EDUCATIONAL OBJECTIVE: Readers will decide whether to restart anticoagulation after a bleeding event on the basis of individual patient factors

Resuming anticoagulation after hemorrhage: A practical approach

ABSTRACT

Most patients who suffer a hemorrhage while on long-term anticoagulant therapy continue to be at risk of thrombosis. Physicians often need to reconsider the need for anticoagulation in view of the risk of recurrent bleeding, and when anticoagulation needs to be resumed, they must also consider the timing and strategy. Since there are no evidence-based guidelines for these situations, the authors of this paper offer a practical framework for individualizing the resumption of anticoagulation after hemorrhage.

KEY POINTS

Not all patients on anticoagulation at the time of a bleeding event have a strong indication to continue anticoagulation afterward.

Important considerations when deciding whether to resume anticoagulation after hemorrhage are whether the source of bleeding has been found and controlled and, if the patient is receiving warfarin, whether he or she can be expected to maintain the target international normalized ratio.

The newer oral anticoagulants, including factor Xa inhibitors and direct thrombin inhibitors, lack antidotes or reversal agents, and their risk of causing bleeding compared with warfarin varies by site of bleeding.

I F A PATIENT receiving anticoagulant therapy suffers a bleeding event, the patient and physician must decide whether and how soon to restart the therapy, and with what agent.

Foremost on our minds tends to be the risk of another hemorrhage. Subtler to appreciate immediately after an event is the continued risk of thrombosis, often from the same medical condition that prompted anticoagulation therapy in the first place (TABLE 1).

Complicating the decision, there may be a rebound effect: some thrombotic events such as pulmonary embolism and atrial fibrillation-related stroke may be more likely to occur in the first weeks after stopping warfarin than during similar intervals in patients who have not been taking it.1-3 The same thing may happen with the newer, target-specific oral anticoagulants.4-6

Although we have evidence-based guidelines for initiating and managing anticoagulation therapy, ample data on adverse events, and protocols for reversing anticoagulation if bleeding occurs, we do not have clear guidelines on restarting anticoagulation after a hemorrhagic event.

In this article, we outline a practical framework for approaching this clinical dilemma. Used in conjunction with consideration of a patient’s values and preferences as well as input from experts, this framework can help clinicians guide their patients through this challenging clinical decision. It consists of five questions:
• Why is the patient on anticoagulation, and what is the risk of thromboembolism without it?
• What was the clinical impact of the hemorrhage, and what is the risk of rebleeding if anticoagulation is resumed?
RESUMING ANTICOAGULATION

Some indications for anticoagulation pose a higher risk of thromboembolism than others

- What additional patient factors should be taken into consideration?
- How long should we wait before restarting anticoagulation?
- Would a newer drug be a better choice?

**BLEEDING OCCURS IN 2% TO 3% OF PATIENTS PER YEAR**

Most of our information on anticoagulation is about vitamin K antagonists—principally warfarin, in use since the 1950s. Among patients taking warfarin outside of clinical trials, the risk of major bleeding is estimated at 2% to 3% per year.7

However, the target-specific oral anticoagulants rivaroxaban (Xarelto), apixaban (Eliquis), dabigatran (Pradaxa) and edoxaban (Savaysa) are being used more and more, and we include them in our discussion insofar as we have information on them. The rates of bleeding with these new drugs in clinical trials have been comparable to or lower than those with warfarin.5

Postmarketing surveillance is under way.

**WHY IS THE PATIENT ON ANTICOAGULATION? WHAT IS THE RISK WITHOUT IT?**

Common, evidence-based indications for anticoagulation are to prevent complications in patients with venous thromboembolism and to prevent stroke in patients with atrial fibrillation or a mechanical heart valve. Other uses, such as in heart failure and its sequelae, pulmonary hypertension, and splanchnic or hepatic vein thrombosis, have less robust evidence to support them.

When anticoagulation-related bleeding occurs, it is essential to review why the patient is taking the drug and the risk of thromboembolism without it. Some indications pose a higher risk of thromboembolism than others and so argue more strongly for continuing the treatment.

Douketis et al9 developed a risk-stratification scheme for perioperative thromboembolism. We have modified it by adding the CHA2DS2-VASc score (TABLE 2),9–11 and believe it can be used more widely.

**TABLE 1**

Factors influencing the decision to resume anticoagulation after a bleeding event

<table>
<thead>
<tr>
<th>Arguing for resuming anticoagulation</th>
<th>Arguing against resuming anticoagulation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strong or near-absolute indication for anticoagulation</td>
<td>No absolute indication for ongoing anticoagulation</td>
</tr>
<tr>
<td>Mechanical mitral valve</td>
<td>Near completion of planned anticoagulation course</td>
</tr>
<tr>
<td>Hypercoagulable state</td>
<td>High risk of rebleeding or presence of additional risk factors for bleeding</td>
</tr>
<tr>
<td>CHADS2 score &gt; 5 (see TABLE 2)</td>
<td>Anticipated high risk of morbidity or death if rebleeding occurs</td>
</tr>
<tr>
<td>CHA2DS2-VASc score &gt; 6 (see TABLE 2)</td>
<td>Anticipated low risk of rebleeding with successful source control and international normalized ratio target control</td>
</tr>
</tbody>
</table>

**High-risk indications**

Conditions that pose a high risk of thrombosis almost always require restarting anticoagulation. Here, the most appropriate question nearly always is not if anticoagulation should be restarted, but when. Examples:
- A mechanical mitral valve
- Antiphospholipid antibody syndrome with recurrent thromboembolic events.

**Lower-risk indications**

Lower-risk indications allow more leeway in determining if anticoagulation should be resumed. The most straightforward cases fall well within established guidelines. Examples:
- Atrial fibrillation and a CHA2DS2-VASc score of 1. The 2014 guidelines from the American College of Cardiology, American Heart Association, and Heart Rhythm Society10 suggest that patients with nonvalvular atrial fibrillation and a CHA2DS2-VASc score of 1 have three options: an oral anticoagulant, aspirin, and no antithrombotic therapy. If such a patient on anticoagulant therapy subsequently experiences a major gastrointestinal hemorrhage requiring transfusion and intensive care and no definitively treatable source of bleeding is found on endoscopy, one can argue that the risks of continued anticoagulation (recurrent bleeding) now exceed the benefits and that the patient would be better served by aspirin or even no antithrombotic therapy.
After 6 months of anticoagulation for unprovoked deep vein thrombosis.

Several studies showed that aspirin reduced the risk of recurrent venous thromboembolism in patients who completed an initial 6-month course of anticoagulation.12–15 Though these studies did not specifically compare aspirin with warfarin or target-specific oral anticoagulants in preventing recurrent venous thromboembolism after a hemorrhage, it is reasonable to extrapolate their results to this situation.

If the risk of recurrent hemorrhage on anticoagulation is considered to be too great, then aspirin is an alternative to no anticoagulation, as it reduces the risk of recurrent venous thromboembolism.16 However, we advise caution if the bleeding lesion may be specifically exacerbated by aspirin, particularly upper gastrointestinal ulcers.

**TABLE 2**

<table>
<thead>
<tr>
<th>Thromboembolic risk by anticoagulation indication</th>
<th>Mechanical heart valve</th>
<th>Atrial fibrillation</th>
<th>Venous thromboembolism</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>High</strong></td>
<td>Any mitral valve prosthesis</td>
<td>CHADS₂, score of 5 or 6</td>
<td>Venous thromboembolism within past 2 months</td>
</tr>
<tr>
<td></td>
<td>Any caged-ball or tilting-disc aortic valve prosthesis</td>
<td>CHADS₂-VASc score of 6 to 9 (suggesting adjusted stroke rate ≥ 9% per year)</td>
<td>Severe thrombophilia (eg, deficiency of protein C, protein S, or antithrombin; antiphospholipid antibodies; multiple abnormalities)</td>
</tr>
<tr>
<td></td>
<td>Stroke or transient ischemic attack within past 6 months</td>
<td>Stroke or transient ischemic attack within past 3 months</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rheumatic valvular heart disease</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Moderate</strong></td>
<td>Bileaflet aortic valve prosthesis and one or more of the following risk factors: atrial fibrillation, prior stroke or transient ischemic attack, hypertension, diabetes, congestive heart failure, age &gt; 75</td>
<td>CHADS₂, score of 3 or 4</td>
<td>Venous thromboembolism within the past 6 months</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CHADS₂-VASc score of 5 (suggesting adjusted stroke rate of 5%–9% per year)</td>
<td>Nonsevere thrombophilia (eg, heterozygous factor V Leiden or prothrombin gene mutation)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Recurrent venous thromboembolism</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Active cancer (treated within 6 months or palliated)</td>
</tr>
<tr>
<td><strong>Low</strong></td>
<td>Bileaflet aortic valve prosthesis without atrial fibrillation and no other risk factors for stroke</td>
<td>CHADS₂, score of 0 to 2</td>
<td>Venous thromboembolism more than 6 months previously and no other risk factors</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CHADS₂-VASc score of 0 to 4 (suggesting adjusted stroke rate &lt; 5% per year and assuming no prior stroke or transient ischemic attack)</td>
<td></td>
</tr>
</tbody>
</table>

*The valve position affects risk for thromboembolism: the incidence rate for valve thrombosis was 5 times higher in mitral valves than in aortic valves; the incidence rate for embolism was 1.5 times higher in mitral valves than in aortic valves.11*

*High-risk patients may also include those with prior thromboembolism during temporary interruption of vitamin K antagonists (eg, warfarin).*

*Moderate-risk patients may also include those with prior stroke or transient ischemic attack occurring more than 3 months before event.*

CHADS₂ = 1 point each except as noted: Congestive heart failure, Hypertension, Age ≥ 75, Diabetes mellitus, and Stroke or transient ischemic attack (2 points)

CHADS₂-VASc = 1 point each except as noted: Congestive heart failure; Hypertension; Age ≥ 75 (2 points); Diabetes mellitus; prior Stroke, transient ischemic attack, or thromboembolism (2 points); Vascular disease; Age 65–74; Sex category (female)


• After 6 months of anticoagulation for unprovoked deep vein thrombosis. Several studies showed that aspirin reduced the risk of recurrent venous thromboembolism in patients who completed an initial 6-month course of anticoagulation.12–15 Though these studies did not specifically compare aspirin with warfarin or target-specific oral anticoagulants in preventing recurrent venous thromboembolism after a hemorrhage, it is reasonable to extrapolate their results to this situation.

If the risk of recurrent hemorrhage on anticoagulation is considered to be too great, then aspirin is an alternative to no anticoagulation, as it reduces the risk of recurrent venous thromboembolism.16 However, we advise caution if the bleeding lesion may be specifically exacerbated by aspirin, particularly upper gastrointestinal ulcers.
RESUMING ANTICOAGULATION

Moderate-risk indications

• After a partial course of anticoagulation for provoked venous thromboembolism. Suppose a patient in the 10th week of a planned 12-week course of anticoagulation for a surgically provoked, first deep vein thrombosis presents with abdominal pain and is found to have a retroperitoneal hematoma. In light of the risk of recurrent bleeding vs the benefit of resuming anticoagulation for the limited remaining period, her 12-week treatment course can reasonably be shortened to 10 weeks.

The risk of recurrent venous thromboembolism when a patient is off anticoagulation decreases with time from the initial event. The highest risk, estimated at 0.3% to 1.3% per day, is in the first 4 weeks, falling to 0.03% to 0.2% per day in weeks 5 through 12, and 0.05% per day thereafter.17–20

Additionally, a pooled analysis of seven randomized trials suggests that patients with isolated, distal deep vein thrombosis provoked by a temporary risk factor did not have a high risk of recurrence after being treated for 4 to 6 weeks.21 These analyses are based on vitamin K antagonists, though it seems reasonable to extrapolate this information to the target-specific oral anticoagulants.

More challenging are situations in which the evidence supporting the initial or continued need for anticoagulation is less robust, such as in heart failure, pulmonary hypertension, or splanchnic and hepatic vein thrombosis. In these cases, the lack of strong evidence supporting the use of anticoagulation should make us hesitate to resume it after bleeding.

WHAT WAS THE CLINICAL IMPACT?

WHAT IS THE RISK OF REBLEEDING?

Different groups have defined major and minor bleeding in different ways.22,23 Several have proposed criteria to standardize how bleeding events (on warfarin and otherwise) are classified,23–25 but the definitions differ.

Specifically, all agree that a “major” bleeding event is one that is fatal, involves bleeding into a major organ, or leads to a substantial decline in hemoglobin level. However, the Thrombolysis in Myocardial Infarction trials use a decline of more than 5 g/dL in their definition,23,25 while the International Society on Thrombosis and Haemostasis uses 2 g/dL.24

Here, we review the clinical impact of the most common sources of anticoagulation-related hemorrhage—gastrointestinal, soft tissue, and urinary tract—as well as intracerebral hemorrhage, a less common but more uniformly devastating event.27

Clinical impact of gastrointestinal hemorrhage

Each year, about 4.5% of patients taking warfarin have a gastrointestinal hemorrhage, though not all of these events are major.28 Evolving data suggest that the newer agents (particularly dabigatran, rivaroxaban, and edoxaban) pose a higher risk of gastrointestinal bleeding than warfarin.29 Patients may need plasma and blood transfusions and intravenous phytonadione, all of which carry risks, albeit small.

Frequently, endoscopy is needed to find the source of bleeding and to control it. If this does not work, angiographic intervention to infuse vasoconstrictors or embolic coils into the culprit artery may be required, and some patients need surgery. Each intervention carries its own risk.

Clinical impact of soft-tissue hemorrhage

Soft-tissue hemorrhage accounts for more than 20% of warfarin-related bleeding events26; as yet, we know of no data on the rate with the new drugs. Soft-tissue hemorrhage is often localized to the large muscles of the retroperitoneum and legs. Though retroperitoneal hemorrhage accounts for a relatively small portion of soft-tissue hemorrhages, it is associated with high rates of morbidity and death and will therefore be our focus.26

Much of the clinical impact of retroperitoneal hemorrhage is from a mass effect that causes abdominal compartment syndrome, hydroureter, ileus, abscess formation, and acute and chronic pain. At least 20% of cases are associated with femoral neuropathy. It can also lead to deep vein thrombosis from venous compression, coupled with hypercoagulability in response to bleeding. Brisk bleeding can lead to shock and death, and the mortality rate in retroperitoneal hemorrhage is estimated at 20% or higher.30

In many cases, the retroperitoneal hemorrhage will self-tamponade and the blood will be reabsorbed once the bleeding has stopped,
but uncontrolled bleeding may require surgical or angiographic intervention.30

**Clinical impact of urinary tract hemorrhage**

Gross or microscopic hematuria can be found in an estimated 2% to 24% of patients taking warfarin31–33; data are lacking for the target-specific oral anticoagulants. Interventions required to manage urinary tract bleeding include bladder irrigation and, less often, transfusion.31 Since a significant number of cases of hematuria are due to neoplastic disease,32 a diagnostic workup with radiographic imaging of the upper tract and cystoscopy of the lower tract is usually required.31 While life-threatening hematuria is uncommon, complications such as transient urinary obstruction from clots may occur.

**Clinical impact of intracranial hemorrhage**

Intracranial hemorrhage is the most feared and deadly of the bleeding complications of anticoagulation. The incidence in patients on warfarin is estimated at 2% to 3% per year, which is markedly higher than the estimated incidence of 25 per 100,000 person-years in the general population.34 Emerging data indicate that the newer drugs are also associated with a risk of intracranial hemorrhage, though the risk is about half that with vitamin K antagonists.35 Intracranial hemorrhage leads to death or disability in 76% of cases, compared with 3% of cases of bleeding from the gastrointestinal or urinary tract.27

Regardless of the source of bleeding, hospitalization is likely to be required and may be prolonged, with attendant risks of nosocomial harms such as infection.

**Risk of rebleeding**

Given the scope and severity of anticoagulation-related bleeding, there is strong interest in predicting and preventing it. By some estimates, the incidence of recurrent bleeding after resuming vitamin K antagonists is 8% to 13%.22 Although there are several indices for predicting the risk of major bleeding when starting anticoagulation, there are currently no validated tools to estimate a patient’s risk of rebleeding.36

The patient factor that most consistently predicts major bleeding is a history of bleeding, particularly from the gastrointestinal tract. Finding and controlling the source of bleeding is important.26,37 For example, a patient with gross hematuria who is found on cystoscopy to have a urothelial papilloma is unlikely to have rebleeding if the tumor is successfully resected and serial follow-up shows no regrowth. In contrast, consider a patient with a major gastrointestinal hemorrhage, the source of which remains elusive after upper, lower, and capsule endoscopy or, alternatively, is suspected to be from one of multiple angiodysplastic lesions. Without definitive source management, this patient faces a high risk of rebleeding.

With or without anticoagulation, after a first intracranial hemorrhage the risk of another one is estimated at 2% to 4% per year.34 An observational study found a recurrence rate of 7.5% when vitamin K antagonist therapy was started after an intracranial hemorrhage (though not all patients were on a vitamin K antagonist at the time of the first hemorrhage).38

Patients with lobar hemorrhage and those with suspected cerebral amyloid angiopathy may be at particularly high risk if anticoagulation is resumed. Conversely, initial events attributed to uncontrolled hypertension that subsequently can be well controlled may portend a lower risk of rebleeding.34 For other types of intracranial hemorrhage, recurrence rates can be even higher. Irrespective of anticoagulation, one prospective study estimated the crude annual rebleeding rate with untreated arteriovenous malformations to be 7%.39 In chronic subdural hematoma, the recurrence rate after initial drainage has been estimated at 9.2% to 26.5%, with use of anticoagulants (in this case, vitamin K antagonists) being an independent predictor of recurrence.40

**WHAT OTHER PATIENT FACTORS NEED CONSIDERATION?**

**Target INR on warfarin**

An important factor influencing the risk of bleeding with warfarin is the intensity of this therapy.37 A meta-analysis41 found that the risks of major hemorrhage and thromboembolism are minimized if the goal international normalized ratio (INR) is 2.0 to 3.0. When considering resuming anticoagulation after bleeding, make sure the therapeutic target is appropriate.37
The factor that most consistently predicts major bleeding is a history of bleeding, particularly gastrointestinal bleeding.

TABLE 3
Therapeutic ranges for common warfarin indications

<table>
<thead>
<tr>
<th>Indication</th>
<th>Target INR range</th>
<th>Duration of therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Most cases of venous thromboembolism</td>
<td>2–3</td>
<td>3–6 months&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Antiphospholipid antibody syndrome with history of arterial or venous thromboembolic event</td>
<td>2–3</td>
<td>Lifelong</td>
</tr>
<tr>
<td>Bioprosthetic mitral valve</td>
<td>2–3</td>
<td>3 months</td>
</tr>
<tr>
<td>Mechanical aortic valve&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2–3</td>
<td>Lifelong</td>
</tr>
<tr>
<td>Mechanical mitral valve</td>
<td>2.5–3.5</td>
<td>Lifelong</td>
</tr>
<tr>
<td>Mechanical combined mitral and aortic valve</td>
<td>2.5–3.5</td>
<td>Lifelong</td>
</tr>
</tbody>
</table>

<sup>a</sup> Lifelong anticoagulation may be needed for certain conditions such as a first unprovoked episode of venous thromboembolism or recurrent venous thromboembolism, as well as active malignancy. Individualized, patient-specific assessment is required.

<sup>b</sup> Older-generation aortic mechanical valves including Starr-Edwards and mechanical-disc valves other than Medtronic Hall prostheses are thought to be more thrombogenic, and accordingly, many recommend a target INR range of 2.5–3.5. This higher target INR range is also considered for patients with aortic mechanical prostheses who are also at higher risk for thromboembolic events, such as those with atrial fibrillation, previous thromboembolism, and a hypercoagulable state.

INR = international normalized ratio

A patient on anticoagulation for the same indication but who has a history of repeated supratherapeutic levels, poor adherence, or poor access to INR monitoring poses very different concerns about resuming anticoagulation (as well as which agent to use, as we discuss below).

Of note, a high INR alone does not explain bleeding. It is estimated that a workup for gastrointestinal bleeding and gross hematuria uncovers previously undetected lesions in approximately one-third of cases involving warfarin. A similar malignancy-unmasking effect is now recognized in patients using the target-specific oral agents who experience gastrointestinal bleeding. Accordingly, we recommend a comprehensive source evaluation for any anticoagulation-related hemorrhage.

Comorbid conditions
Comorbid conditions associated with bleeding include cancer, end-stage renal disease, liver disease, arterial hypertension, prior stroke, and alcohol abuse. Gait instability, regardless of cause, may also increase the...
risk of trauma-related hemorrhage, but some have estimated that a patient would need to fall multiple times per week to contraindicate anticoagulation on the basis of falls alone.46

Concurrent medications
Concomitant therapies, including antiplatelet drugs and nonsteroidal anti-inflammatory drugs, increase bleeding risk.47,48 Aspirin and the nonsteroidals, in addition to having antiplatelet effects, also can cause gastric erosion.37 In evaluating whether and when to restart anticoagulation, it is advisable to review the role that concomitant therapies may have had in the index bleeding event and to evaluate the risks and benefits of these other agents. Additionally, warfarin has many interactions. Although the newer drugs are lauded for having fewer interactions, they are not completely free of them, and the potential for interactions must always be reviewed.49 Further, unlike warfarin therapy, therapy with the newer agents is not routinely monitored with laboratory tests, so toxicity (or underdosing) may not be recognized until an adverse clinical event occurs. Ultimately, it may be safer to resume anticoagulation after a contributing drug can be safely discontinued.

Advanced age
The influence that the patient’s age should have on the decision to restart anticoagulation is unclear. Although the risk of intracranial hemorrhage increases with age, particularly after age 80, limited data exist in this population, particularly with regard to rebleeding. Further, age is a major risk factor for most thrombotic events, including venous thromboembolism and stroke from atrial fibrillation, so although the risks of anticoagulation may be higher, the benefits may also be higher than in younger patients.37,46 We discourage using age alone as a reason to withhold anticoagulation after a hemorrhage.

■ HOW LONG SHOULD WE WAIT TO RESTART ANTICOAGULATION?

We lack conclusive data on how long to wait to restart anticoagulation after an anticoagulation-associated hemorrhage.

The decision is complicated by evidence suggesting a rebound effect, with an increased risk of pulmonary embolism and atrial fibrillation-related stroke during the first 90 days of interruption of therapy with warfarin as well as with target-specific oral anticoagulants.3–8 In anticoagulation-associated retroperitoneal bleeding, there is increased risk of deep vein thrombosis from compression, even if venous thromboembolism was not the initial indication for anticoagulation.10

In patients with intracranial hemorrhage, evidence suggests that the intracranial hemorrhage itself increases the risk of arterial and venous thromboembolic events. Irrespective of whether a patient was previously on anticoagulation, the risk of arterial and venous thromboembolic events approaches 7% during the initial intracranial hemorrhage-related hospitalization and 9% during the first 90 days.43,10,51

To date, the only information we have about when to resume anticoagulation comes from patients taking vitamin K antagonists.

Timing after gastrointestinal bleeding
Small case series suggest that in the first 2 months after warfarin-associated gastrointestinal bleeding, there is substantial risk of rebleeding when anticoagulation is resumed—and of thrombosis when it is not.52,53 Two retrospective cohort studies may provide some guidance in this dilemma.28,54

Witt et al28 followed 442 patients who presented with gastrointestinal bleeding from any site during warfarin therapy for varied indications for up to 90 days after the index bleeding event. The risk of death was three times lower in patients who restarted warfarin than in those who did not, and their rate of thrombotic events was 10 times lower. The risk of recurrent gastrointestinal bleeding was statistically insignificant, and there were no fatal bleeding events. Anticoagulant therapy was generally resumed within 1 week of the bleeding event, at a median of 4 days.28,55

Qureshi et al54 performed a retrospective cohort study of 1,329 patients with nonvalvular atrial fibrillation who had experienced a gastrointestinal hemorrhage while taking warfarin. They found that resuming warfarin after 7 days was not associated with a higher risk of recurrent gastrointestinal bleeding and that the rates of death and thromboembolism were lower than in patients who resumed warfarin...
RESUMING ANTICOAGULATION

after 30 days. On the other hand, the risk of recurrent gastrointestinal bleeding was significantly greater if therapy was resumed within the first week.

In view of these studies, we believe that most patients should resume anticoagulation after 4 to 7 days of interruption after gastrointestinal bleeding.55

Timing after soft-tissue hemorrhage

The literature on resuming anticoagulation after soft-tissue hemorrhage is sparse. A retrospective study52 looked at this question in patients with spontaneous rectal sheath hematoma who had been receiving antiplatelet drugs, intravenous heparin, vitamin K antagonists, or a combination of these, but not target-specific agents. More than half of the patients were on vitamin K antagonists at the time of hemorrhage. Analysis suggested that when benefits of resuming anticoagulation are believed to outweigh risks, it is reasonable to resume anticoagulation 4 days after the index event.56

Timing after intracranial hemorrhage

Anticoagulation should not be considered within the first 24 hours after intracranial hemorrhage, as over 70% of patients develop some amount of hematoma expansion during this time.34,57 The period thereafter poses a challenge, as the risk of hematoma expansion decreases while the risk of arterial and venous thromboembolism is ongoing and cumulative.50

Perhaps surprisingly, national guidelines suggest starting prophylactic-dosed anticoagulation early in all intracranial hemorrhage patients, including those not previously on warfarin.58,59 In a randomized trial, Boeer et al60 concluded that starting low-dose subcutaneous heparin the day after an intracranial hemorrhage decreased the risk of thromboembolism without increasing the risk of rebleeding.60 Dickmann et al61 similarly concluded that there was no increased risk of rebleeding with early prophylactic-dosed subcutaneous heparin.61 Optimal mechanical thromboprophylaxis, including graduated compression stockings and intermittent pneumatic compression stockings, is also encouraged.34

Expert opinion remains divided on when and if anticoagulants should be resumed.34,62 The American Heart Association suggests that in nonvalvular atrial fibrillation, long-term anticoagulation should be avoided after spontaneous lobar hemorrhage; antiplatelet agents can be considered instead.39 In nonlobar hemorrhage, the American Heart Association suggests that anticoagulation be considered, depending on strength of indication, 7 to 10 days after the onset.58 The European Stroke Initiative suggests patients with strong indications for anticoagulation be restarted on warfarin 10 to 14 days after the event, depending on the risk of thromboembolism and recurrent intracranial hemorrhage.59 Others suggest delaying resumption to 10 to 30 weeks after an index intracranial hemorrhage.63

Overall, in the immediate acute period of intracranial hemorrhage, most patients will likely benefit from acute reversal of anticoagulation, followed by institution of prophylactic-dose anticoagulation after the first 24 hours. Going forward, patients who remain at higher risk of a recurrence of anticoagulant-related intracranial hemorrhage (such as those with lobar hemorrhage, suspected cerebral amyloid angiopathy, and other high-risk factors) than of thromboembolic events may be best managed without anticoagulants. Alternatively, patients with deep hemispheric intracranial hemorrhage, hypertension that can be well controlled, and a high risk of serious thromboembolism may experience net benefit from restarting anticoagulation.34

We recommend considering restarting anticoagulation 7 days after the onset of intracranial hemorrhage in patients at high risk of thromboembolism and after at least 14 days for patients at lower risk (TABLE 2). Discussions with neurologic and neurosurgical consultants should also inform this timing decision.

WOULD A NEWER DRUG BE A BETTER CHOICE?

The emergence of target-specific oral anticoagulants, including factor Xa inhibitors such as rivaroxaban, apixaban, and edoxaban and the direct thrombin inhibitor dabigatran etexilate, presents further challenges in managing anticoagulation after hemorrhage. TABLE 4 summarizes the current FDA-approved indications.64-67

These newer agents are attractive because, compared with warfarin, they have wider therapeutic windows, faster onset and offset...
### TABLE 4
Uses of oral anticoagulants approved by the US Food and Drug Administration

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage</th>
<th>For postoperative prophylaxis after total knee or hip arthroplasty</th>
<th>To treat and reduce risk of recurrence of deep vein thrombosis and pulmonary embolism</th>
<th>Cost ($) a</th>
</tr>
</thead>
<tbody>
<tr>
<td>Warfarin (vitamin K antagonist)</td>
<td>Dosed to target international normalized ratio (INR)</td>
<td>Not recommended for this use</td>
<td>Dosed to target INR</td>
<td>6.00 (generic) 43.00 (brand name)</td>
</tr>
<tr>
<td>Apixaban (factor Xa inhibitor)</td>
<td>5 mg twice daily</td>
<td>2.5 mg twice daily</td>
<td>Not FDA-approved for this use</td>
<td>265.00</td>
</tr>
<tr>
<td></td>
<td>2.5 mg twice daily in those with at least 2 of the following:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>age &gt; 80, body weight &lt; 60 kg, or serum creatinine &gt; 1.5 mg/dL</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Edoxaban (factor Xa inhibitor)</td>
<td>60 mg once daily if creatinine clearance is &gt; 50 to ≤ 95 mL/min (do not use if higher)</td>
<td>Not recommended for this use</td>
<td>60 mg once daily</td>
<td>333.00</td>
</tr>
<tr>
<td></td>
<td>30 mg once daily if creatinine clearance is 15–50 mL/min</td>
<td></td>
<td>30 mg once daily if creatinine clearance is 15–50 mL/min or if body weight is ≤ 60 kg or if using certain P-glycoprotein inhibitors</td>
<td></td>
</tr>
<tr>
<td>Rivaroxaban (factor Xa inhibitor)</td>
<td>20 mg once daily with evening meal</td>
<td>10 mg once daily with or without food</td>
<td>15 mg twice daily with food for the first 21 days for initial treatment of acute deep vein thrombosis or pulmonary embolism</td>
<td>265.00</td>
</tr>
<tr>
<td></td>
<td>15 mg once daily with evening meal if creatinine clearance is 15–50 mL/min</td>
<td></td>
<td>After the initial treatment period, 20 mg once daily with food for the remaining treatment</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Avoid for this indication if creatinine clearance is &lt; 30 mL/min</td>
<td></td>
</tr>
<tr>
<td>Dabigatran (direct thrombin inhibitor)</td>
<td>150 mg twice daily</td>
<td>Not FDA-approved for this use</td>
<td>150 mg twice daily</td>
<td>265.00</td>
</tr>
<tr>
<td></td>
<td>75 mg twice daily if creatinine clearance is 15–30 mL/min</td>
<td></td>
<td>Recommend initiation after 5–10 days of parenteral anticoagulation</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Avoid for this indication if creatinine clearance is &lt; 30 mL/min</td>
<td></td>
</tr>
</tbody>
</table>

a Approximate wholesale acquisition cost of 30 days of treatment at the lowest daily dose; actual retail prices may be higher. Information from references 64–67 and www.fdbhealth.com.
RESUMING ANTICOAGULATION

of action, and fewer drug and food interactions. A meta-analysis of data available to date suggests that the new drugs, compared with warfarin, show a favorable risk-benefit profile with reductions in stroke, intracranial hemorrhage, and mortality with similar overall major bleeding rates, except for a possible increase in gastrointestinal bleeding.

However, when managing anticoagulation after a bleeding event, the newer agents are challenging for two reasons: they may be associated with a higher incidence of gastrointestinal bleeding than warfarin, and they lack the typical reversal agents that can be used to manage an acute bleeding event.

In individual studies comparing warfarin with dabigatran, rivaroxaban, apixaban, or edoxaban for stroke prevention in patients with atrial fibrillation, there was no significant difference in the rate of major bleeding between dabigatran in its higher dose (150 mg twice a day) or rivaroxaban compared with warfarin. The risk of major bleeding was actually lower with apixaban and edoxaban.

In regard to specific types of major bleeding, the rate of intracranial hemorrhage was significantly lower with dabigatran, rivaroxaban, apixaban, and edoxaban than with warfarin. Some have proposed that since the brain is high in tissue factor, inhibition of tissue factor-factor VIIa complexes by vitamin K antagonists leaves the brain vulnerable to hemorrhage. Others suggest that the targeted mechanism of target-specific agents, as opposed to the multiple pathways in both the intrinsic and extrinsic coagulation cascade that vitamin K antagonists affect, may explain this difference.

However, some studies suggest that rivaroxaban and the higher doses of dabigatran and edoxaban are associated with higher rates of major gastrointestinal bleeding compared with warfarin. But apixaban demonstrated no significant difference in gastrointestinal bleeding, and instead demonstrated rates of gastrointestinal bleeding comparable to that with aspirin for stroke prevention in atrial fibrillation.

The new oral anticoagulants lack antidotes or reversal agents such as phytonadione and fresh-frozen plasma that are available to manage warfarin-associated bleeding events. Other proposed reversal options for the new agents include activated charcoal (if the drugs were taken recently enough to remain in the gastrointestinal tract) and concentrated clotting factor product, though research is ongoing in regards to the most appropriate use in clinical practice. Unlike rivaroxaban and apixaban, dabigatran has low plasma protein binding and is dialyzable, which provides another strategy in managing dabigatran-related bleeding.

Of note, the above bleeding risk calculations relate to the first anticoagulant-related bleeding event, though presumably the same risk comparison across agents may be applicable to rebleeding events. Given the data above, when anticoagulation is to be resumed after an intracranial hemorrhage, the risk of rebleeding, particularly in the form of recurrent intracranial hemorrhage, may be lower if a target-specific oral anticoagulant is used. Similarly, when anticoagulation is to be resumed after a gastrointestinal bleeding event, reinitiation with warfarin or apixaban therapy may present the lowest risk of recurrent gastrointestinal rebleeding. In other sources of bleeding, such as retroperitoneal bleeding, we suggest consideration of transitioning to warfarin, given the availability of reversal agents in the event of recurrent bleeding.

Other important drug-specific factors that must be noted when selecting an agent with which to resume anticoagulation after a hemorrhage include the following:

- In patients with significant renal impairment, the choice of agent will be limited to a vitamin K antagonist.
- A meta-analysis of randomized clinical trials suggests that in the elderly (age 75 and older) target-specific oral anticoagulants did not cause excess bleeding and were associated with at least equal efficacy compared with vitamin K antagonists.
- Target-specific oral anticoagulants may be beneficial in patients who have challenges in achieving INR targets, as evidence suggests that switching to them is associated with a reduction in bleeding for patients who struggle to maintain an appropriately therapeutic INR. On the other hand, if there is concern that a patient may occasionally miss doses of an anticoagulant, given the rapid onset and offset of action of target-specific agents compared with warfarin, a missed dose of a target-
specific agent may result in faster dissolution of anticoagulant effect and increased risk of thrombotic events, and lapses in anticoagulation will not be identified by routine drug monitoring. As such, it is vital to have a frank discussion with any patient who has difficulty maintaining therapeutic INRs on warfarin treatment to make sure that he or she is not missing doses.

- If there is no clear and compelling reason to select a particular agent, cost considerations should be taken into account. We have included estimated 30-day pricing for the various agents in Table 4.

**Table 4.**

<table>
<thead>
<tr>
<th>Agent</th>
<th>Estimated 30-day Pricing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eliquis (apixaban)</td>
<td>$125.00</td>
</tr>
<tr>
<td>Xarelto (rivaroxaban)</td>
<td>$135.00</td>
</tr>
<tr>
<td>Pradaxa (dabigatran)</td>
<td>$150.00</td>
</tr>
</tbody>
</table>

**REFERENCES**

RESUMING ANTICOAGULATION


55. Brotman DJ, Jaffer AK. Resuming anticoagulation in the first week following gastrointestinal tract hemorrhage: should we adopt a 4-day rule? Arch Intern Med 2012; 172:1492–1493.
