Brief Review

Factor XI as a Therapeutic Target

David Gailani, Andras Gruber

Abstract—Factor XIa is a plasma serine protease that contributes to thrombin generation primarily through proteolytic activation of factor IX. Traditionally considered part of the intrinsic pathway of coagulation, several lines of evidence now suggest that factor XIa serves as an interface between the vitamin-K–dependent thrombin generation mechanism and the proinflammatory kallikrein–kinin system, allowing the 2 systems to influence each other. Work with animal models and results from epidemiological surveys of human populations support a role for factor XIa in thromboembolic disease. These data and the clinical observation that deficiency of factor XI, the zymogen of factor XIa, produces a relatively mildly bleeding disorder suggest that drugs targeting factor XI or XIa could produce an antithrombotic effect while leaving hemostasis largely intact. Results of a recent trial comparing antisense-induced factor XI reduction to standard-dose low molecular–weight heparin as prophylaxis for venous thrombosis during knee replacement are encouraging in these regards. Here, we discuss recent findings on the biochemistry, physiology, and pathology of factor XI because they relate to thromboembolic disease. 

Key Words: factor XI ■ factor XII ■ hemorrhage ■ thrombin ■ thrombosis

Vitamin K antagonists such as warfarin have been mainstays of antithrombotic therapy for >50 years. Although these drugs demonstrate efficacy across a spectrum of clinical settings, they increase the risk of major bleeding, and frequent monitoring is required to maintain the drug effect within a narrow therapeutic window.1 In 2005, Hirsh et al2 described features of an ideal anticoagulant that included a high efficacy-to-safety index and a predictable dose response that obviates the need for laboratory monitoring. Direct oral anticoagulants (DOACs) that target thrombin or factor Xa are improvements over warfarin in these regards. They are at least as effective as warfarin for preventing stroke in patients with atrial fibrillation and treating venous thromboembolism (VTE) and are as effective as low molecular–weight heparin for VTE prophylaxis in hip or knee arthroplasty.3 They also seem to be safer than warfarin with lower rates of intracranial bleeding and are given in fixed doses that do not require monitoring. However, the available DOACs, such as warfarin, target proteins that are central to the body’s response to injury, and this places limits on how they can be used.

There is interest in targeting the plasma zymogen factor XI (fXI) and its protease form factor XIa (fXIa) for prevention or treatment of thrombosis. FXI seems to contribute to VTE and ischemic stroke in humans4,5 and is required for formation of occlusive clots in animal thrombosis models.6,7 Because congenital absence of fXI is associated with a relatively mild bleeding disorder,8 it is anticipated that neutralizing fXI or fXIa would cause a smaller defect in hemostasis than would warfarin or a DOAC. Results from a recent clinical trial involving patients undergoing knee replacement support the premise that an antithrombotic effect can be achieved by targeting fXI without precipitating severe bleeding.9 Here, we discuss recent findings on the biochemistry and physiology of fXI, the preclinical and clinical evidence supporting a role for this protein in thrombosis, and the mechanisms by which it may contribute to thromboembolism.

The Factor XI Molecule

FXI arose from a duplication of the gene for prekallikrein, the zymogen of α-kallikrein.10 Prekallikrein, factor XII (fXII), and high molecular–weight kininogen comprise the plasma kallikrein–kinin system (Figure 1).11,12 Among its functions, the kallikrein–kinin system may contribute to the host response to infection by assembling on microorganisms, and generating inflammatory kinins and antimicrobial peptides. A similar process, contact activation, leads to coagulation when blood is exposed to artificial surfaces, such as medical devices used in extracorporeal blood oxygenation.11,12 During contact activation, reciprocal conversion of prekallikrein and fXII to α-kallikrein and factor XIIa (fXIIa) occurs on a surface (Figure 1). FXI, like its homolog prekallikrein, is activated by fXIIa and, similar to α-kallikrein, fXla has some capacity to activate fXII.5,13 However, fXI has features distinguishing it from prekallikrein that facilitate interactions with the thrombin generation mechanism (Figure 1).10 Prekallikrein and fXI polypeptides each have 4 apple domains and a trypsin-like catalytic domain. The fXI apple 3 domain contains a factor IX-binding exosite not present on prekallikrein. The amino acid sequence adjacent to the fXI activation site (Arg89-Ile90) also differs from corresponding prekallikrein...
Figure 1. The relationship of factor XI to thrombin generation and contact activation. Proteolytic reactions required for thrombin generation at an injury site are shown in the gray oval on the left, with each reaction indicated by a yellow arrow. The factor VIIa/tissue factor (TF) complex initiates thrombin generation by activating factors X and IX. Activated factor X (factor Xa) is responsible for cleaving prothrombin to form thrombin. Protease zymogens are indicated in black, and their active forms are indicated by a lower case “a”. Cofactors are shown as red ovals. Calcium (Ca2+) and phospholipid (PL)-dependent reactions are indicated. Thrombin generated early in coagulation converts FXI to FXIa, which sustains thrombin production through factor X activation (green arrows). Note that FXI activation does not require FXIIa, explaining why FXII deficiency does not cause bleeding. Proteolytic reactions involved in contact activation are shown in the gray oval on the right, with each reaction indicated by a black arrow. Artificial or abnormal surfaces facilitate FXII autoactivation. FXIIa converts prekallikrein (PK) to α-kallikrein, which activates additional FXII and cleaves high molecular-weight kininogen (HK), liberating bradykinin (BK), and antimicrobial peptides (AMPs). Contact activation can promote thrombin generation through FXIIa-mediated activation of FXI.11,12 There is evidence that FXIa, in turn, can activate FXII,13 although this is not a standard part of contact activation models. In plasma, PK and FXI circulate as complexes with HK, which may serve as a cofactor for PK and FXI activation. Activation of FXI by FXIIa is not required for hemostasis but contributes to thrombosis in animal models. FXI is considered a component of contact activation (kallikrein–kinin) and thrombin generation in the scheme shown here, functioning as a bidirectional interface between the 2 systems. Hypothetically, activation of either system could activate the other through FXI conversion to FXIIa. Image modified from Gailani et al4 and Bane et al13 with permission.

sequence, permitting FXI to be activated by thrombin, as well as by FXIIa. The combination of prekallikrein-like and novel features permits FXI to promote thrombin generation through FXIIa-dependent and FXIIa-independent processes (Figure 1).4

Factor XI in Hemostasis

The phenotype associated with congenital FXI deficiency indicates that FXI has a role in limiting trauma-induced bleeding. In humans, severe deficiency (≤15% normal level) may exacerbate post-traumatic bleeding, particularly in areas with high fibrinolytic activity (urinary tract, nose, and mouth).7 Hemorrhage in other tissues is less frequent, and procedures such as appendectomy and cholecystectomy may be well tolerated without factor replacement.7,14 Because deficiencies of FXII, prekallikrein, or high molecular–weight kininogen are not associated with abnormal bleeding, FXI is probably activated by FXIIa-independent processes during hemostasis. In the model in Figure 1, FXI is activated by thrombin after the VIIa/tissue factor complex initiates coagulation, with FXIa sustaining thrombin generation through factor IX activation.15 In addition to promoting fibrin formation, FXI-dependent thrombin generation may promote activation of TAFI (thrombin-activatable fibrinolysis inhibitor), a metalloprotease that modifies fibrin by removing binding sites for fibrinolytic proteins, rendering it resistant to fibrinolytic degradation.16 Although severe FXI deficiency delays clot formation in surface-dependent assays such as the activated partial thromboplastin time, the magnitude of the abnormality correlates poorly with symptoms, and some patients with severe deficiency may not bleed abnormally, even with trauma. It is conceivable that some individuals have relatively robust factor VIIa/tissue factor activity or weak fibrinolytic activity that tips the balance in favor of clot stability, rendering FXIa unnecessary. The clinical experience suggests that inhibitors targeting FXI/FXIa would leave some patients more prone to trauma-induced bleeding. However, spontaneous soft tissue bleeding is not part of the phenotype of FXI deficiency, and such drugs would not be expected to precipitate severe bleeding as frequently as would warfarin or DOACs.

Factor XI and Thrombosis in Humans

Despite its modest role in hemostasis, there is substantial evidence supporting a role for FXI in thrombosis. Plasma FXI levels at the upper end of the normal range are linked to modest increases in VTE and ischemic stroke risk.4,5 The 10% of subjects with the highest FXI levels in the Leiden Thrombophilia Study had a 2-fold higher risk of VTE than the 90% of subjects with lowest FXI levels.17 This result is supported by data from the Longitudinal Investigation of Thromboembolism Etiology cohort,18 and the observation that
FXI deficiency reduces the incidence of VTE. High FXI or FXIa levels were associated with increased risk for ischemic stroke in several studies, including the Atherosclerosis Risk in Communities study and the Risk of Arterial Thrombosis In relation to Oral contraceptives study. Severe FXI deficiency reduces incidence of stroke. A role for FXI in myocardial infarction is less clear. FXI levels correlated with myocardial infarction risk in the Study of Myocardial Infarction Leiden but not in the Atherosclerosis Risk in Communities or Risk of Arterial Thrombosis In relation to Oral contraceptives studies, and the incidence of myocardial infarction in FXI-deficient people is similar to the expected incidence in age-matched controls. These data indicate that FXI participates in thrombosis in humans, but suggest that the contribution varies between vascular beds.

**Factor XI and Thrombosis—Animal Models**

Mice lacking coagulation factors have been compared for resistance to thrombosis by a variety of techniques. Our data for ferric chloride (FeCl₃)-induced carotid artery occlusion are summarized in Figure 2. Factor IX-deficient mice have a significant bleeding disorder (hemophilia B), and high FeCl₃ concentrations are required to induce thrombosis in these animals. Mice lacking FXI or FXII do not have obvious hemostatic abnormalities. Despite this, they are at least as resistant to FeCl₃-induced thrombosis as factor IX-deficient mice, showing that a bleeding phenotype is not a prerequisite for resistance to thrombus formation. Mice lacking prekallikrein or high molecular-weight kininogen, proteins required for optimal FXII, are also resistant to FeCl₃-induced thrombosis, suggesting that a contact activation-like process drives thrombosis in mice.

In a primate model, FXI and FXII contributed to thrombosis, but in contrast to mice, FXI inhibition had a greater effect than FXII inhibition. Perhaps this reflects stronger thrombin-mediated FXI activation in primates. When tissue factor–coated or collagen-coated vascular grafts are inserted into arteriovenous shunts in baboons, platelets and fibrin deposit within the coated graft segment, followed by clot extension downstream into the uncoated graft. Polyclonal anti-FXI IgG or a monoclonal IgG that blocks FXIa activation of factor IX (O1A6) reduces platelet and fibrin accumulation within coated graft portions and blocks downstream growth. Monoclonal IgG inhibiting FXI activation by FXIIa (14E11/3G3) or FXII activation (15H8) has little effect on platelet accumulation, and a modest effect on fibrin deposition, in coated portions of grafts (Figure 3) but do limit clot extension. The results support a larger role for FXI than FXII in the primate model, in agreement with data indicating that FXI makes a greater contribution to VTE and stroke than does FXII in humans.

In the mouse and primate models, lack of FXI activity interferes with thrombus growth. Although this could reflect a decreased rate of clot formation, increased clot breakdown may also be a factor. Fibrinolytic degradation of plasma thrombi introduced into the jugular veins of rabbits was ≈2-fold greater in the presence of neutralizing anti-FXI antibody than in controls, consistent with in vitro results demonstrating increased clot sensitivity to fibrinolysis in the absence of FXI. A role for FXI in enhanced clot resistance to fibrinolysis is consistent with the observation that FXI-deficient individuals bleed most frequently from tissues with high intrinsic fibrinolytic activity.

**Factor XI and Inflammation—Animal Models**

Although the contributions of FXI to coagulation have received considerable study, data from mouse models suggest that FXI influences inflammatory processes in a manner that impacts blood vessel biology. Absence of protein C (PC), the zymogen of a protease that regulates coagulation and inflammation, results in perinatal death with intravascular thrombus formation and tissue inflammation prominently featured. FXI deficiency reduces the severity of this phenotype, with some PC⁻/⁻/FXI⁻/⁻ mice living to adulthood. FXI deficiency reduces ischemia–reperfusion injury after temporary middle cerebral artery occlusion. Although reduced fibrin deposition may have a role in the tissue-sparing effect, alteration of the inflammatory response could be involved. Shnerb Ganor et al reported that FXI deficiency reduces atherosclerotic plaque growth and plaque infiltration by macrophages in ApoE-deficient mice. van Montfoort et al also noted reduced macrophage infiltration and an absence of

![Figure 2](http://atvb.ahajournals.org/)

**Figure 2.** Ferric chloride (FeCl₃)-induced carotid artery occlusion. Carotid artery occlusion was induced in wild-type (WT) C57Bl/6 mice, and in mice lacking factor IX (IX⁻/⁻), factor XI (XI⁻/⁻), prekallikrein (PK⁻/⁻), or high molecular-weight kininogen (HK⁻/⁻) with varying concentrations of FeCl₃ as indicated at the bottom of the graph. The percent of animals with patent arteries 30 minutes after FeCl₃ exposure is shown (n=10 for each bar). HK⁻/⁻ mice are homozygous null for deletions of the Kng1 gene. Mice, unlike humans, have 2 kinogen genes (Kng1 and Kng2). Kng1 is thought to be responsible for most, if not all, of the HK in plasma. Image derived from data in references 25–27.
neutrophils in arteries in ApoE-null mice after knockdown of fXI expression. FXI-deficient mice have a survival advantage over wild-type mice after sepsis induced by cecal ligation and puncture. Although initial studies pointed to reduced consumptive coagulation in FXI-deficient animals, we noted that a coagulopathy is not a consistent feature after cecal ligation and puncture and that FXI deficiency improves survival in the absence of a coagulopathy. The early cytokine response and rise in markers of systemic inflammation after cecal ligation and puncture were reduced in FXI-deficient mice. The mechanism by which FXI promotes inflammation is just starting to be investigated. Certainly, a contribution based on its role in thrombin and fibrin formation seems likely. However, in the cecal ligation and puncture model, FXI deficiency also decreased FXII and prekallikrein turnover, suggesting that it contributes to activation of these proteins (Figure 1). If this is the case, FXI deficiency could reduce kallikrein generation and subsequent bradykinin production. In addition, FXI modulates neutrophil migration in vitro, and its absence could alter neutrophil behavior during inflammatory processes.

Therapeutic Targeting of Factor XI and Factor Xla

Several therapeutic strategies to target FXI and FXIa are under development. Antisense oligonucleotide (ASO) knockdown of FXI expression has undergone phase II testing. 2'-Methoxethyl DNA ASOs are avidly taken up by hepatocytes and bind to mRNAs through complementary base pairing, followed by RNase H–dependent degradation of the ASO–mRNA complex. The anti-FXI ASO IONIS-FXIa (formerly ISIS-416858) was compared with standard-dose enoxaparin for prevention of VTE in patients undergoing knee replacement. IONIS-FXIRX was given in 200 or 300 mg ASO. Interestingly, thrombi were not only rarer but also smaller in the 300 mg ASO group, suggesting that thrombus growth is compromised when FXI is reduced below a threshold. Alternatively, because venography was performed more than a week after surgery, the results could reflect greater fibrinolytic degradation of clot formed intraoperatively in the absence of FXI. Although clinically relevant bleeding was not statistically different in ASO- and enoxaparin-treated patients, the study was not powered to show a difference in bleeding. It is worth noting that patients started ASO treatment 5 weeks before surgery and were under the full drug effect during surgery. Despite this, abnormal intraoperative hemostasis was
not observed, and postoperative bleeding was rare, even with FXI levels <10% of normal.

Because of the slow onset of action, ASO therapy cannot be applied in situations where an antithrombotic effect is required rapidly. Anti-FXI antibodies have shown promise in preclinical studies, and an antibody specifically targeting FXIa is entering clinical evaluation. Antibody-based therapies are highly specific and provide rapid onset of action. Their half-lives make them better suited for long-term inhibition than for situations requiring brief treatment. Small molecule active site inhibitors of FXIa are efficacious in rodent and rabbit thrombosis models, and some are entering phase I testing. Most have relatively short half-lives, facilitating dose adjustment and more rapid dissipation of effect after discontinuation. Specificity is an issue with these agents, as the FXIa active site is structurally similar to those of other proteases. Some anti-FXIa compounds demonstrate activity toward the FXI homolog α-kallikrein. This might be beneficial, because it may produce anti-inflammatory and antithrombotic effects distinct from those attributable to FXIa inhibition.

Conclusions, Conjecture, and Future Considerations

FXI was identified as a plasma constituent missing in patients with abnormal surface-dependent clotting and a mild bleeding disorder. As part of the intrinsic pathway, the protein provides a link between contact activation and factor IX, driving thrombin generation in activated partial thromboplastin time assays. Subsequent work identified FXIa-independent mechanisms for FXI activation, explaining the phenotypic differences between FXI and FXII deficiencies. Genomic studies have shed additional light on the relationship between FXI and the traditional coagulation cascade. Most terrestrial vertebrates have a vitamin K-dependent mechanism for thrombin generation that is required for hemostasis, and a kallikrein-kinin system (Figure 1). However, in most lineages, these 2 ancient systems lack the connection provided by FXI. The duplication of the prekallikrein gene that led to FXI seems to have occurred during mammalian evolution. FXI retains features of prekallikrein that allow it to interact with FXII/FXIIa, while possessing adaptations that facilitate interactions with factor IX and thrombin. On the basis of the structural, biochemical, and genomic data, it seems reasonable to classify FXI as a component of contact activation and as a coagulation factor. The dual role may be central to its ability to promote thrombosis, because it allows the kallikrein–kinin system and coagulation mechanisms to activate each other (Figure 1). This strategic location indicates that FXI may be an excellent target for controlling thromboinflammatory processes initiated by either system.

The prominent roles for FXI and FXII in thrombosis models stand in stark contrast to their limited importance for hemostasis, indicating fundamental differences in processes that form hemostatic and thrombotic clots. In mice, FXI or FXII deficiency do not prevent clot formation on surfaces of injured vessels but do cause an impressive defect in clot propagation into the vessel lumen. Platelet deposition in thrombogenic grafts in baboons treated with anti-FXI IgG (Figure 3) is also primarily on the graft surface. Intraluminal thrombus fragment in the absence of FXI or FXII in flowing blood. In this environment, these proteins may support thrombin generation to maintain thrombus stability and growth. Hemostatic clots may not require FXII and in most cases require FXI because they form largely outside of damaged vessels and within vessel walls, away from the higher shear environment in the lumen. Perhaps unrecognized features of thrombotic clots allow them to recruit FXI and FXII more efficiently than do hemostatic clots. We need to identify and characterize factors that promote FXI and FXII activation during thrombosis because they may turn out to be targets for antithrombotic therapy in their own right. Polyanions that facilitate FXI and FXII activation such as polyphosphate, DNA-containing neutrophil extracellular traps, and RNA, are candidates for cofactors that enhance FXI and FXII activation and may contribute to thrombus formation in vivo.

The recent ASO trial provided proof of concept that targeting FXI can produce a useful therapeutic effect in patients at risk of thrombosis. Future work will be directed at determining whether these promising results are relevant to other clinical settings. It seems prudent to first test FXIa inhibitors for prophylaxis, such as in primary or secondary prevention of VTE or prevention of stroke in patients with atrial fibrillation. At this point, we do not have preclinical data to indicate that FXI/FXIa inhibition will be useful in treating active thrombosis. FXIa inhibition may be useful in patients who are poor candidates for warfarin or DOACs, or as short-term prophylaxis after procedures, where even modest bleeding is problematic (eg, neurosurgery). Inhibitors of FXI/FXIa should also be considered in situations where blood is exposed to artificial surfaces (dialysis, extracorporeal membrane oxygenation, cardiopulmonary bypass, and artificial heart valves).

The advantage of strategies targeting FXI compared with current therapies likely lies in the area of safety. Preclinical and epidemiological data indicate that FXI/FXIa inhibitors will not promote bleeding after injury to the same extent as warfarin or DOAC therapy and, therefore, may not need to be discontinued before certain invasive procedures. That patients successfully underwent knee replacement with reduced FXI levels is encouraging but does not establish the safety profile for drugs targeting this protease for all hemostatic challenges. Considering the phenotype of FXI deficiency, we should anticipate that FXI inhibitors will exacerbate bleeding in some individuals undergoing surgery, particularly when the oropharynx or urinary tract is involved, and in some women with a propensity for menorrhagia or who are postpartum. An ideal antithrombotic agent could be defined as a drug that, at its effective dose, completely and specifically blocks a prothrombotic activity while leaving hemostasis intact, eliminating concerns with overdose or monitoring. Along this line of thinking, FXIIa seems to be an ideal target. Preclinical studies in rodents and rabbits support a role for FXII in thrombus formation and selective blockade of FXI activation by FXIIa without inhibiting other functions of FXIIa or FXI demonstrated antithrombotic efficacy in mice and primates. However, enthusiasm must be tempered by observations that FXII does not seem to contribute to VTE, stroke, or myocardial infarction in...
humans and that FXII inhibition is less effective than FXI inhibition in the primate model.\textsuperscript{29,30} Also, FXII likely contributes to several homeostatic and host-defense processes,\textsuperscript{11,12} and the repercussions of long-term inhibition of FXII are uncertain. Still, brief treatment with FXII inhibitors may be effective for procedures in which blood is exposed to artificial surfaces.\textsuperscript{11} If inhibitors of FXI/FXIa and FXII/FXIIa prove to be effective in the clinic, their safety profiles should broaden the spectrum of clinical situations in which antithrombotic therapy can be applied.

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**References**


The plasma protease factor Xla contributes to thrombin generation by activating factor IX. Traditionally considered part of the intrinsic pathway of coagulation, factor Xla may function as an interface between thrombin generation and the proinflammatory kallikrein–kinin system.

Data from animal models and epidemiological surveys indicate that factor XI contributes to thromboembolic disease.

Therapeutic strategies designed to neutralize factor Xla (antibodies, RNA aptamers, and small molecule active site inhibitors) or its zymogen form factor XI (antisense oligonucleotides and antibodies) are being developed, with the goal of testing them in prevention or treatment of thromboembolic disease.

Congenital factor XI deficiency is associated with a mild-to-moderate bleeding disorder, and it is anticipated that therapies targeting this protein will be associated with a relatively low risk of serious bleeding compared with currently available anticoagulants.

Results of a recent trial of antisense oligonucleotide-mediated factor XI reduction for venous thrombosis prophylaxis in knee replacement surgery indicate that factor XI is an important contributor to thrombosis in this setting and that targeting factor XI can produce a useful therapeutic effect.
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Graphic Abstract

**Thrombin Generation**
- IX → IXa
- IXa → VIIIa
- VIIIa → Xa
- Xa → Va
- Va → Ca²⁺
- Ca²⁺ → PL
- PL → Thrombin

**Contact Activation**
- (Kallikrein-Kinin System)
- XII → XIl
- XIl → α-Kal
- α-Kal → HK
- HK → PK

Additionally:
- Prothrombin → X
- Xa → Ca²⁺
- Ca²⁺ → PL
- PL → Thrombin

**Recruitment Pathways**
- TF → VIIa
- VIIa → Xa
- Xa → Ca²⁺