide consistency between laboratories and facilitate the adoption of common therapeutic ranges.

Anti-Xa assays have some limitations and drawbacks. First, the anti-Xa assay is more expensive than the PTT. Second, it may not be financially or technically feasible to offer the anti-Xa assay in small laboratories, depending on test volume and instrumentation. Third, different UFH preparations may yield different standard curves. It is generally not feasible to establish a unique standard curve for every UFH preparation, so the potential exists for the standard curve of an anti-Xa assay to not be representative of the anticoagulant response for some UFH preparations. The same limitation applies to LWMH preparations.

Two variations of the anti-Xa assay deserve mentioning. The first variation is a clot-based rather than chromogenic assay. Factor V, fibrinogen, activator, and calcium serve as the “substrate” and the clotting time serves as the “signal.” The clotting time is proportional to the heparin concentration. Clot-based assays tend to be less precise than chromogenic assays and may be affected by more biological variables. The second variation involves an initial step of adding excess AT to the test specimens in a quantity that assures all heparin becomes complexed with AT. The theoretical appeal of this approach is that it detects “free” heparin and provides a truer estimate of heparin concentration. The alternate view is that the value of interest is the anticoagulant effect, not the heparin concentration. In AT deficiency, the anticoagulant effect may be muted physiologically due to a decrease in AT-heparin complexes, but the anti-Xa assay results may appear to show a therapeutic response due to the inclusion of “free” heparin. Currently, there is no consensus of the use of AT-supplemented versus non-supplemented anti-Xa assays.

ACT Basics

The activated clotting time (ACT) is a whole-blood clotting test commonly used to monitor high-dose UFH therapy, particularly cardiac catheterization and cardiopulmonary bypass procedures. The ACT consists of the following reaction²²:

Whole Blood + Particulate Activator → Fibrin Clot

Celite and kaolin are commonly used as activators. Typically, the ACT is a point-of-care test because non-anticoagulated blood specimens cannot be transported to a central laboratory quickly enough to avoid pre-test activation of clotting.

The ACT offers several advantages for UFH monitoring. First, the ACT has a wide dose-response range so it can be used for assessing high-dose UFH therapy. Second, as a point-of-care test, the ACT is simple to perform and results are available with rapid turnaround time. The rapid turnaround facilitates its use in assessing both induction and reversal of UFH anticoagulation in near-real-time during procedures. Third, extensive clinical experience over several decades provides a high level of familiarity and comfort with its use. Fourth, suitable alternatives are not readily available. Consequently, the ACT is widely used for high-dose UFH management.

The ACT also has several important limitations and drawbacks. First, the ACT is non-standard and imprecise. The inter-analyzer variance is generally large, even with analyzers of the same make and model using the same lot of activator. Even duplicate simultaneous results from the same analyzer may vary considerably²³. Second, because non-anticoagulated blood is used, results are affected by deficiencies in specimen collection and application technique, including delayed application of specimens. Third, the ACT reaction follows the same pathways as the PTT reaction, so the ACT is affected by the same biological variables. In addition, because whole blood is used, the ACT is also affected by variations in platelet count inasmuch as platelets provide phospholipid surfaces for reactions of the intrinsic and common pathways. By contrast, in the PTT assay phospholipid is provided as a reagent in a controlled amount. Fourth, although the dose-response of the ACT extends into the standard-dose UFH range, it is less precise and offers no advantages over the PTT for this use. Fifth, many ACT analyzers have only basic functionality and cannot be interfaced with information systems for data management.

A few issues concerning ACT testing should be considered. First, many different activator reagents are available commercially, the most common being celite, kaolin, and combinations of celite and kaolin. Different reagents generate different ACT values. One of the more important differences is the effect of aprotinin, a serine protease inhibitor commonly used in coronary artery bypass graft surgery. Celite is considerably more responsive to aprotinin than is kaolin. Controversy remains about the value of including or excluding the aprotinin effect

in ACT monitoring, and depending on one's view, either celite or kaolin becomes the preferred activator^{22,24}. Second, despite the ACT's high imprecision and poor inter-analyzer concordance, it has a well-established role in the management of high-dose UFH therapy. The ACT's utility in this setting is due to the fact that the goal of UFH dosing is not to hit some precise target but rather to assure that the anticoagulation level is above some threshold. The ACT's coarse estimate of clotting time is sufficient in this regard, but likely would not be if more precise dosing were required. Finally, while most ACT methods detect formation of a clot, one method (i-STAT, Abbott Laboratories, Abbott Park, Illinois) detects thrombin generation rather than clot formation. This method has been shown to correlate reasonably well (within the context of a very imprecise test) with the widely used Hemochron ACT (ITC, Edison, New Jersey)²⁵. However, one potential source of major discrepancies is fibrinogen deficiency or dysfibrinogenemia because fibrinogen is measured in the clot-based ACT's pathway, but not the i-STAT ACT's pathway.

Therapeutic Monitoring

The selection of tests and therapeutic ranges for UFH monitoring are listed in Table 10.1. For information about UFH administration and frequency of testing, the interested reader is referred to an excellent review by Hirsh and colleagues¹⁸.

The therapeutic range of the anti-Xa UFH assay for standard IV UFH therapy is 0.3 to 0.7 IU/mL¹⁸, and this serves as the basis for determining the PTT therapeutic range. The PTT therapeutic range should be established for each analyzer/reagent combination by determining from regression analysis the PTT values that correlate with the anti-Xa UFH assay's therapeutic range limits. Where feasible, a minimum of 30 fresh patient specimens should be used rather than frozen aliquots or UFH-spiked samples¹⁸. The PTT response to *in vitro* heparin spiking overestimates the anti-Xa levels in patients on UFH therapy²⁶, so patients' specimens are considered the gold standard for determining the PTT therapeutic range. One note of caution is that the correlation of anti-Xa and PTT values is low, so there is large uncertainty in the estimates of the PTT therapeutic range limits; however, no better alternative currently exists.

Once the PTT therapeutic range has been established, the implementation of a new PTT reagent or reagent lot does not necessarily

require that a new therapeutic range be determined, provided that the heparin response does not change significantly. In the appendix of an article²², Olson and colleagues described a method for comparing the responsiveness of PTT reagents with fresh patient specimens and monitoring the cumulative difference of means. As long as the cumulative difference remains within $+/-5$ seconds, the PTT therapeutic range does not need to be reestablished.

Some have questioned whether the PTT therapeutic range needs to be verified or established with each lot of UFH used therapeutically. In addition to being impractical, there is evidence suggesting it is unnecessary²⁷.

Historical therapeutic ranges have been superseded by ranges corresponding to anti-Xa UFH assay values. Fixed ranges such as 1.5 to 2.5 times the PTT reference mean or 1.5 to 2.5 times the patient's baseline are inappropriate because of the wide variation in dose-response among PTT methods, and the formerly widely used heparin therapeutic range of 0.2–0.4 U/mL was based on protamine titration assays and is now outdated.

Target ACT ranges are established by clinical protocols for each type of procedure.

Low-Molecular-Weight Heparin

Low-molecular-weight heparin (LMWH) is a subcutaneously administered anticoagulant, used for the treatment and prevention of thromboembolic events, including deep venous thrombosis, pulmonary embolism, stroke, myocardial infarction, and unstable angina^{18,28,29}.

Mechanism of Action

LMWH is prepared from UFH by chemical or enzymatic depolymerization. The molecular weight ranges from 1500 to 10,000 daltons, averaging 4000–5000 daltons, around 15 saccharides¹⁸. As does UFH, LMWH exerts its principal anticoagulant effect by the potentiation of AT via binding of a specific pentasaccharide sequence present on only one-third of LMWH molecules. A comparison of LMWH with UFH is given in Table 10.5. The differences between LMWH and UFH are virtually all attributable to the different binding properties of the shorter saccharide chains. Significant differences include less non-specific binding to plasma proteins, endothelial cells, macrophages, and

Table 10.5. Comparison of Unfractionated Heparin, Low-Molecular-Weight Heparin, and Fondaparinux

| Property | UFH | LMWH | Fondaparinux |
|---|---|--|--|
| Mode of action | Activation of AT | Activation of AT | Activation of AT |
| Molecular weight, daltons | 3000–30,000 | 1500–10,000 | 1728 |
| Molecules containing high-affinity pentasaccharide, % | 33 | 33 | 100 |
| Administration | IV or SC | SC | SC |
| Plasma half-life, h | 0.5–2.5 | 3–4 | 17–21 |
| Binding to plasma proteins, endothelial cells, platelets, and macrophages | Much | Little | None |
| Dose-response | Unpredictable | Predictable | Predictable |
| Dosage | Titrated to lab value | Fixed dose or weight-based | Fixed dose or weight-based |
| Anti-Xa/anti-IIa ratio | 1.2 | 2–4 | No anti-IIa activity |
| PTT at therapeutic levels | Prolonged | Not prolonged or only slightly prolonged | Not prolonged or only slightly prolonged |
| Thrombocytopenia (<50,000–100,000/ μ L, incidence, %) | 1.0–1.5 | 1.0–1.5 | 0 |
| Thrombocytopenia (<50,000/ μ L, incidence, %) | 0.2 | 0.1 | 0 |
| HIT (immune-mediated thrombocytopenia), % | 0.1–5 | < 0.1–1 | None |
| Bleeding complications, % | 3–5 | Same or lower | Same or higher |
| Protamine reversal | Effective for IV, less effective for SC | Less effective | Not effective |

Abbreviations: AT = antithrombin; UFH = unfractionated heparin; LMWH = low-molecular-weight heparin; IV = intravenous; SC = subcutaneous; PTT = partial thromboplastin time; HIT = heparin-induced thrombocytopenia.

platelets, resulting in a much more predictable dose-response; longer plasma half-life, allowing once- or twice-daily dosing; and reduced anti-IIa activity, of uncertain significance. In addition, reduced binding to platelets is believed to be responsible for a lower incidence of HIT compared to UFH.

Rationale for Laboratory Monitoring

Routine laboratory monitoring of the anticoagulant effect of LMWH is not required for most patients because of the highly predictable dose-response. Monitoring is indicated in some special situations, including renal failure, morbid obesity, low body weight, newborns, young children, and patients receiving prolonged therapy^{18,28}. As with UFH, HIT occurs with LMWH therapy, although at a much lower frequency, so routine monitoring of platelet counts may be required only for surgical patients or for medical patients who received UFH prior to LMWH²⁰.

Therapeutic Monitoring

When monitoring is required, the test of choice is the anti-Xa assay. Because each brand of LMWH has unique properties, the anti-Xa LMWH assay's standard curve should be established with LMWH preparations used in the facilities served by the laboratory. A standard curve established with one brand of LMWH should not be considered valid for other brands. The therapeutic targets of anti-Xa activity measured 4 hours after administration of enoxaparin, the best-studied LMWH preparation, are shown in Table 10.1.

Fondaparinux

Fondaparinux is a subcutaneously administered synthetic analog of the AT-binding pentasaccharide found in active molecules of UFH and LMWH, used in the treatment of acute venous thromboembolism and prevention of thromboembolism^{30,31}.

Mechanism of Action

Fondaparinux exerts its anticoagulant effect by the potentiation of AT. A comparison of fondaparinux with LMWH and UFH is shown in

Table 10.5. Significant properties of fondaparinux include an absence of anti-IIa activity because it is too small to bridge AT to thrombin; no detectable binding to plasma proteins or cells, providing a highly predictable dose-response with excellent bioavailability following SC injection; and a longer half-life than either UFH or LMWH, 17–21 hours, allowing once-daily dosing. In addition, fondaparinux has not been associated with HIT, cannot be reversed by protamine sulfate, and is not effective in patients with moderate or severe AT deficiency. Fondaparinux is excreted unchanged in the urine, so renal insufficiency may be a contraindication to its use.

Rationale for Laboratory Monitoring

Routine laboratory monitoring of the anticoagulant effect of fondaparinux is not required for most patients because of the highly predictable dose-response. Monitoring is indicated in renal impairment. Although clinical experience with fondaparinux is relatively limited as of this writing and indications for laboratory monitoring have not been fully elucidated, it seems prudent to consider monitoring in extremes of body weight, young age, and prolonged use.

Therapeutic Monitoring

When monitoring is desirable, the test of choice is the anti-Xa assay whose standard curve is established using fondaparinux. The therapeutic target is shown in Table 10.1. It should be noted that the anti-Xa fondaparinux assay does not have clinical validation at this point in time.

Lepirudin

Lepirudin is an IV-administered direct thrombin inhibitor (DTI) used in patients with HIT in the treatment of thromboembolism and in cardiopulmonary bypass procedures^{17,28,30}. Hirudin has been studied in a variety of additional uses and appears at least as effective as UFH³⁰.

Mechanism of Action

Hirudin is a naturally occurring polypeptide of 65 amino acids, produced in trace amounts by the leech *Hirudo medicinalis*, with thrombin-inhibiting activity. Lepirudin is a recombinant hirudin, derived from

yeast cells. Lepirudin differs from hirudin by one amino acid substitution and the absence of a sulfate group on another amino acid. Lepirudin has only 1/10th of hirudin's affinity for thrombin. Lepirudin binds irreversibly to thrombin's active catalytic and substrate-binding sites. It inhibits both free and clot-bound thrombin. It has a predictable dose-response and its efficacy is unaltered by AT deficiency. No antidotes are available and bleeding complications are hard to treat, but this situation is mitigated to some degree by its short half-life of 40–45 minutes. Lepirudin is excreted in the urine, so its use should be limited in renal insufficiency. Reactivity with heparin-induced antibodies has not been reported, but allergic reactions with cough, bronchospasm, stridor, or dyspnea are relatively common, occurring or suspected in 10% of recipients. Hirudin may be more effective than UFH in preventing extension of thrombi³⁰.

Rationale for Laboratory Monitoring

Lepirudin has a relatively narrow therapeutic window. Overdosage is of particular concern because of the absence of an antidote.

Therapeutic Monitoring

The PTT is used for routine laboratory monitoring with a manufacturer-recommended therapeutic range of 1.5–2.5 times the reference mean¹⁷. The therapeutic range was determined in clinical trials, but because of the wide variability of PTT assays, the transferability of this range to all laboratories seems questionable. See the PTT section above for other considerations of the PTT for therapeutic monitoring.

Lepirudin increases the INR, and this effect can complicate the conversion to warfarin therapy. Lepirudin's manufacturer recommends that before initiating warfarin therapy the PTT ratio be reduced to just above 1.5, then standard INR therapeutic ranges be used for decisions about warfarin dosing and discontinuation of lepirudin¹⁷.

Bivalirudin

Bivalirudin is an IV-administered DTI used for anticoagulation of patients with unstable angina undergoing percutaneous transluminal coronary angioplasty (PTCA) and receiving concomitant aspirin³².

Other uses have been studied showing bivalirudin to have efficacy similar to or better than UFH or lepirudin³⁰.

Mechanism of Action

Bivalirudin, like lepirudin, is a derivative of hirudin. Bivalirudin is a synthetic combination of 8- and 12-peptide fragments of hirudin that bind to thrombin's active catalytic site and substrate-binding sites, respectively. Thrombin slowly cleaves bivalirudin with recovery of thrombin's active-site function. Bivalirudin has a predictable dose-response and a short half-life of only 25 minutes. Its efficacy is unaltered by AT deficiency. Bivalirudin is not extensively excreted in the urine. Bivalirudin has not been shown to react with heparin-induced antibodies. No antidotes are available, but the short half-life and reversible binding limit bleeding complications. In PTCA, the 4% risk of a major bleed with bivalirudin is less than half the risk with UFH³⁰.

Therapeutic Monitoring

Because of the predictable dose-response and short half-life, routine laboratory monitoring of bivalirudin is not required.

Argatroban

Argatroban is an IV-administered DTI used in patients with HIT in the treatment or prophylaxis of thromboembolism and in percutaneous coronary intervention procedures^{16,28,30}.

Mechanism of Action

Argatroban, a synthetic derivative of L-arginine, is a competitive inhibitor of thrombin that binds reversibly to thrombin's active catalytic site. It inhibits both clot-bound and free thrombin. It has a predictable dose-response and its efficacy is unaltered by AT deficiency. No antidotes are available and bleeding complications may be hard to treat, but the short plasma half-life of 24 minutes somewhat mitigates this disadvantage²⁸. Argatroban is metabolized primarily in the liver, so its use should be limited in hepatic impairment. Argatroban does not react with heparin-induced antibodies^{16,28}.

Rationale for Laboratory Monitoring

Argatroban has a relatively narrow therapeutic window. Overdosage is of particular concern because of the absence of an antidote.

Therapeutic Monitoring

The PTT is used for routine laboratory monitoring with a manufacturer-recommended therapeutic range of 1.5–3.0 times the patient's baseline PTT, but < 100 seconds¹⁶. Although this therapeutic range was determined in clinical trials, its transferability to all laboratories seems doubtful given the wide inter-laboratory variation of the PTT. See the PTT section above for other considerations of the PTT for therapeutic monitoring.

Drotrecogin Alfa (Activated Protein C)

Drotrecogin alfa is an IV-administered anticoagulant, used in the treatment of severe sepsis^{30,33}.

Mechanism of Action

Drotrecogin alfa is a recombinant form of human activated protein C (APC), with the same amino acid sequence. Drotrecogin alfa exerts its anticoagulant effect by inhibiting factors Va and VIIIa. Drotrecogin alfa is inactivated by endogenous plasma protease inhibitors and has a mean half-life of 30 minutes³⁴. Compared to placebo, drotrecogin alfa reduces mortality by 19% in a subset of patients with severe sepsis³⁰. Bleeding is the most serious adverse event, and no known antidote is available. The short half-life helps in the management of drotrecogin alfa-associated bleeding.

Therapeutic Monitoring

Routine laboratory monitoring of drotrecogin alfa is not indicated. The underlying coagulopathy of severe sepsis affects most coagulation tests, making them unsuitable for assessing the specific effects of drotrecogin alfa.

Other Anticoagulants

Many other new anticoagulant drugs are in development or clinical trials. For an overview, the interested reader is referred to an excellent article by Weitz and colleagues³⁰.

Effects of Anticoagulants on Coagulation Tests

Anticoagulant drugs are commonly used in both inpatients and outpatients. Different anticoagulants affect coagulation tests to varying degrees. One of the challenges for both clinicians and laboratories is in assessing whether abnormal coagulation test results are due to the presence of anticoagulants, either as therapeutic agents or specimen contaminants, or abnormalities in the hemostatic system. Table 10.6 summarizes the effects on coagulation tests of the anticoagulants discussed in this chapter^{16,17,29,31–33,35,36}.

Common Pitfalls

Therapeutic monitoring of anticoagulation requires laboratories to produce reliable test results and, for some assays, therapeutic ranges. Some of the more common problems include:

- The PT thromboplastin manufacturer does not provide the correct ISI for the analyzer.
- The laboratory does not correctly determine the MNPT for the INR calculation.
- The laboratory does not assess the accuracy of its INRs by comparison with prior results, other laboratories, or plasma calibrators.
- The INR is calculated incorrectly.
- The PTT therapeutic range for UFH therapy is not determined initially by the laboratory or is not verified with each reagent lot change.
- Incorrect materials are used to establish the standard curves for anti-Xa assays; for example, LMWH heparin is monitored using an anti-Xa assay set up with UFH.
- The poor interchangeability of ACT results between analyzers is not recognized.
- The influence of anticoagulants on common coagulation tests is not recognized.

Table 10.6. Effects of Anticoagulant Medications on Coagulation Tests^{a,b}

| Anticoagulant | PT ^c | PTT | TT | Fib | DD | Anti-Xa | ACT | DRVVT | VWF act | APC Res | Plsm act | PC act | PC Ag | PS act | AT act | F.II | F.IX | F.X | PLT Agg |
|---|-----------------|-----|-----|-----|----|---------|-----|-------|---------|----------------|----------|--------|-------|--------|--------|------|------|-----|---------|
| Argatroban | ↑ | ↑↑ | ↑↑ | ↓↓ | 0 | 0 | ↑↑ | ↑↑ | 0 | ? | 0 | ↑ | 0 | ↑↑ | 0 | ↓↓ | ↓↓ | ↓ | 0 |
| Bivalirudin | 0-↑ | ↑↑ | ↑↑ | ↓ | 0 | 0 | ↑↑ | ↑↑ | 0 | ? | 0 | ↑ | 0 | ↑↑ | 0 | ↓ | ↓↓ | 0 | 0 |
| Lepirudin | 0-↑ | ↑↑ | ↑↑ | ↓ | 0 | 0 | ↑↑ | ↑↑ | 0 | ? | 0 | ↑↑ | 0 | ↑↑ | 0 | 0 | ↓↓ | 0 | 0 |
| Fondaparinux | 0 | 0-↑ | ↑ | 0 | 0 | ↑↑ | ↑↑ | 0 | 0 | ? | 0 | 0 | 0 | ↑ | 0 | 0 | 0 | 0 | 0 |
| Warfarin | ↑↑ | 0-↑ | 0-↑ | 0 | 0 | 0 | 0-↑ | ↑↑ | 0 | 0 ^d | 0 | ↓ | ↓ | ↓ | 0 | ↓↓ | ↓↓ | ↓↓ | 0 |
| Heparin, unfractionated, standard dose IV or high-dose SC | 0-↑ | ↑↑ | ↑↑ | 0 | 0 | ↑↑ | ↑↑ | ↑↑ | 0 | 0 ^d | 0 | ↑ | 0 | ↑ | 0 | ↓ | ↓ | ↓ | 0 |
| Heparin, unfractionated, low-dose SC | 0 | 0-↑ | ↑ | 0 | 0 | 0-↑ | 0-↑ | 0 | 0 | ? | 0 | 0-↑ | 0 | 0-↑ | 0 | 0 | 0 | 0 | 0 |
| Heparin, unfractionated, high-dose IV | ↑ | ↑↑ | ↑↑ | 0 | 0 | ↑↑ | ↑↑ | ↑↑ | 0 | ↑↑ | 0 | ↑ | 0 | ↑ | 0 | ↓ | ↓ | ↓ | 0 |

(Continued)

Table 10.6. (Continued)

| Anticoagulant | PT ^c | PTT | TT | Fib | DD | Anti-Xa | ACT | DRVVT | VWF act | APC Res | Psm act | PC act | PC Ag | PS act | AT act | F.II | F.IX | F.X | PLT Agg |
|--|-----------------|-----|----|-----|----|---------|-----|-------|---------|---------|---------|--------|-------|--------|--------|------|------|-----|---------|
| Heparin, LWM, therapeutic | 0 | 0-↑ | ↑ | 0 | 0 | ↑↑ | 0-↑ | ↑ | 0 | ↑ | 0 | ↑ | 0 | ↑ | 0 | ↓ | ↓ | ↓ | 0 |
| Heparin, LWM, prophylactic | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| Drotrecogin alfa (activated protein C) | 0 | ↑↑ | ? | ? | ? | ? | ? | ? | ? | ? | ? | ? | ? | ? | ? | 0 | ↓↓ | 0 | ? |

^a References are noted in the text.

^b Effects are dose-dependent and reagent-dependent

^c Abbreviations and symbols:

- ACT activated clotting time, clotting
- Anti-Xa anti-Xa heparin, LMW heparin, or fondaparinux assay, chromogenic
- APC Res activated protein C resistance, clotting
- AT act antithrombin activity (functional), chromogenic
- DD D-dimer, EIA or immunoturbidimetric
- DRVVT dilute Russell viper venom time
- F.II factor II (prothrombin), clotting (PT-based)
- F.IX factor IX, clotting (PTT-based)
- F.X factor X, clotting (PT-based)

(Continued)

Table 10.6. (Continued)

| | |
|---------|---|
| Fib | fibrinogen, clotting |
| PC | act protein C activity (functional), clotting |
| PC Ag | protein C antigenic, EIA |
| Plsm | plasminogen activity (functional), clotting |
| PLT Agg | platelet aggregation |
| PT | prothrombin time, clotting |
| PTT | activated partial thromboplastin time, clotting |
| TT | thrombin time, clotting |
| VWF act | von Willebrand factor activity (functional), platelet agglutination |
| ? | unknown |
| 0 | unaffected or not significantly affected |
| ↑ | mild to moderate increase or prolongation |
| ↑↑ | moderate to marked increase or prolongation |
| ↓ | mild to moderate decrease |
| ↓↓ | moderate to marked decrease |

^d No effect if method uses dilution with factor V-deficient plasma; clotting times are prolonged if method uses undiluted specimen

Summary and Key Points

Laboratory monitoring of anticoagulation therapy is critical for the efficacious and safe use of anticoagulants. Some anticoagulants require monitoring of their specific anticoagulant effects, but many do not. All anticoagulant therapy requires some level of general monitoring for complications such as occult bleeding. This chapter has focused on specific monitoring. Some key points are:

- Warfarin therapy is widely used and monitored with the INR, a mathematical transformation of the PT involving a manufacturer-assigned ISI and laboratory-determined MNPT or involving a calibration procedure in the laboratory. The purpose of the INR is to transform PT values into the PT ratio that would have been obtained had the specimen been tested by a reference method. Use of the INR has facilitated the evolution of global therapeutic ranges, but the INR schema does not, as yet, assure full comparability of results between laboratories.
- UFH therapy is widely used. A major limitation is a highly variable dose-response between patients due largely to non-specific binding of large heparin molecules to plasma proteins, endothelial cells, monocytes, and platelets. The PTT is the principal test used for therapeutic monitoring, but it also exhibits a highly variable response to heparin. It is incumbent on each laboratory to establish therapeutic range limits and to verify or re-establish the limits whenever a reagent lot change is made or a new reagent is implemented. The PTT is not useful for monitoring LMWH or high-dose UFH therapy.
- The ACT is used for monitoring high-dose UFH therapy, generally for cardiac catheterization, cardiopulmonary bypass, or related procedures. The ACT has a proven clinical record, but suffers from poor precision and poor comparability between analyzers. Therapeutic ranges are generally determined from published clinical protocols.
- LMWH and fondaparinux exert their anticoagulant effects through the activation of AT, as does UFH, but with far more highly predictable dose-responses. Routine monitoring is not required except in special circumstances including body weight extremes, young age, renal dysfunction, and prolonged therapy.
- The anti-Xa assay is increasingly used for monitoring UFH, LMWH, and fondaparinux therapy. A specific standard curve must be established for each type of anticoagulant. PTT therapeutic ranges are defined by anti-Xa therapeutic range limits.

- Direct thrombin inhibitors, including argatroban, lepirudin, and bivalirudin, selectively inhibit thrombin and do not require AT activity. Argatroban and lepirudin require therapeutic monitoring with the PTT with limits defined in terms of PTT ratios. Laboratories do not have the means of assessing whether these ratios are suitable for their PTT methods.

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11

Coagulation Testing and Transfusion Medicine

Robert C. Blaylock*
and Christopher M. Lehman

Coagulation Testing and Transfusion Medicine

Transfusion Services are presented with many challenging coagulation problems. Most cases gain attention when out of the ordinary requests for blood products are received. The Transfusion Medicine Physician is often brought into a clinical scenario that has already become an emergency and quick response is imperative¹⁻⁸. Decisions often have to be made empirically, or with the help of a very limited coagulation test menu. This chapter will focus on maximizing information from a short coagulation test menu, reviewing the tools available to correct coagulopathies, and applying these tests to emergency situations.

Essential Coagulation Assays

The basic coagulation assays available in most hospitals include a platelet count, prothrombin time (PT), partial thromboplastin time (PTT), fibrinogen, mixing study, D-Dimer, and heparin assay. These tests have already been discussed in detail, so what follows is a brief discussion on some misconceptions about certain assays and helpful hints regarding result interpretation in an emergency. Ultimately, the coagulation assays used to direct transfusion therapy are not as important as selecting, and implementing an appropriate algorithm to guide clinician practice⁹.

*Associate Professor of Pathology, University of Utah Health Sciences Center, Salt Lake City, UT, Medical Director, Blood Bank, The University of Utah School of Medicine, Salt Lake City, UT, and Director of University Hospital Clinical Laboratories, ARUP Laboratories, Salt Lake City, UT

Prothrombin Time

The PT is used to monitor the extrinsic and common soluble coagulation pathways. It is also used to monitor warfarin therapy via the international normalized ratio (INR) – a value calculated employing the PT, the mean normal PT and the international sensitivity index (see chapter 10, Monitoring of Anticoagulant Therapy). The INR was developed to standardize PT results between laboratories using reagents with differing sensitivities to the anticoagulant effects of warfarin. The INR can be used to guide transfusion of fresh frozen plasma in bleeding patients receiving warfarin therapy. The INR has not been evaluated as a tool for guiding transfusion in other clinical situations (e.g., massive transfusion with multiple coagulation factor deficiencies).

A common misconception among clinicians is that heparin cannot cause a prolongation of the PT. Most PT reagents now contain substances (e.g. heparinase, polybrene) that will neutralize up to 1.0–2.0 U/mL of heparin in a patient sample resulting in a PT result unaffected by heparin. However, greater plasma heparin concentrations (e.g. heparin overdose, or status-post hemodialysis or cardiopulmonary bypass) will cause a prolongation of the PT (as well as the PTT) resulting in confusion as to the cause of clinical bleeding. An example will be presented later.

Partial Thromboplastin Time

The PTT is used to measure the intrinsic and common coagulation pathways. It is commonly used to monitor heparin therapy. The PTT is not as sensitive to warfarin therapy as the PT, but overdoses of warfarin can cause a prolongation of the PTT since the intrinsic or common pathways include factors II, IX, and X.

Fibrinogen Concentration

Fibrinogen levels are extremely valuable for diagnostic purposes and in determining therapy. However, many different methodologies for measuring fibrinogen levels are available and all have their pros and cons. Clinical laboratories generally require a rapid turn-around-time and rely on functional (rather than antigenic) fibrinogen assays. The Clauss assay, which derives the fibrinogen concentration from the clotting time of diluted plasma after addition of excess thrombin, is the most commonly used method. However, fibrinogen levels can also

be calculated from the total change in optical signal observed during a PT reaction. Antigenic assays are mainly used in a reference laboratory setting. The type of assay used in a hospital is not as important as understanding how the assay performs in certain clinical situations. For example, understanding how different heparin concentrations affect fibrinogen results is critical.

Fibrinogen levels of 80–100 mg/dL are required to achieve hemostasis in a bleeding patient. Routine fibrinogen determinations in emergency situations may seem unnecessary because the PT and PTT might be expected to become prolonged with significant fibrinogen deficiency. Unfortunately, this is not necessarily true. An example of a fibrinogen deficient response curve for PT and PTT is represented in Table 11.1. The curve is created by maintaining normal levels of all clotting factors except fibrinogen. Fibrinogen levels are then varied and the PT and PTT are measured. This curve demonstrates that fibrinogen levels can drop to approximately 58 mg/dL, while the PT and PTT remain in the normal range. It is not until fibrinogen levels drop to approximately 29 mg/dL that the PT prolongs to 1.6 seconds above the upper limit of normal, but the PTT is still at the upper limit of normal. A clinical condition that results in destruction of fibrinogen may not be

Table 11.1. Effect of Decreasing Fibrinogen Concentration on the PT and PTT

| Fibrinogen Deficient Curve | | | |
|-----------------------------------|-----------------------------|--|---|
| % Activity | Fibrinogen mg/dL | PT (s) (normal 10.7–15.0 seconds) | PTT (s) (normal 25–40 seconds) |
| 100 | 288 | 11 | 28 |
| 80 | 230.4 | 11.4 | 29.8 |
| 70 | 201.6 | 11.6 | 29.8 |
| 40 | 115.2 | 12.6 | 31.9 |
| 33 | 95.04 | 13.8 | 34.5 |
| 30 | 86.4 | 13.1 | 32.6 |
| 20 | 57.6 | 14.2 | 34.3 |
| 10 | 28.8 | 16.6 | 40 |
| 5 | 14.4 | 23.4 | 44.5 |
| 0 | 0 | 54.1 | 150 |

Abbreviations: PT = prothrombin time; PTT = partial thromboplastin time.

detected if the PT and PTT are the only assays relied on to detect a low fibrinogen.

Prothrombin Time and Partial Thromboplastin Time Mixing Studies

Mixing studies are used to determine if a prolongation in a PT or PTT is due to a deficiency of clotting factors, heparin or inhibitors. The test is performed by mixing patient plasma, with pooled normal plasma in a 1:1 ratio. After mixing, the PT and PTT are repeated and the times should shorten close to, or within the normal range, if the prolonged results are due to a deficiency of clotting factors. In emergency situations, mixing studies can be useful in identifying the presence of heparin as a cause of unexpected bleeding, since a lack of correction suggests the presence of heparin or an inhibitor. However, correction of the PTT does not rule out the presence of heparin in the sample (Figure 11.1). Therefore, heparin assays should replace mixing studies for this purpose.

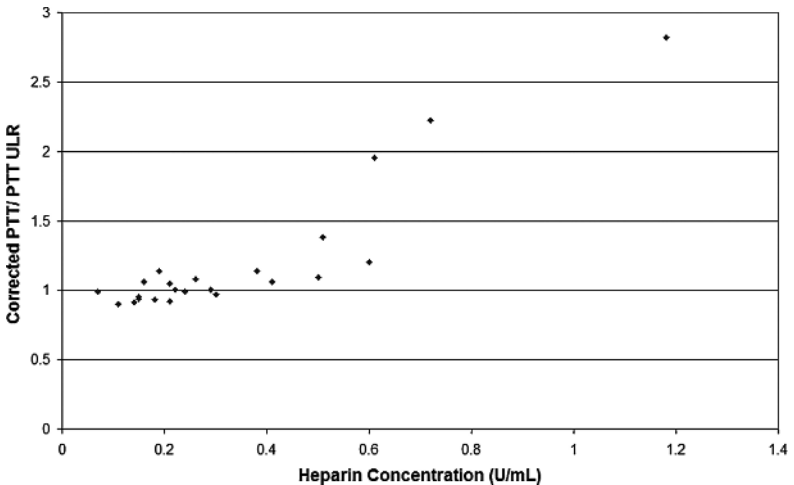


Figure 11.1. Plot of the ratio of the corrected PTT (1:1 dilution of patient sample with pooled normal plasma) to the uncorrected PTT versus the concentration of heparin in specimens collected from patients receiving unfractionated heparin therapy. Results for corrected samples were not consistently above the upper limit of the reference interval until the heparin concentration reached 0.4 U/mL. Abbreviations: ULR = upper limit of reference interval; U = units.

D-dimer Assay

D-dimer assays are very specific for the diagnosis of disseminated intravascular coagulation (DIC)¹⁰. The diagnosis of severe, acute DIC must be made quickly to find, and treat, the underlying cause. Early identification of DIC will also direct appropriate transfusion therapy as will be discussed later. Minor elevations in D-dimer are seen following routine surgical procedures, and in trauma patients, and should not be considered diagnostic for DIC^{10–12}.

Platelet Counts

Platelet counts are essential for identifying bleeding that may be secondary to thrombocytopenia and for directing transfusion therapy in a bleeding patient. This is an extremely valuable, but underutilized assay.

Thromboelastography

The search for the “Holy Grail” of coagulation assays has been ongoing for decades. The perfect assay would assess platelet number, platelet function and the soluble coagulation system with a single, rapid, inexpensive test. The thromboelastograph (TEG[®]) is the prototype example of an attempt at such an assay. TEG[®] has been discussed previously in this book, but in short, is a whole blood clotting assay that gives a real time graph of clot formation and stability. Standard TEG[®] analysis was expected to be the ultimate coagulation assay because of the ability to assess all components necessary for clot formation. Subsequent studies demonstrated that the standard TEG[®] assay was not able to differentiate between the effects of viable platelets versus cryodisrupted platelets that were added to the test system. Therefore, standard TEG[®] analysis appears to be sensitive to the amount of phospholipid in the test system, but unable to differentiate between functional and non-functional platelets¹³. New TEG[®] assays have been developed that use platelet agonists as initiators of clot formation in addition to the original phospholipid reagent¹⁴. This modification was made to better assess platelet function. These new assays have significantly increased the cost of TEG[®] without convincing evidence of clinical benefit (see Chapter 7, Testing for Acquired Platelet Disorders).

Transfusion Medicine Tools: Blood Components

Platelets

Platelet concentrates should be transfused in a systematic fashion. This requires consideration of three questions:

- 1) Is thrombocytopenia causing life-threatening hemorrhaging?
- 2) What is the cause of the thrombocytopenia?
- 3) Did the patient's platelet count increase after platelet transfusion?

The primary sites of life-threatening hemorrhage due to thrombocytopenia include the brain and the lungs. If thrombocytopenia is the suspected cause of life-threatening hemorrhage, platelets must be transfused immediately.

Bleeding patients who are not in jeopardy of death or significant morbidity should not be transfused until the cause of the low platelet count is evaluated. Understanding the etiology of the thrombocytopenia is critical because transfusion of platelets in certain disease states can actually cause morbidity and even death. Two clinical conditions where platelet transfusions should be avoided are thrombotic thrombocytopenic purpura (TTP) and heparin-induced thrombocytopenia. Platelet transfusions administered to patients with these conditions can cause an increase in platelet thrombi in tissues resulting in end organ damage.

After platelets are transfused, it is important to check a platelet count to ascertain the success of the transfusion. Many clinical conditions can lead to the immediate destruction of platelets including autoimmune thrombocytopenic purpura, sepsis, fever or splenomegaly. The administration of medications such as amphotericin or anti-thymocyte globulin can also result in platelet destruction. Patients with a prior history of pregnancy and/or transfusion are at risk for developing HLA antibodies and may need HLA matched platelets to have an adequate response to platelet transfusion.

The best way to monitor platelet transfusion success is by determining a pre-transfusion platelet count and checking the count again one hour after completing the platelet transfusion. If time does not permit waiting for an hour (e.g. the need for an invasive procedure), a post-transfusion platelet count can be checked after 10 minutes. The transfusion of an adult dose of platelets (one apheresis platelet unit or 6 pooled, random donor platelet units derived from whole blood) should

increase the platelet count by 30,000 to 60,000/ μL . Factors determining the magnitude of the response include the number of platelets in the product, the size of the patient, and whether the patient has any clinical conditions leading to platelet refractoriness. Calculations such as the corrected count increment or recovery rate can best quantify the success of a platelet transfusion; however, these tools are usually not necessary in routine practice.

The goal of platelet transfusion is to provide adequate circulating platelet numbers to prevent or minimize bleeding. Patients with platelet counts below 5,000/ μL are generally considered to be at risk for life-threatening, spontaneous hemorrhage in the central nervous system. Prophylactic platelet transfusions for patients undergoing chemotherapy are usually given when counts are between 10,000 and 20,000/ μL at the discretion of the clinician. Some clinician's prefer to limit patient exposure to blood products and will tolerate minor mucosal bleeding and petechiae to avoid the risks of transfusion. Others transfuse platelets at counts closer to 20,000/ μL to limit bleeding as much as possible.

Most patients are not at risk for severe bleeding when platelet counts are above 20,000 per μL , unless they are scheduled for an invasive procedure or have received anti-platelet medications. Most invasive procedures, or surgery, can be safely performed when counts are above 50–75,000 per/ μL , but many surgeons aren't comfortable until counts are above 100,000/ μL . Striving for platelet counts significantly above 100,000/ μL is neither a prudent, nor necessary use of a precious resource. Additional platelet transfusions during an invasive procedure may be necessary, and should be guided by platelet counts obtained during surgery.

The following case history demonstrates why a systematic approach to platelet transfusion is essential to optimize patient outcome. A sixty year old male was scheduled for a total hip replacement. He had a long history of aplastic anemia that required red cell transfusions. The patient's platelet count usually ran around 30,000 per μL . He was transfused two units of apheresis platelets and taken to surgery without checking a post-transfusion platelet count. The patient required more than 70 blood product units during the procedure and almost exsanguinated. Later investigation revealed the patient did not respond to the pre-operative platelet transfusions because of preformed HLA antibodies that immediately destroyed the transfused platelets.

A systematic approach to platelet transfusion could have prevented this near disaster. The patient did not have life-threatening bleeding prior to surgery, but performing a total hip replacement with a platelet count of 30,000/ μL put him at risk. Prior to platelet transfusion, the cause of the patient's thrombocytopenia should have been investigated to determine if there was a contraindication to platelet transfusion. In this case the cause was aplastic anemia and platelet transfusion was indicated. Finally, a post-transfusion platelet count should have been checked to see if surgery could have been performed safely. Unfortunately, this was not done and the surgeon was not aware that the platelet transfusions had not increased the pre-operative platelet count. If the post-transfusion count had been performed, surgery would have been canceled and re-scheduled for a day when HLA matched platelets would have been available to raise the patient's platelet count to a level adequate for a safe procedure.

Fresh Frozen Plasma (FFP)

Plasma is frozen soon after collection to preserve levels of the labile clotting factors V and VIII. FFP not only contains all factors in the soluble coagulation system, but also natural inhibitors of the soluble coagulation system (e.g. anti-thrombin), and proteins of the fibrinolytic system (e.g. plasminogen). FFP provides all factors necessary for the delicate balance between hemostasis and thrombosis. However, FFP is used mainly in hemorrhaging patients to replace deficiencies of multiple coagulation factors.

Hospitals are required to have laboratory guidelines for the use of FFP. The most commonly used cutoff to indicate the need for FFP transfusion is a PT and/or PTT that exceeds 1.5 times the mid-range of normal³. The problem with this universally applied guideline is that it does not result in a standard that can be compared between different hospitals and coagulation assay methodologies. Note that an INR of 1.5 is not necessarily equivalent to a PT of 1.5 times the mid-range of normal.

An alternative to the above calculation involves using a dilution curve to provide a guideline for FFP use (Table 11.2). A dilution curve is performed by diluting pooled, normal plasma with a known amount of crystalloid solution (e.g. normal saline) and performing PT and PTT assays on the dilutions. For example, the 50% PT and PTT times were run on a sample of 50% normal saline and 50% normal plasma. The dilution curve is an in-vitro simulation of what happens

Table 11.2. Sensitivity of the PT and PTT to Dilution of Pooled Normal Plasma with Normal Saline

| Dilution Sensitivity (upper limit of normal in seconds: PT = 15.5 s, PTT = 37 s) Prothrombin Activity Curve | | |
|--|--------|---------|
| % Activity | PT (s) | PTT (s) |
| 100 | 13.6 | 29 |
| 90 | 14.5 | 30 |
| 80 | 15.2 | 31 |
| 70 | 15.9 | 33 |
| 60 | 17.2 | 36 |
| 50 | 19.2 | 40 |
| 40 | 22.4 | 48 |
| 30 | 27.7 | 64 |
| 20 | 40.3 | 99 |
| 15 | 52.8 | >150 |
| 10 | 102.8 | >150 |
| 5 | >150 | >150 |
| 1 | >150 | >150 |

Abbreviations: PT = prothrombin time; PTT = partial thromboplastin time.

when a bleeding patient is resuscitated with normal saline prior to FFP availability. The PT and PTT at the 50 to 40% level are used as the guideline for FFP transfusion in the case of dilutional coagulopathy. The advantage to this approach is that it assures that the PT and PTT that are used as a guideline in a given institution fall at the same place on a dilution curve. This is not always the case when using the 1.5 times midrange of normal guideline, where the calculation may lead to a 10–20% difference between where the PT and PTT fall on the dilution curve. While the dilution curve gives more consistency between the PT and PTT guideline values, it does not solve the problem of standardization between hospitals. For example, the 50% cutoff will not result in the same number of seconds for PT and PTT assays in different hospitals.

Many clinicians attempt to use the INR as a guideline for FFP use. While it is true the INR will increase with a worsening coagulopathy, it has not been thoroughly evaluated as a method to direct FFP therapy. The exception would be in the setting of rapid warfarin reversal.

Cryoprecipitate

Historically, cryoprecipitate has been used to treat hemophilia A and von Willebrand disease, but now is used almost exclusively as a source of fibrinogen. The content, and indications for use of cryoprecipitate are not understood by many clinicians. Cryoprecipitate is produced from FFP by allowing FFP to thaw in a 1–6 degree Celsius environment. When kept at this temperature, the FFP melts, but 10–20 mL of precipitate will form. Refrigerated centrifugation separates the precipitate from the liquid. The precipitate can then be isolated, and when warmed, will go into solution. The result is a liquid, which contains fibrinogen, factor VIII, von Willebrand factor and factor XIII. Myths surrounding the use of cryoprecipitate include that it is a concentrated form of FFP and that it contains more fibrinogen than FFP. In fact, cryoprecipitate contains only the factors listed above, and only about 50% of the fibrinogen contained in the original unit of FFP. Cryoprecipitate does not contain as much fibrinogen as FFP, but there is more fibrinogen per unit volume compared with FFP (i.e., it is a concentrated source of fibrinogen). Each bag of cryoprecipitate contains, on average, about 200 mg of fibrinogen. As noted earlier in this chapter, fibrinogen levels of 80–100 mg/dL are required to achieve hemostasis in a bleeding patient. To raise the fibrinogen level in an adult by 100 mg/dL requires between 10–30 bags of cryoprecipitate depending on the size of the patient (about 2 bags per 10 kilograms of body weight). The most common error in cryoprecipitate transfusion is delivery of a dose insufficient to significantly increase fibrinogen levels in an adult (e.g. 4 bags). Patient fibrinogen levels should always be re-evaluated after cryoprecipitate infusion.

Clinical Scenarios and Case Studies

Dilutional Coagulopathy

Massive transfusion is defined as replacement of a patient's blood volume in a 24-hour period¹. Some prefer the definition of transfusion of 10 units of packed red blood cells in an adult², or even four packed red blood cells in a one-hour period¹. Most massively transfused patients are initially resuscitated with crystalloids followed by packed red blood cells. Red cells contain negligible amounts of the soluble coagulation system and no viable platelets, while crystalloids contain nothing involved in hemostasis. The result is dilution of the soluble

coagulation system and platelets. Each will be discussed separately, but must be evaluated together in a bleeding patient.

Dilution of the Soluble Coagulation System

Therapeutic plasma exchange (TPE) represents the best example of pure dilution of the soluble coagulation system. In TPE, whole blood is removed from the patient and directed into an automated cell separator. The various components of the blood are separated by differential centrifugation and the desired blood fraction (e.g., plasma) is removed. TPE is used in many clinical conditions where a pathologic substance in plasma (often an auto-antibody) produces a disease state. The rate of removal of the pathologic substance is dependent upon the number of plasma volumes exchanged¹⁵ (Table 11.3).

Unfortunately, TPE does not exclusively remove the pathologic substance, and beneficial contents of plasma, such as clotting factors, are also removed. FFP should be the ideal replacement fluid because it contains clotting factors and immunoglobulins, but it carries risks of transfusion transmitted disease and adverse reactions, such as transfusion related acute lung injury. Therefore, FFP is only used as a replacement fluid for patients at risk of bleeding, or who need a specific factor contained in FFP such as the metalloprotease required for patients with TTP.

Colloids, or a combination of colloid and crystalloids, are the usual replacement fluid selected for TPE. These solutions contain no clotting factors, and patients must be monitored closely to avoid spontaneous hemorrhage and to ensure the safety of invasive procedures if they become necessary in the course of patient management. Routine laboratory testing prior to TPE should include a platelet count, PT, PTT and fibrinogen. These assays should be in the normal range before a

Table 11.3. Effect of the Number of Plasma Volumes Exchanged on the Removal of Pathologic Substances in Plasma

| Plasma volumes exchanged | Fraction Removed |
|---------------------------------|-------------------------|
| 0.5 | .39 |
| 1.0 | .43 |
| 1.5 | .78 |
| 2.0 | .86 |

TPE procedure is initiated. Clinicians rarely appreciate the severity of the transient coagulopathy produced by a TPE procedure. They must be reminded that all clotting factors are depleted by the procedure, and this can result in a coagulopathy more severe than that seen in hemophilia patients lacking a single clotting factor. TPE with colloid replacement is usually performed on an every other day basis to allow for recovery of clotting factors through endogenous production prior to the next procedure. However, recovery rates vary between patients, and coagulation testing prior to each procedure is prudent.

The following is a poignant case illustrating the severity of a pure, dilutional coagulopathy caused by TPE. A 77 year-old male was admitted for TPE to treat myasthenia gravis. The platelet count, PT, PTT and fibrinogen were all within the normal range prior to the procedure. The patient underwent a 1.3 plasma volume exchange utilizing a left femoral vein, dual-lumen catheter, and using a 5% albumin solution as the replacement fluid. The procedure would have predicted approximately 70% removal of the original clotting factors, but should not have resulted in spontaneous hemorrhage.

Three to four hours after completion of the procedure, the apheresis team was called by an intensive care doctor wondering why the patient was found hypotensive with a 4–5 unit hematoma in the right groin. A hematoma in the right groin was perplexing since the catheter used for the procedure was in the left groin, and venous lines rarely bleed after TPE. The following coagulation assay results were obtained:

- | | | |
|-----------------------|----------|------------|
| • PT (11–15.5 sec) | 29 sec | Mix 16 sec |
| • PTT (25–37 sec) | 72 sec | Mix 34 sec |
| • FIB (150–350 mg/dL) | 70 mg/dL | |
| • D-dimer | normal | |

(Where FIB equals fibrinogen).

The PT, PTT and fibrinogen were about what would be expected following a TPE procedure. The PT and PTT were prolonged but the mixing study corrected almost to normal, and the platelet count was above 100,000/ μ L. The fibrinogen was decreased as expected, and a normal D-dimer ruled-out DIC. Comparing the PT and PTT after the procedure with the dilution curve (Table 11.2) shows the results landing on the curve between the 30% and 20% level. This means that 70% to 80% of the original clotting factors were removed. The predicted removal rate was around 70%, but further investigation showed that

the patient had received some additional crystalloid fluid when he was found hypotensive, explaining the slightly more severe coagulopathy than predicted. The patient was given 8 units of FFP. The coagulopathy was corrected and the bleeding stopped.

The mystery surrounding this case involved why bleeding occurred in the right groin when the line was placed in the left femoral vein. Further investigation revealed that the physician inserting the line first attempted to place it in the right femoral vein, but aborted the attempt when he cannulated the femoral artery by mistake. He had pulled the line out, and held pressure until the bleeding had stopped, but failed to inform anyone of the mishap. A platelet plug had stopped the initial bleeding in the artery, but once the TPE was initiated, clotting factor levels were dropped to levels that would not allow adequate fibrin formation to cement the platelet plug in place.

All cases of massive transfusion result in a similar coagulopathy when crystalloid resuscitation is used prior to FFP availability. Blood banks must have protocols in place that allow for thawing of FFP early in the course of massive transfusion. Often, FFP must be given empirically, since time may be insufficient for the laboratory to turn around basic coagulation results. When possible, thawed FFP should be available before the patient receives a complete blood volume of red cells and crystalloid.

Dilutional Thrombocytopenia

Dilutional thrombocytopenia occurs later in the course of transfusion of resuscitated patients than what would be predicted by “wash-out” mathematical calculations¹. This probably reflects mobilization of endogenous platelet reserves including marginated platelets, platelets contained in the spleen and lungs, and platelets released early from the bone marrow. Platelet transfusions are usually given to massively transfused patients after initial transfusion of red cells and plasma, and are usually effective in maintaining platelet counts sufficient for primary hemostasis. As already noted, the effectiveness of platelet transfusions should be monitored with frequent platelet counts, a fast and inexpensive assay.

Inadvertent Heparinization

Heparin can be very problematic for the Transfusion Service, particularly when it is present in a sample due to contamination from a

line draw, or, more importantly, due to unintentional heparinization of the patient resulting in unexpected bleeding. However, the presence of heparin in a sample must be suspected for the heparin assay to be ordered. Routine coagulation assay patterns can help identify samples containing heparin. The first clue to the presence of small concentrations of heparin will be a greatly prolonged PTT with a PT that is normal or close to normal. The discrepancy between the two assays is due to the heparinase or heparin-binding agent contained in the PT reagent that neutralizes the effects of the heparin. If heparin concentrations are very high, both the PT and PTT will be prolonged, usually greater than the time where laboratories stop reporting results (e.g. > 150 seconds). Mixing studies were historically used to provide evidence for the presence of heparin, but now, once heparin is suspected, a quantitative heparin assay should be performed.

Once the presence of heparin is confirmed, the investigation must continue to determine if it is sample contamination, or if the patient had been given heparin. This may be as easy as asking the phlebotomist if the sample was obtained with a peripheral stick, or was collected from a line. The next step should be to contact the nurse to see if the patient has been placed on heparin and if the patient shows any signs of bleeding. The investigation must continue even if the floor denies overt heparin administration because of ubiquitous heparin availability in hospitals. Patients may receive heparin for dialysis, invasive diagnostic procedures or may be status post-cardiopulmonary bypass. Patients may have large concentrations of heparin in central lines, which can be inadvertently flushed into the patient. Heparin comes in many different concentrations that can lead to an inadvertent overdose.

The following case illustrates the difficulties heparin can cause, both in diagnosis and in delaying treatment. A two-day old full term male was in the well-baby nursery of a small community hospital when the nurse noticed an increased respiratory rate. The pediatrician was concerned that the baby was developing pneumonia and gave the child a bolus of antibiotics while arranging transportation to a children's hospital. Upon arrival at the children's hospital the child had confluent ecchymoses and signs consistent with meningitis. A lumbar puncture was performed which resulted in severe bleeding, spinal cord compression and lower extremity paralysis. An ultrasound demonstrated intra-ventricular hemorrhage as well. Fearing DIC, clinicians ordered FFP, cryoprecipitate and platelets, which led to an investigation by

the blood bank physician. Laboratory results on the baby showed the following:

- PT > 150 seconds, mixing study > 150 seconds
- PTT > 150 seconds, mixing study > 150 seconds
- Fibrinogen = 0 milligrams/dL
- D-dimer = normal
- *Plateletcount* = 170,000/ μ L

The blood bank physician covering the pediatric hospital was perplexed by the results. The PT, PTT and fibrinogen were consistent with DIC, but the platelet count, D-dimer and mixing studies were more consistent with heparinization. The physician had the assays repeated at an adjoining adult hospital where he was more familiar with the coagulation assay characteristics and the following results were obtained:

- PT > 150 seconds, mixing study > 150 seconds
- PTT > 150 seconds, mixing study > 150 seconds
- Fibrinogen = 142 mg/dL
- D-dimer = normal
- Platelet count = 165,000/ μ L

These results were consistent with systemic heparinization of the baby given the severe bleeding history. The nurse was contacted with an explanation of the laboratory results and to recommend protamine administration. The nurse did not believe the interpretation of the results, so a heparin assay was performed which was 5.2 U/mL (recommended heparin therapeutic range equals 0.3–0.7 U/mL). The attending physician was convinced of the heparin overdose and protamine was given to the child correcting the coagulopathy.

This case makes some salient points. The first is the importance of understanding the strengths and weaknesses of coagulation assays in the hospital where you practice. The assays obtained at the children's hospital were confusing because of the fibrinogen level of zero. The blood bank physician later discovered that fibrinogen levels at the children's hospital were calculated from the rate of clot formation on the PT assay. Heparin had prevented the PT from clotting; therefore a zero fibrinogen level was calculated. The fibrinogen assay at the adult hospital was resistant to the effects of heparin. The physician knew heparin would falsely reduce the true fibrinogen level by 10–20%, but that fibrinogen would be measured. The second point is that a

heparin assay result is often required to convince clinicians that the bleeding problem is related to heparin. The final question to answer was where did the heparin come from? In this case, as in most mysterious cases of inadvertent heparin administration, the transfusion medicine physician must perform the investigation. This investigation revealed that the community hospital had decided to heparinize the patient's I.V. after an antibiotic bolus was given to the child. Unfortunately, instead of 10 U/mL of heparin, the nurse was handed a concentration of 10,000 U/mL, and the child was inadvertently, systemically anticoagulated.

Heparin can complicate patient care at anytime. Know how coagulation assays in your hospital respond to heparin, and have a low threshold for performing heparin assays when unexpected bleeding occurs.

Disseminated Intravascular Coagulation

DIC is caused by a stimulus that leads to abnormal and constant thrombin generation¹⁶. Many etiologies for DIC exist, but the most severe, acute cases of DIC are obstetrically related. Causes of DIC in pregnant women include amniotic fluid embolus, infection and retained products of conception. While clinicians search for the underlying cause of the DIC, the blood bank must provide product support before and after the underlying cause is treated.

In addition to thrombin generation, DIC also results in activation of the fibrinolytic system and abnormal amounts of plasmin are generated¹⁶. Thrombin converts fibrinogen to fibrin, and plasmin not only digests cross-linked fibrin clots, but also destroys fibrinogen before it is converted to fibrin. Therefore, DIC results in a rapid, and disproportionate reduction in fibrinogen levels relative to other clotting factors. Treatment must include rapid replacement of fibrinogen, and cryoprecipitate is the ideal blood product for this purpose, as demonstrated below.

A 27 year-old female, with a pregnancy estimated at 29 weeks gestation, was admitted with premature rupture of membranes. Fetal heart tracings deteriorated and the patient was taken to the operating room for a successful cesarean delivery without immediate complications. A post-operative check on the patient was performed 1.5 hours after delivery. She was pale, tachycardic, hypotensive, and oozing from the incision site with approximately one liter of blood found in

Table 11.4. Selected Laboratory Values from a Patient Diagnosed with DIC

| Time ^a (hours) | HCT 39–47% | PLT | | | Fibrinogen | D-dimer <.5 mg/dL |
|------------------------------|---------------|-------------------|---------------|----------------|------------------|----------------------|
| | | 140–440,000 μL | PT 13–16 s | PTT 25–36 s | 150–350 mg/dL | |
| Admit | 39 | 183 | | | | |
| 1.5 | 17 | 53 | >150 | >150 | <7 | >256 |
| 4.5 | 18 | 19 | 21 | 48 | 152 | 64 |
| 6 | 29 | 134 | 17 | 31 | | |

^a Time in hours after admission to the hospital

Abbreviations: HCT = hematocrit; PLT = platelet count; PT = prothrombin time; PTT = partial thromboplastin time; s = seconds; DIC = disseminated intravascular coagulation.

the bed. She was taken emergently back to the operating room for a hysterectomy. Laboratory results at admission and hours after the original cesarean section are shown in Table 11.4.

The laboratory findings 1.5 hours after delivery, when the patient was found in shock, show the hallmarks of DIC. Most notably, no detectable fibrinogen and extremely high D-dimer levels. Thirty units of cryoprecipitate were given to rapidly correct the fibrinogen level. As previously noted, cryoprecipitate provides the most concentrated form of fibrinogen and can be rapidly thawed, pooled and infused into the patient. In addition to cryoprecipitate, the patient received nine units of packed red blood cells, two apheresis packs of platelets and seven units of FFP. The laboratory results drawn 4.5-hours after the hysterectomy show dramatic improvement. By the six-hour mark, the PT and PTT had returned to normal and the patient was stable and survived without permanent sequelae.

DIC is one of the most feared clinical conditions facing Transfusion Medicine Services. Severe, acute DIC is most commonly related to obstetrical cases and has a high mortality rate. Successful treatment must be directed at the underlying cause and the blood bank must focus on rapid replacement of fibrinogen. This is best accomplished with the use of adequate amounts of cryoprecipitate. Plasma and platelets will also be required to provide the factors not contained in cryoprecipitate.

Summary and Key Points

Transfusion services commonly encounter cases of catastrophic bleeding. The key to successful management of the cases is rapid

diagnosis of the cause, using routine coagulation assays. Once the coagulopathy is understood, the appropriate component therapy can be provided. Many algorithms directing blood product transfusion exist. The key to success is thorough understanding of the strengths and weaknesses of the assays used to support the algorithm.

- Have an early warning system in place for catastrophic bleeding and treat empirically with blood products if necessary
- Develop a limited coagulation test menu, and algorithm, to guide blood product administration during massive transfusion
- Understand the strengths and weaknesses of the coagulation assays in your hospital
- Understand how heparin affects coagulations assays, and have a low threshold for ordering a heparin assay
- Use a systematic approach for platelet transfusions
- Treat severe, acute DIC aggressively with blood products. Early focus should be on improving low fibrinogen levels using cryoprecipitate

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