Identification and Basic Management of Bleeding Disorders in Adults

Rebecca Kruse-Jarres, MD, Tammuella C. Singleton, MD, and Cindy A. Leissinger, MD

Adults with bleeding disorders may present to their family physician with minor bleeding symptoms or hematologic laboratory abnormalities discovered during evaluation for surgery or another purpose. Identifying the small proportion of adults who have an underlying bleeding disorder as the cause for such signs or symptoms may be challenging. In cases of asymptomatic hematologic laboratory abnormalities, the particular abnormality should narrow down the potentially affected hemostatic component(s), ideally streamlining subsequent investigation. In patients presenting with bleeding symptoms, a thorough history and physical examination are critical for first identifying bleeding as pathologic, then performing the appropriate diagnostic evaluation after excluding identifiable causes. Knowledge of the pathophysiologic processes contributing to impaired hemostasis in any given bleeding disorder ensures proper treatment and avoids therapies that are unnecessary or even contraindicated. Management is further determined by bleeding phenotype and, for invasive procedures, the anticipated risk for bleeding. Consultation with a hematologist may facilitate proper evaluation and treatment, particularly in adults with rare bleeding disorders or no identifiable cause for bleeding. This article reviews the diagnostic approach to hematologic laboratory abnormalities and abnormal bleeding in adults, as well as basic preventive care and hemostatic management of adults with bleeding disorders. (J Am Board Fam Med 2014;27:549–564.)

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From the Department of Medicine (RKJ, CAL), and the Department of Pediatrics (TCS), Tulane University School of Medicine, New Orleans, Louisiana.
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Corresponding author: Rebecca Kruse-Jarres, MD, Section of Hematology and Medical Oncology, Tulane University School of Medicine, 1430 Tulane Ave., Box SL-78, New Orleans, LA 70112-2699 (E-mail: rkruseja@tulane.edu).

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preventive care and hemostatic management of those ultimately diagnosed with bleeding disorders also is discussed.

Diagnostic Considerations in an Adult With a Potential Bleeding Disorder

Abnormal Bleeding Symptoms

Bleeding is a common symptom and does not always indicate an underlying bleeding disorder. Symptoms such as gum bleeding, epistaxis, menorrhagia, petechiae, and bruising are especially common; in one study they were reported by anywhere from 22% to 85% of men and women without bleeding disorders. Identification of pathologic bleeding may, therefore, prove challenging. Clinically significant mucocutaneous bleeding is defined as any of the following: spontaneous or provoked bleeding from 2 or more distinct mucocutaneous sites; bleeding from a single site warranting blood transfusions; or bleeding from a single site on 3 or more separate occasions. Bleeding scoring systems have shown promise in retrospectively predicting bleeding phenotype in type 1 VWD and prospectively excluding mild bleeding disorders in patients presenting with bleeding symptoms or abnormal coagulation study results, but they require further investigation and validation for broader clinical use.

A thorough history and physical examination often provides clues as to whether bleeding is pathologic and may even point to potential underlying causes. Historical factors to explore are outlined in Table 1. During physical examination, the skin and mucous membranes should be inspected for stigmata of bleeding (eg, bruising, petechiae) and other findings suggestive of potential underlying causes of bleeding (eg, jaundice, telangiectasia). The presence of hepatomegaly, splenomegaly, or joint hypermobility may suggest potential diagnoses associated with bleeding. Skin or conjunctival pallor, tachycardia, or a cardiac flow murmur may indicate associated anemia. Historical and physical findings may suggest an abnormality of either primary hemostasis, which culminates in the formation of a platelet plug; secondary hemostasis, in which fibrin is formed via the coagulation “cascade”; or fibrinolysis, the normal breakdown of clots. A simplified schematic of the coagulation cascade and the corresponding laboratory assays for each pathway are provided in Figure 1. A more thorough review of the complex process of hemostasis and its various components is beyond the scope of this article but can be found elsewhere.

Excessive bruising, epistaxis, bleeding after dental extraction, and menorrhagia are symptoms suggestive of quantitative or qualitative platelet disorders. Patients with platelet abnormalities may also experience excessive bleeding after hemostatic challenges. The presence of petechiae in particular suggests a platelet defect. Quantitative platelet abnormalities in adults are most often acquired; autoimmune (ie, idiopathic thrombocytopenic purpura [ITP]) and drug-induced thrombocytopenia account for the vast majority of cases of isolated thrombocytopenia. Certain acquired thrombocytopenic conditions (eg, disseminated intravascular coagulation [DIC] and thrombotic thrombocytopenic purpura [TTP]/hemolytic
uremic syndrome) may present with bleeding in acutely ill patients (Table 2) but may also be the underlying reason for asymptomatic thrombocytopenia in an ambulatory patient. Congenital (or inherited) thrombocytopenias are usually diagnosed during childhood, but diagnosis could be delayed into adulthood, especially in individuals who do not regularly obtain health care. Numerous inherited thrombocytopenias exist; individual conditions may be identified based on platelet size, coexisting physical or laboratory abnormalities, and the presence of defective platelet function as well as an abnormal platelet count.8,17,18 In most inherited forms, thrombocytopenia is mild: bleeding occurs only occasionally or after hemostatic challenge.17 A family history of thrombocytopenia (including thrombocytopenia erroneously attributed to other causes such as ITP16) may suggest an inherited thrombocytopenia.19

Qualitative platelet disorders presenting in adults can be caused by medication (eg, aspirin and nonsteroidal anti-inflammatory drugs [NSAIDs]), uremia, cirrhosis, and myeloproliferative disorders.3,20 Several inherited disorders of platelet function exist as well and are classified in various ways (by defective platelet function [eg, adhesion or aggregation] or platelet component [eg, receptors, granules, or membrane phospholipids]).8,15 Although severe inherited disorders of platelet receptors (eg, Glanzmann thrombasthenia and Bernard-Soulier syndrome) and some platelet granule disorders typically present earlier in life, the majority of inherited platelet function disorders present during adulthood, often after hemostatic challenge.8,21

VWD may also be considered a disorder of platelet function, given the role of von Willebrand factor (VWF) in platelet adhesion and aggregation.22 Consequently, VWD tends to present with clinical signs and symptoms similar to those of
<table>
<thead>
<tr>
<th>Mechanism of Thrombocytopenia</th>
<th>Differential Diagnosis</th>
<th>Comments</th>
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<tr>
<td>Impaired production</td>
<td>Hematologic malignancies&lt;sup&gt;20&lt;/sup&gt;</td>
<td>Often accompanied by abnormalities in other marrow cell lines (i.e., red and white blood cells)</td>
</tr>
<tr>
<td></td>
<td>Aplastic anemia&lt;sup&gt;41&lt;/sup&gt;</td>
<td>Other marrow cell lines affected as well</td>
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<td></td>
<td>Myelodysplasia&lt;sup&gt;20&lt;/sup&gt;</td>
<td>May be accompanied by abnormalities in other marrow cell lines; bleeding may occur at higher platelet counts than expected</td>
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<td></td>
<td>Drugs/toxins&lt;sup&gt;41,42&lt;/sup&gt;</td>
<td>For example, alcohol, chemotherapeutic agents, radiation</td>
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<td></td>
<td>Viral marrow suppression or damage&lt;sup&gt;42&lt;/sup&gt;</td>
<td>For example, because of EBV, parvovirus, HCV, or HIV</td>
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<td></td>
<td>Gestational thrombocytopenia&lt;sup&gt;43&lt;/sup&gt;</td>
<td>Generally mild [i.e., platelet count &gt;70,000/mL] and self-limited, resolving after delivery; exact mechanism of thrombocytopenia unknown—hemodilution and increased platelet turnover may also contribute</td>
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<td></td>
<td>Liver disease&lt;sup&gt;42&lt;/sup&gt;</td>
<td>Because of reduced levels of thrombopoietin, which is produced by the liver</td>
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<td></td>
<td>Nutritional deficiencies&lt;sup&gt;42&lt;/sup&gt;</td>
<td>For example, folate, vitamin B&lt;sub&gt;12&lt;/sub&gt;</td>
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<td></td>
<td>Inherited thrombocytopenias&lt;sup&gt;16,17,19,41&lt;/sup&gt;</td>
<td>Often present with incidental thrombocytopenia in adulthood; may have family history of thrombocytopenia or personal history of low platelet counts</td>
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<tr>
<td>Destruction or consumption</td>
<td>Immune</td>
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<td></td>
<td>Medication, including heparin (most common), various antimicrobial, antiarrhythmic, anticonvulsant, and antifungal agents, and H2 receptor antagonists&lt;sup&gt;91&lt;/sup&gt;</td>
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<td></td>
<td>ITP</td>
<td>Along with drug-induced thrombocytopenia, accounts for majority of isolated thrombocytopenia in adults&lt;sup&gt;16&lt;/sup&gt;; typically chronic in adults&lt;sup&gt;41&lt;/sup&gt;; may occasionally be accompanied by Coombs positive hemolytic anemia [Evans syndrome]&lt;sup&gt;41&lt;/sup&gt;</td>
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<td>Autoimmune disease&lt;sup&gt;41&lt;/sup&gt;</td>
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<td>Infection&lt;sup&gt;41,45&lt;/sup&gt;</td>
<td>For example, HIV</td>
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<td></td>
<td>TTP in presence of ADAMTS13 autoantibodies&lt;sup&gt;41,46&lt;/sup&gt;</td>
<td>Coombs-negative hemolytic anemia and thrombocytopenia; may or may not have associated renal insufficiency, fever, and mental status changes; neurological symptoms vary, ranging from headache and confusion to seizures and stroke-like symptoms</td>
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<tr>
<td>Nonimmune</td>
<td>HUS</td>
<td>Relatively uncommon but life-threatening cause of thrombocytopenia; classic form consists of microangiopathic hemolytic anemia, thrombocytopenia, and renal failure</td>
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<td>DIC&lt;sup&gt;41&lt;/sup&gt;</td>
<td>Other hallmark laboratory findings include decreased fibrinogen, elevated fibrin degradation products, and a positive D-dimer</td>
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<td>Sepsis&lt;sup&gt;42&lt;/sup&gt;</td>
<td>Serious intrapartum condition characterized by hemolysis and elevated liver enzymes in addition to thrombocytopenia; frequently coexists with preeclampsia; may recur in subsequent pregnancies</td>
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<td>HELLP syndrome&lt;sup&gt;44&lt;/sup&gt;</td>
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<td>Physical destruction&lt;sup&gt;41,42&lt;/sup&gt;</td>
<td>For example, valvar disease, cardiopulmonary bypass, cavernous hemangiomas (e.g., in Kassabach-Merritt syndrome)</td>
</tr>
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<td>Sequestration</td>
<td>Splenomegaly&lt;sup&gt;41,42&lt;/sup&gt;</td>
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ADAMTS, a disintegrin and metalloproteinase with thrombospondin type 1 motives; DIC, disseminated intravascular coagulation; EBV, Epstein-Barr virus; HCV, hepatitis C virus; HELLP, hemolysis, elevated liver enzymes, and low platelets; HIV, human immunodeficiency virus; HUS, hemolytic uremic syndrome; ITP, idiopathic thrombocytopenic purpura; TTP, thrombotic thrombocytopenic purpura.
platelet disorders. In patients with a relevant history (eg, significant mucocutaneous bleeding, family history), initial hematologic laboratory evaluation should include specific testing for VWD (VWF antigen, VWF ristocetin cofactor activity, and factor VIII activity assays); other screening tests such as activated partial thromboplastin time (aPTT) and bleeding time or a platelet function analyzer (PFA-100; Siemens Health care Diagnostics Inc., Tarrytown, NY) may miss VWD. Routine hemostatic screening laboratories may likewise miss other platelet function defects. Consequently, in patients with mucocutaneous bleeding who do not have thrombocytopenia or VWD, platelet aggre-gometry should be considered as an initial test for assessing platelet function, in addition to evaluation of a peripheral blood smear for abnormalities in platelet morphology that are specific to certain conditions (eg, gray platelet syndrome).

In contrast to the superficial bleeding associated with platelet defects, coagulation factor defects result in delayed, deep bleeding, for example, into muscles or joints, as well as deep soft-tissue and mucocutaneous bleeding. Patients with milder congenital deficiencies or those with certain specific congenital deficiencies (eg, factor XI deficiency) are more likely to bleed after hemostatic challenge.

Menorrhagia is a common bleeding symptom in women, both those with and without bleeding disorders. Menorrhagia is the most common bleeding symptom in women with inherited bleeding disorders, particularly menorrhagia that begins at menarche and persists into adulthood. When menorrhagia begins after the age of 20 years, acquired bleeding diatheses should be considered, as should nonhematologic causes such as uterine pathology (eg, fibroids), hypothyroidism, and, in women older than 40 years of age, perimenopausal anovulation. Consensus recommendations for the evaluation of acute menorrhagia have been published. They include assessments of the patient’s menstrual, bleeding, medication, and family histories; speculum and pelvic examinations with subsequent Papanicolaou test and endometrial biopsy, as appropriate, based on such factors as patient age and feasibility of performing these interventions through heavy menstrual bleeding; and (preferably intravaginal) ultrasound. Recommended initial laboratory testing includes a complete blood count, pregnancy test, prothrombin time (PT), aPTT, fibrinogen, and, if feasible, VWF levels. Additional studies may include tests of liver or platelet function or specific factor levels, as clinically indicated. Samples may also be drawn for storage for future testing, especially if administration of transfusional therapies (eg, fresh frozen plasma [FFP]) is anticipated.

**Hematologic Laboratory Abnormalities**

Once a significant bleeding history is identified, an initial laboratory evaluation is generally undertaken to determine the underlying cause. Alternatively, an adult with an undiagnosed bleeding disorder may present with abnormal hematologic laboratory studies obtained as part of an evaluation for surgery or for some other reason. Increased sensitivity of the reagents used in coagulation assays, most notably PT and aPTT, has led to an increased incidence in abnormal results for these tests. Screening coagulation laboratory studies have a low yield overall in the absence of any symptoms or family history of an underlying bleeding disorder. Even among patients with a high pretest probability of having a bleeding disorder, only a minority of abnormalities correspond with a clinically significant bleeding diathesis. Failure to identify the subset of individuals whose abnormal coagulation studies signify an as yet undiagnosed bleeding disorder, however, may have serious, if not grave, consequences, particularly if an invasive intervention is planned.

**Abnormal Coagulation Assays**

Prolongation of the aPTT or PT may indicate an acquired or congenital clotting factor deficiency or an inhibitor of one or more coagulation factors. Potential inhibitors include medication (namely anticoagulants), antibodies directed against specific coagulation factors, and nonspecific inhibitors (eg, lupus anticoagulants). A mixing study can be used to differentiate a deficiency from an inhibitor. In a mixing study, equal volumes of normal and patient plasma are combined, and then the coagulation study is repeated. In cases of coagulation factor deficiency, the presence of normal plasma replaces the missing factor(s), thereby normalizing the abnormal coagulation study. In contrast, when an inhibitor is present the abnormality persists after the addition of normal plasma. In some cases, a prolonged incubation period after mixing is necessary for accurate interpretation of results; therefore, it is imperative that the mixing study be incubated at 37°C for 2 hours. Readily identifiable
causes of coagulation study abnormalities (eg, anticoagulant medication, systemic diseases such as liver disease, or artifactual prolongation [eg, caused by sample “contamination” with heparin]) should ideally be excluded before proceeding to a mixing study.29

The potential coagulation factors involved and, therefore, possible diagnoses can be narrowed down based on which coagulation study is abnormal29–32 (Figures 1 and 2). Isolated prolongation of the aPTT indicates an abnormality of the intrinsic pathway (ie, of prekallikrein, high-molecular-weight kininogen, factor VIII [FVIII], factor IX, factor XI, or factor XII [FXII]).29,30 An isolated prolonged aPTT can also indicate a lupus anticoagulant. An isolated prolonged PT indicates an abnormality of the extrinsic pathway (ie, factor VII [FVII]).29,30 On occasion, congenital deficiencies of the final common pathway factors II (FII, also known as prothrombin), V (FV), and X (FX) and fibrinogen

Figure 2.29–32 Differential diagnosis for abnormalities of aPTT and PT. Once the coagulation laboratory study abnormality has been identified, the differential diagnosis may be further narrowed down based on the specific coagulation study abnormalities (activated partial thromboplastin time [aPTT], prothrombin time [PT], or both); the presence or absence of bleeding symptoms; and the results of the mixing study. Note that prolonged incubation may be required for accurate mixing study results. *PT may also be prolonged by heparin (at high doses) or direct thrombin inhibitor (DTIs). †PT may also be prolonged by FII and FX deficiencies. ‡aPTT may also be prolonged in the setting of advanced liver disease or vitamin K deficiency. §Applies to 10% of lupus anticoagulants. DIC, disseminated intravascular coagulation; FII, factor II; FIX, factor IX; FV, factor V; FVII, factor VII; FVIII, factor VIII; FX, factor X; FXI, factor XI; FXII, factor XII; HMWK, high-molecular-weight kininogen; VWD, von Willebrand disease.
present with an isolated prolonged PT and a normal aPTT. Not all conditions that prolong aPTT or PT are associated with a bleeding phenotype (Figure 2). For example, lupus anticoagulant is more likely to be associated with thrombosis than with bleeding, except in rare cases of associated antiprothrombin antibodies, which lead to bleeding symptoms and a prolonged PT in addition to prolonged aPTT.\textsuperscript{3,2} Deficiencies of the contact-activating factors (FXII, prekallikrein, and high-molecular-weight kininogen) are rare and do not cause easy bleeding; however, they are associated with markedly prolonged aPTT\textsuperscript{s}. When asked to evaluate an asymptomatic patient who has a markedly prolonged aPTT, testing for lupus anticoagulant should be considered first when the mixing study fails to correct and for FXII deficiency when the aPTT corrects in the mixing study.

Prolongation of both aPTT and PT isolates the abnormality to the final common pathway, consisting of FV, FX, FII, and fibrinogen.\textsuperscript{2,3,2} Congenital or acquired deficiencies of any of these factors may present with a prolonged PT and aPTT. Acquired deficiencies of single coagulation factors may occur in the setting of systemic diseases such as amyloidosis (FX) and myeloproliferative disease (FV) and must be differentiated from congenital deficiencies.\textsuperscript{2} Deficiencies of multiple factors from both the intrinsic and extrinsic pathways or from all 3 pathways may also simultaneously prolong aPTT and PT.\textsuperscript{2} Multiple factor deficiencies may occur as a result of severe liver disease, supratherapeutic warfarin doses resulting in deficiency of vitamin K–dependent factors, or consumptive coagulopathy (ie, DIC, which generally occurs in the setting of systemic illness and is therefore unlikely to present solely with asymptomatic coagulation study abnormalities). Potential inhibitors that may present with prolonged aPTT and PT include heparin, direct thrombin inhibitors, potent lupus anticoagulants, and other nonspecific inhibitors such as those associated with lymphoproliferative disorders or monoclonal protein disorders. Bleeding symptoms are generally a feature of all conditions that simultaneously prolong PT and aPTT.

Vitamin K deficiency and liver disease both may result in prolongation of PT or, in more advanced stages, of both PT and aPTT. The vitamin K–dependent coagulation factors (II, VII, IX, and X) may become depleted because of malabsorption, prolonged antibiotic use, or warfarin therapy.\textsuperscript{2} Liver disease is distinguished from vitamin K deficiency by a deficiency of FV in addition to the vitamin K–dependent factors.\textsuperscript{2} Liver disease is often quite advanced by the time abnormalities in coagulation laboratory studies are present; therefore, patients with liver disease are unlikely to present solely with asymptomatic coagulation laboratory abnormalities and often have concurrent physical signs (eg, jaundice, hepatomegaly) or other laboratory abnormalities indicative of impaired hepatic function (eg, thrombocytopenia, hypoalbuminemia, transaminitis).

Acquired coagulation factor inhibitors (or autoantibodies), most commonly directed against FVIII (a condition referred to as acquired hemophilia), deserve special mention because they may be associated with serious bleeding in adults with no history of bleeding. Acquired hemophilia is a rare condition (incidence of 1 to 4 per million per year\textsuperscript{3} ) that predominately affects older adults. In the largest collection of affected patients to date (n = 501), the median age at diagnosis was 74 years; however, younger women in particular may be affected as well because of an association with pregnancy.\textsuperscript{3,2} In approximately half of cases, a coexisting underlying condition such as pregnancy is identified,\textsuperscript{3} some of which (eg, cancer, autoimmune disease) are characterized by immune dysregulation. Acquired hemophilia should be suspected in an adult with new- or recent-onset bleeding who has no personal or family history of bleeding and presents with an isolated prolonged aPTT that does not correct in a mixing study.\textsuperscript{3,2} Acquired hemophilia was associated with an especially high mortality—up to 41% in untreated patients\textsuperscript{3,2} and 6% to 8% among effectively treated patients,\textsuperscript{3,2} mostly because of rebleeding. Because acquired hemophilia requires specialized treatment, prompt diagnosis is important, particularly when an invasive procedure is necessary.\textsuperscript{3,2}

**Thrombocytopenia**

Thrombocytopenia is defined as a platelet count below the lower limit of the normal range (ie, <150,000/μL in most laboratories). In asymptomatic patients, artificial thrombocytopenia as a result of platelet clumping may first be excluded by examining the peripheral blood smear.\textsuperscript{4} Thrombocytopenia occurs because of impaired production,\textsuperscript{1,3,3} destruction or consumption,\textsuperscript{1,4,2,4,4} or sequestration\textsuperscript{4,4} of platelets.
Immune thrombocytopenias, either idiopathic (primary) or secondary to an autoimmune disease, may present with asymptomatic, isolated thrombocytopenia. Other potential causes in ambulatory patients include medication; infectious agents such as Epstein-Barr, human immunodeficiency, or hepatitis C virus; or primary marrow failure. Conversely, certain thrombocytopenic conditions may be excluded based on the lack of specific predisposing factors or acute illness (e.g., shiga toxin–induced hemolytic uremic syndrome or DIC). Bleeding propensity in thrombocytopenic conditions usually depends on the platelet count. Bleeding is generally mild and limited to easy bruising when platelet counts exceed 20,000/μL. The risk for spontaneous bleeding increases only after the platelet count decreases to 10,000/μL, except in ITP, in which the increased presence of young, hyperfunctional platelets may preserve hemostasis even at platelet counts below this level.

Abnormal Platelet Function
Historically, bleeding time was used as a screening test for qualitative platelet abnormalities, including in patients undergoing invasive procedures, particularly those who were recently exposed to medication that might alter platelet function (e.g., aspirin or NSAIDs). This test was also used to screen for certain bleeding disorders, including VWD. However, bleeding time has been shown to be relatively insensitive and poorly reproducible. The test has not been shown to predict excessive surgical bleeding (particularly without other history suggesting a bleeding disorder) or to reliably identify aspirin- or NSAID-induced platelet dysfunction. In addition, some patients with VWD have a normal bleeding time. Therefore, this test is no longer routinely recommended as part of an initial hemostatic laboratory evaluation, especially to screen for VWD. For assessing platelet function, the PFA-100 has supplanted the bleeding time in many laboratories. In this technique, a sample of whole blood is passed through an aperture in a membrane coated with collagen and either epinephrine or adenosine diphosphate, and the amount of time it takes for the membrane to become occluded (i.e., the closure time) is measured. Although the PFA-100 has been shown to have a relatively high sensitivity for detecting moderate and severe VWD, aspirin- and NSAID-related platelet dysfunction, and severe platelet function disorders, closure times may be normal in milder VWD and platelet function disorders, including relatively common platelet storage pool deficiencies, thereby limiting the usefulness of the PFA-100 as a screening tool for all platelet function disorders. If a patient is suspected of having a significant platelet function disorder, consultation with a hematologist is suggested to obtain more extensive platelet aggregation and release studies in a specialized laboratory setting.

Normal Initial Hemostatic Laboratory Studies
Of note, several bleeding disorders are associated with normal routine initial hemostatic laboratory studies (i.e., platelet count, PT, aPTT, and PFA-100) (Table 3). In some cases, the sensitivity of these studies for detecting certain conditions, such

<table>
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<th>Condition</th>
<th>Recommended Screening Tests</th>
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<td>VWD*</td>
<td>VWF antigen, VWF ristocetin cofactor activity, and FVIII activity assays</td>
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<tr>
<td>FXIII deficiency</td>
<td>Quantitative functional FXIII assay (ammonia-release or amine-incorporation assay)</td>
</tr>
<tr>
<td>Fibrinolytic disorders (α2-antiplasmin and PAI-1 deficiencies)</td>
<td>Specific functional (activity) and antigen assays</td>
</tr>
<tr>
<td>Collagen disorders (e.g., EDS, BJHS)</td>
<td>Hypermobility assessment tools (e.g., Beighton score, Brighton criteria)</td>
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*Initial hematologic laboratory studies will be normal in only some cases of von Willebrand disease (VWD). Nevertheless, additional screening tests should be performed before excluding VWD based on a normal activated partial thromboplastin time or Platelet Function Analyzer, when clinically indicated. BJHS, benign joint hypermobility syndrome; EDS, Ehlers-Danlos syndrome; FXIII, factor XIII; FVIII, factor VIII; PAI, plasminogen activator inhibitor; VWF, von Willebrand factor.
as some types of VWD or milder factor deficiencies, may be limited. In other cases, neither fibrin generation nor platelet function is impaired; therefore, coagulation studies and quantitative and qualitative platelet test results are normal. Alternative screening tests are indicated when these bleeding disorders are suspected based on clinical grounds.11,23,56–60 (Table 3).

Therapeutic Strategies in Patients With Bleeding Disorders

Preventive Care and Measures

Some basic guidelines apply for all patients with bleeding disorders in the primary care setting. In general, any medication that may impair hemostatic function, such as aspirin and NSAIDs, should be avoided, particularly during bleeding episodes. Proper preventive care, including screening for common age-related comorbidities and associated risk factors, is important in this population. Certain conditions may be particularly problematic in patients with bleeding disorders; for example, untreated severe hypertension may lead to intracranial bleeding.61 Invasive preventive or screening procedures should not necessarily be deferred because of the risk for bleeding. Routine dental care may be especially important in patients with bleeding disorders to avoid the need for more extensive dental procedures down the line.62,63 Invasive screening procedures such as colonoscopy may be performed with minimal bleeding risk.64 Consultation with a hematologist should be considered before such invasive procedures to determine whether systemic hemostatic coverage is indicated and, if so, what this coverage should consist of. Hemostatic therapy often is not needed for minimally invasive procedures, particularly if no tissue incision or excision is required.65 Meticulous surgical technique and topical measures such as suturing and use of fibrin glue, oxidized cellulose (eg, Surgicel [Ethicon, LLC, San Lorenzo, Puerto Rico]), or topical antifibrinolytics (eg, tranexamic acid) may be sufficient to maintain hemostasis or at least minimize the use of systemic hemostatic treatments in some dental procedures.65–67 Such topical measures may also be used for bleeding from wounds, in addition to closing (when appropriate), applying pressure to, or packing the wound. Packing can also be used to control epistaxis.68 With diligent attention to local hemostasis, systemic treatments can often be avoided in cases of minor bleeding.

Hemostatic Therapies

Consultation with a hematologist is strongly recommended before initiating systemic therapies to discuss treatment options and to ensure that samples for all indicated laboratory studies are collected before any blood products are administered, since this may affect the reliability of any results. Transfusion of the missing or defective hemostatic component may not be indicated or effective in all cases. For example, transfusion of platelets may be ineffective in ITP and, in the absence of life-threatening hemorrhage, is contraindicated in microangiopathic conditions such as TTP and in heparin-induced thrombocytopenia, given the risk for exacerbating these conditions and fueling thrombosis.45,69,70 Similarly, in the setting of liver disease, infusion of large volumes of FFP to treat presumptive coagulopathy may exacerbate bleeding that is a result of elevated venous pressure (eg, variceal bleeding) by further increasing this pressure.71 In addition, factor replacement may be ineffective in factor deficiencies because of an acquired inhibitor (antibody against the clotting factor concentrate). Other specialized treatments or interventions may be indicated or advised for immediate management of specific bleeding disorders (eg, plasma exchange for TTP or immunomodulatory therapies for immune-mediated thrombocytopenia).

Platelet Transfusion

Indications for platelet transfusion in patients with thrombocytopenia vary based on the underlying mechanism of thrombocytopenia.41,69,70 Thrombocytopenia as a result of reduced production is an indication for platelet transfusion, and the thresholds for transfusion vary based on the indication (Table 4). There are no such thresholds for platelet transfusion in patients with immune-mediated thrombocytopenia or platelet function defects.69 Platelet transfusion should only be undertaken in patients with autoimmune thrombocytopenia in instances of serious or life-threatening bleeding (eg, gastrointestinal or intracranial hemorrhage).69,70 Large amounts of platelets may be required to boost the platelet count in the setting of autoimmune thrombocytopenia, given the shortened survival of the transfused platelets; concomitant ad-
administration of immune-modulating therapies such as steroids or intravenous immunoglobulin may attenuate this process, resulting in a more rapid increase in platelet count. Patients with inherited and acquired platelet function abnormalities rarely require platelet transfusion. Alternative measures are proposed as first-line therapies in patients with platelet function defects who are actively bleeding or who require surgery (Table 4), except in cases of severe inherited disorders such as Glanzmann thrombasthenia or Bernard-Soulier syndrome.

Human leukocyte antigen–matched platelets may be used in patients requiring multiple transfusions, given the possibility for alloimmunization. If human leukocyte antigen–matched platelets are not available, leukocyte-depleted concentrates may be used.

**Coagulation Factor Replacement**

For known or suspected coagulation factor deficiencies, factor replacement may be used to treat active bleeding or for hemostatic coverage before surgery. The indications for and necessity of factor replacement vary based on the specific bleeding disorder, the patient’s bleeding tendency, the severity of bleeding, and, in the case of invasive procedures, the anticipated risk for bleeding. Because coagulation studies do not necessarily predict surgical bleeding risk in patients without an apparent bleeding disorder, the prophylactic use of factor replacement sources such as FFP simply to correct coagulation study abnormalities before surgery is inappropriate. In fact, prophylactic transfusion of FFP has not been shown to correct coagulation study abnormalities or reduce bleeding risk.

Several options exist for replacing coagulation factors when indicated for treatment or prevention of bleeding (Table 5). Given their wide availability, FFP and cryoprecipitate are mainstays of coagulation factor replacement worldwide. FFP is obtained from whole blood or by apheresis and contains all clotting factors. Cryoprecipitate is derived from thawing a single donor unit of FFP at 4°C and is rich in FVIII, VWF, FXIII, and fibrinogen.
ogen. FFP and cryoprecipitate are primarily indicated for replacement of multiple deficient clotting factors (eg, in DIC) and should be used only to replace single clotting factors when no specific concentrate is available. Large volumes of FFP are required to sufficiently boost a single coagulation factor, putting some patients at risk for fluid overload. In addition, despite substantial improvements in donor and plasma screening, transmission of blood-borne pathogens continues to be a theoretical concern with FFP and cryoprecipitate because these products are not subjected to a specific viral inactivation process. Therefore, virally inactivated FFP should be considered when available. For single-factor deficiencies, specific plasma-derived or recombinant factor concentrates are the treatment of choice, when available (Table 5). There are currently no single-factor concentrates for FII, FV, or FX. Prothrombin complex concentrates (PCCs), which are highly purified contraceptive, have a high risk of thrombogenicity and are indicated for warfarin reversal.

### Table 5. Factor Replacement Sources

<table>
<thead>
<tr>
<th>Product</th>
<th>Composition</th>
<th>Indications</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>FFP</td>
<td>All coagulation factors</td>
<td>Multiple factor deficiencies (eg, in DIC, liver disease)</td>
<td>May be required in large volumes to sufficiently boost levels of a single coagulation factor</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FV deficiency (may supplement with platelet transfusions)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>FXI deficiency (in lieu of FXI concentrate when there is a heightened risk of thrombogenicity [eg, in the peripartum period])</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Other single-factor deficiencies when single-factor concentrates or PCCs are unavailable</td>
<td></td>
</tr>
<tr>
<td>Cryoprecipitate</td>
<td>Concentrate precipitated from FFP; rich in FVIII, VWF, FXIII, and fibrinogen</td>
<td>Used most often to replace fibrinogen</td>
<td>May be required in large volumes</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Not virally inactivated</td>
</tr>
<tr>
<td>PCCs</td>
<td>Highly purified concentrates from pooled normal plasma containing FII, FIX, and FX (±FVII)</td>
<td>FII or FX deficiency</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>FIX or FVII deficiencies when single-factor concentrates are unavailable</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>4-factor PCCs are indicated for warfarin reversal</td>
<td></td>
</tr>
<tr>
<td>Activated PCCs</td>
<td>PCC with FII, FIX, and FX (mainly nonactivated) and FVII (mainly in the activated form)</td>
<td>Coagulation factor inhibitors</td>
<td></td>
</tr>
<tr>
<td>Single-factor concentrates</td>
<td>Plasma derived</td>
<td>Respective single-factor deficiencies</td>
<td>Treatment of choice for single-factor deficiencies, when available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rFVIIa is indicated as a bypassing agent for FVIII and FIX inhibitors, as well as for replacing FVII in FVII deficiency</td>
<td>Risk for human viral contamination is present but exceedingly low in plasma-derived concentrates because of screening and viral inactivation procedures; it is probably nonexistent in recombinant concentrates</td>
</tr>
</tbody>
</table>

*Only 4-factor prothrombin complex concentrates (PCCs) may be used for replacement of factor VII (FVII); however, 4-factor PCCs are not universally available. Although single-factor concentrates may be used for respective low-titer coagulation factor inhibitors (eg, plasma-derived or recombinant factor VIII [FVIII] for FVIII inhibitors), bypassing agents are generally recommended for active bleeding (for which they are considered first-line therapy) and for hemostatic coverage during invasive procedures in patients with inhibitors. DIC, disseminated intravascular coagulation; FFP, fresh frozen plasma; F, factor; PCC, prothrombin complex concentrate; rF, recombinant factor; VWF, von Willebrand factor.*
trates of specific coagulation factors (FII, factor IX, and FX ± FVII) obtained from pooled normal plasma, should be used in lieu of FFP for deficiencies of FII or FX, when available, given that PCCs are virally inactivated and contain known amounts of each factor.74,75 Absent any other source, FFP is the only option for FV replacement; platelet transfusions may be used adjunctively with platelet transfusion and other nontransfusional therapies in these patients63.

Theoretical risk for thrombogenesis: may be increased in patients who receive concurrent factor replacement or who have certain bleeding disorders (eg, dysfibrinogenemia, in which antifibrinolytics are contraindicated)

**Desmopressin**

Transiently increases VWF and FVIII levels89 and augments platelet adhesiveness and aggregation68

Agent of choice for treatment or prevention of bleeding in most patients with type 1 VWD; however, patients with type 3 VWD and most patients with type 2 VWD typically require factor concentrates containing FVIII and VWF90

Other desmopressin-responsive bleeding disorders include mild hemophilia and platelet function disorders, in which the use of desmopressin for surgical coverage has been described11; also commonly used for uremic bleeding98

Specifically used for treatment of menorrhagia in women with bleeding disorders25,80,91

Patients, especially children, receiving desmopressin should be closely monitored for hyponatremia22,87

**Vitamin K**

May be used in vitamin K deficiency and for bleeding caused by overmedication with warfarin

Widely available and inexpensive

Should be given intravenously or orally (not subcutaneously) for best absorption and fastest response92–94

FVIII, factor VIII; PAI, plasminogen activator inhibitor; VWD, von Willebrand disease; VWF, von Willebrand factor.

Nontransfusional Hemostatic Therapies

Nontransfusional therapies such as antifibrinolytics (ε-aminocaproic acid or tranexamic acid),1,2,5,35,56–68,73,78–86 desmopressin,21,22,25,35,56,68,77,80,87–91 and vitamin K92–94 may be used for the treatment or prevention of bleeding in patients with bleeding disorders (Table 6) and occasionally as sole hemostatic agents for the treatment of mild mucosal bleeding or for hemostatic coverage during minor (eg, dental) procedures, thus sparing the patient exposure to transfusional therapies. Nontransfusional therapies can also be used as an adjunct to transfusional therapies in cases of more severe bleeding or for hemostatic coverage during major surgery.

Menorrhagia in women with bleeding disorders often is successfully managed with antifibrinolytics and hormone therapy,85 including combination or progestin-only contraceptives for maintenance ther-
apy and conjugated estrogens for acute menorrhagia. Transfusional therapies may be used in refractory cases. Invasive options for controlling menorrhagia include the levonorgestrel-releasing intrauterine system, endometrial balloon tamponade, dilatation and curettage, and non-fertility-sparing procedures such as endometrial ablation, uterine artery embolization, and hysterectomy.

Conclusions
Adults with undiagnosed bleeding disorders may initially present to their primary care provider with asymptomatic hematologic laboratory abnormalities or with abnormal bleeding symptoms in the outpatient or inpatient setting. Accordingly, family physicians should be prepared to recognize potential hallmarks of an underlying bleeding disorder and to initiate a proper workup and, in some cases, treatment, particularly if there is active bleeding. Consultation with a hematologist may be considered to assist with definitive diagnosis and, for patients who are ultimately diagnosed with a bleeding disorder, individualized long-term hemostatic management. The family physician serves as a crucial member of the multidisciplinary team providing care to adults with bleeding disorders and may play an especially important role in prioritizing routine and disease-related preventive care in this population.

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References

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