Factor XI Inhibitors for Prevention and Treatment of Venous Thromboembolism: A Review on the Rationale and Update on Current Evidence

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Although anticoagulation therapy has evolved from non-specific drugs (i.e., heparins and vitamin K antagonists) to agents that directly target specific coagulation factors (i.e., direct oral anticoagulants, argatroban, fondaparinux), thrombosis remains a leading cause of death worldwide. Direct oral anticoagulants (i.e., factor IIa- and factor Xa-inhibitors) now dominate clinical practice because of their favorable pharmacological profile and ease of use, particularly in venous thromboembolism (VTE) treatment and stroke prevention in atrial fibrillation. However, despite having a better safety profile than vitamin K antagonists, their bleeding risk is not insignificant. This is true for all currently available anticoagulants, and a high bleeding risk is considered a contraindication to anticoagulation. As a result, ongoing research focuses on developing future anticoagulants with an improved safety profile. Several promising approaches to reduce the bleeding risk involve targeting the intrinsic (or contact activation) pathway of coagulation, with the ultimate goal of preventing thrombosis without impairing hemostasis. Based on epidemiological data on hereditary factor deficiencies and preclinical studies factor XI (FXI) emerged as the most promising candidate target. In this review, we highlight unmet clinical needs of anticoagulation therapy, outlay the rationale and evidence for inhibiting FXI, discuss FXI inhibitors in current clinical trials, conduct an exploratory meta-analysis on their efficacy and safety, and provide an outlook on the potential clinical application of these novel anticoagulants.

Keywords: factor XI, anticoagulants, thrombosis, venous thromboembolism, hemorrhage, hemostasis

INTRODUCTION

Evolution of Anticoagulation

Anticoagulation represents one of three forms of antithrombotic therapy, next to antiplatelet and thrombolytic therapy, and is the mainstay of prevention and treatment of venous thromboembolism (VTE), a disease entity covering pulmonary embolism and deep vein thrombosis. The landscape of anticoagulant drugs in use have shifted over the last decades from unspecific drugs with multiple pharmacological targets to agents specifically targeting a factor of
the coagulation cascade. Long before the pharmacological mechanisms had been revealed, heparin and vitamin K-antagonists (VKA) were the first drugs in clinical use (1, 2). However, their narrow therapeutic window, unpredictable pharmacodynamic response, and serious, albeit rare, adverse events (e.g., heparin-induced thrombocytopenia, coumarin-induced skin necrosis) led to the search for safer anticoagulants.

The first evolutionary step in anticoagulant treatment involved the development of derivatives of heparin and coumarin. Low molecular weight heparin (LMWH) and coumarin derivatives such as warfarin and phenprocoumon were developed. Yet, these newer drugs still rely on a multi-targeted approach that impairs physiological hemostasis. Heparin is mediated by antithrombin and heparin cofactor II (3), and coumarins disrupt the synthesis of vitamin-K-dependent coagulation factors (4). The first selective inhibitor of a coagulation factor was fondaparinux, a synthetic heparin pentasaccharide inhibiting factor Xa (FXa) with a favorable pharmacokinetic profile, which, similar to LMWH, did not require coagulation monitoring (5). Development of anticoagulant therapy continued with the invention of selective inhibitors of thrombin (FIIa): bivalirudin and the small molecule argatroban. However, long-term application was still limited due to their need of parenteral administration. But the new era was finally ushered in with the invention of the direct oral anticoagulants (DOAC). Already in 2004, ximelagatran, a direct oral FIIa inhibitor, was developed but withdrawn from the market due to hepatotoxicity. Subsequently, the new class of DOAC (dabigatran, rivaroxaban, apixaban, edoxaban, and betrixaban), which selectively and reversibly inhibit FIIa or FXa in the common pathway of the coagulation cascade, was introduced in clinical practice in 2010, 2011, 2012, 2015, and 2017, respectively, and led to a revolution in anticoagulant therapy (6).

**The Current State of Anticoagulant Therapy**

Predictable pharmacokinetics, ease of use, and a wide therapeutic window are the most important advantages of DOAC, leading to a quick uptake in prescriptions and broad use in clinical practice (6, 7). Today, DOAC are the most commonly used anticoagulants in the western world (8, 9), and two DOAC rank among the top 10 best-selling drugs worldwide (10). In a review on anticoagulants prescription trends in the primary care setting in England, the proportion of DOAC have increased from 9% in 2014 to 74% in 2019 (11). However, an analysis of electronic database searches suggests that VKA still remain the most popular anticoagulants worldwide (12). Possible reasons include limited access to DOAC in some parts of the world, considerably lower costs of VKA, which is particularly relevant in health systems without drug reimbursement, and physician and patient resistance to changing the standard of care. DOAC are most frequently prescribed for stroke prevention in patients with atrial fibrillation, followed by treatment of VTE and prevention of its recurrence (8). In VTE patients, DOAC are now recommended as the first choice for anticoagulant therapy (13–15). Compared to VKA, they have proven to be of similar efficacy (16–21) and are associated with a lower bleeding risk, particularly with respect to intracranial hemorrhage (22). However, higher rates of gastrointestinal bleeding have been reported for dabigatran, rivaroxaban, and edoxaban in special patient populations including patients with atrial fibrillation or cancer (23, 24). To mitigate bleeding risk, reduced-dose DOAC regimens are used in patients with specific dose adjustment criteria and for long-term prevention of VTE (i.e., for rivaroxaban and apixaban) (25). Moreover, specific reversal agents have been developed, and in 2015 idarizumab for dabigatran and in 2018 andexanet alfa for reversal of the FXa inhibitors apixaban and rivaroxaban have been approved and introduced for use in clinical practice (26, 27). Nevertheless, bleeding remains the most frequent complication of anticoagulation. Fear of bleeding contributes to underuse and underdosing of eligible patients, and a high risk of bleeding is the most important contraindication to long-term anticoagulation (28, 29).

**Unmet Clinical Needs**

Patients in need of a better anticoagulation strategy can be divided into two groups. The first group represents patients with an increased risk of bleeding (e.g., patients with kidney impairment, prior history of bleedings, cancer, or the elderly). Notably, bleeding risk assessment is challenging and no single approach has been proved to be consistently superior (30). Thus, the decision to initiate or continue anticoagulation remains at the discretion of the treating physician based on an individual assessment of risk of thrombosis and bleeding. Accordingly, anticoagulants with an improved safety profile could serve as a novel and important treatment option in patients in whom the risk of bleeding is higher than the risk of thrombosis. A second group that might benefit from new treatment options includes patients in whom the efficacy of DOAC has been tested but has shown to be inferior to VKA, or in whom there is insufficient evidence to support the use of DOAC, such as patients whose cardiovascular system is exposed to artificial surfaces. In patients with mechanical heart valves, dabigatran was inferior to VKA (31), and randomized trials of other DOAC against VKA are lacking. In dialysis patients, where the risk of bleeding is exceptionally high (32), the benefit-to-risk ratio of oral anticoagulants, in general, is still debated as observational studies report excess bleeding rates with VKA and DOAC (33, 34). In patients with left ventricular assist devices or extracorporeal membrane oxygenation, DOAC have not been assessed in randomized-controlled trials. Further, VKA are recommended over DOAC in patients with antiphospholipid syndrome as two randomized trials were prematurely terminated due to excess thromboembolic events in the DOAC compared to the VKA treatment arm (35, 36). Taken together, a significant proportion of patients need safer anticoagulants or anticoagulants designed to prevent and treat thrombosis triggered by artificial surfaces.

**Thrombogenesis—Revisiting the Coagulation Cascade**

The formation of a blood clot, i.e., thrombogenesis, is the combined result of increased coagulation, hemodynamic...
changes in the circulation, and vascular endothelial injury (37). Anticoagulation is effective to prevent and treat thrombosis but also interferes with hemostasis, and leads to increased bleeding risk (38).

Different models to explain the coagulation process in vitro and in vivo have been proposed. The classical clotting cascade model consists of the intrinsic and extrinsic pathways that converge in the common pathway (FX, FV, FII) (Figure 1) (39, 40). The extrinsic coagulation pathway [tissue factor (TF), FVII], which is essential for fibrin formation at a site of vascular injury, is initiated by exposure of TF to blood and formation of the TF:FVIIa complex (41). The intrinsic coagulation pathway (FXII, FIX, FVIII) is initiated by contact activation mediated by FXII when blood interacts with certain negatively charged surfaces. This strict separation of both pathways is based on in vitro observations. In the last two decades, the cell-based coagulation model, which interconnects both pathways, gained acceptance and better reflects the hemostatic processes in vivo (42–44). Based on this theory, coagulation or pathological thrombosis is initiated by the exposure of TF from TF-bearing cells (representing the extrinsic pathway). Importantly, large amounts of thrombin are needed thereafter for both thrombosis and hemostasis, which are generated by subsequent activation of the intrinsic pathway (44). In vivo, activation of the intrinsic pathway through the contact system is initiated by FXII activation via exposure to polyanionic compounds originating from injured cells, activated platelets, or pathogens (45). However, in this cell-based model, activation of the intrinsic pathway also occurs independently of the contact system. FXI is activated by FIXa generated from the extrinsic pathway and FIX by the TF:FVIIa complex (extrinsic FXase) (46). Subsequently, FIXa and FXIa contribute to the amplification of thrombin production via the “truncated” intrinsic pathway.

Modern coagulation theory hypothesizes that the mechanisms of pathological thrombosis and physiological hemostasis are distinguishable, but all currently available anticoagulants in clinical use either target coagulation FII or FX, both essential factors for physiological hemostasis. Inhibition of either one of the two factors interrupts the common pathway and disrupts the positive feedback activation of FV, FVIII, and FXI. Upstream the common pathway, initiation of hemostasis mainly depends on the TF-mediated extrinsic pathway, and deficiency or increased activity of this pathway leads to bleeding or thrombosis. The intrinsic pathway, which is activated by the contact activation system and resembles an interface between inflammation, coagulation, and innate immunity, however, contributes weakly to hemostasis but is closely associated with pathological thrombosis (47–50). In the presence of excessive TF, thrombin
generation is likely sufficient for hemostasis without a necessary involvement of the intrinsic pathway. But in the presence of limited TF, activation of FX through the intrinsic FXase is required for rapid and sufficient production of thrombin as seen in various pathological processes (e.g., immunothrombosis, atherothrombosis) (51, 52). Thus, revisiting the intrinsic pathway with its contact activation system in the effort to inhibit the coagulation cascade upstream of FX is a noteworthy approach in the desire to prevent thrombosis without the cost of impairing hemostatic functions.

The Intrinsic Pathway, Contact System, and Rationale for Targeting Factor XI

Anticoagulant-related bleeding might be avoided if coagulation factors that are more important for pathological thrombosis than for physiological hemostasis are targeted selectively (53). Accordingly, the detrimental role of contact system-dependent coagulation activation in thrombus formation proposes the intrinsic coagulation pathway as a potential target for novel anticoagulants. The intrinsic pathway begins with activation through the contact system, which consists of FXII, prekallikrein (PK), and a cofactor high-molecular-weight kininogen (HK) (54). This contact activation is triggered when plasma encounters certain negatively charged macromolecules, leading to the activation of FXII. Subsequently, FXI and FIX get activated. Therefore, factors of the intrinsic pathway represent promising candidate targets for antithrombotic therapy (55–57). Notably, FXI research has proceeded furthest in clinical trials having shown the most promising evidence from epidemiological studies and animal models.

FXI, discovered in 1953, is a serine protease that originated from duplication of the prekallikrein gene. It can be activated by FXIIa through the contact activation pathway but also acts independently of contact activation when activated by thrombin (FIIa) via the positive feedback loop (Figure 1). FXI is considered essential for thrombus growth and stabilization but might have a subsidiary role in physiological hemostasis (58, 59). In epidemiological and Mendelian randomization studies, elevated FXI has been linked to stroke and VTE (60–62). Furthermore, studies on lower levels of FXI or FXI-deficiency showed that decreased FXI levels were associated with a lower risk of VTE and cardiovascular events (63–66). However, patients with FXI-deficiency also appear to be at higher risk for post-traumatic bleeding, whereas patients deficient in contact factors (FXII, PK, HK) have no apparent bleeding diathesis (67). Such findings could be viewed as disproof to the theory of separating thrombosis from hemostasis by FXI inhibition. However, FXI activity levels only poorly correlate with bleeding risk (67), and congenital FXI deficiency is rarely associated with spontaneous bleeding (68). It has been reported that bleeding most likely occurs in individuals with FXI levels lower than 20% (69, 70). Thus, a threshold of 20% of FXI activity levels is typically used to define severe FXI deficiency (66, 71, 72) and moderate inhibition of FXI at or above this threshold might pose a solution to the coagulation dilemma. This hypothesis is further supported by findings from large genome-wide association studies. Genetic disposition to lower FXI levels was associated with a reduced risk for VTE and ischemic stroke but without increased risk for major bleeding (73). Evidence to inhibit FXI is further supported by animal studies. In animal models of thrombosis, FXI-deficient mice were protected from occlusive thrombosis in the vena cava (74), as well as from thrombosis in the arterial system (75). Interestingly, in an animal model of tail or surgical bleeding, mice with FXI deficiency displayed a normal bleeding phenotype, whereas a saphenous vein injury model revealed reduced hemostasis (76). Furthermore, inhibition of FXI in rats was effective in preventing venous thrombosis without increased bleeding in one study and minimal bleeding risk in another (77, 78). Various other studies have also shown similar findings for FXII (79, 80), but preclinical studies on inhibition of FXI and FXII in non-human primates suggested a greater antithrombotic effect when targeting FXI (81–84), providing the basis for FXI as the primary target for a new generation of anticoagulants in clinical trials.

Clinical Trials of Factor XI Inhibition

Early clinical evaluation of novel anticoagulants usually starts with dose-finding studies in patients undergoing elective knee arthroplasty. This is considered a time-efficient way due to the high number of VTE events after surgery and the objective and efficient evaluation of drug efficacy using venography to determine the incidence of deep vein thrombosis of the lower extremity. Four of these studies on FXI inhibition have already been completed, in which clinically relevant bleeding was determined as the primary safety endpoint by combining major bleeding and clinically relevant non-major bleeding (85–88). The second type of FXI inhibitor studies focused on patients with end-stage kidney disease on hemodialysis (89, 90). High risk of bleeding and thrombosis provoked by the extracorporeal device makes them the prime target for potentially safer anticoagulants inhibiting the intrinsic pathway. In the following, all FXI inhibitors that have already completed phase II in clinical trial evaluation are discussed in more detail ordered by appearance of publication (Table 1). In Figure 1, FXI inhibitors and their mechanisms of action are displayed.

IONIS-FXI<sub>RX</sub>/FXI-ASO, an antisense oligonucleotide, was the first agent investigated in a phase II trial in patients undergoing total knee arthroplasty. Antisense oligonucleotides are attractive novel therapeutics that are highly bound to plasma proteins limiting glomerular filtration and urinary excretion, have no known drug-to-drug interaction and require subcutaneous application. IONIS-FXI<sub>RX</sub> blocks the hepatic synthesis of FXI by specifically reducing FXI messenger RNA levels in the liver. As such, it takes several weeks to lower FXI to therapeutic levels, and thus the restoration to normal levels is also delayed for several weeks after treatment discontinuation (81). In a randomized controlled trial of 300 patients, efficacy and safety of IONIS-FXI<sub>RX</sub> at doses of 100, 200, and 300 mg were evaluated and compared to enoxaparin 40 mg once daily (85). Treatment with IONIS-FXI<sub>RX</sub> was initiated 36 days before surgery and patients received 9 subcutaneous doses over a total of 39 days. Shortly after the start of the trial, the 100 mg dose regimen was discontinued due to insufficient reduction of FXI
lower rates in IONIS-FXI treatment group. Bleeding rates did not statistically differ with 200 mg arm, 4% in the 300 mg arm, and 30% in the enoxaparin as assessed by venography. The rates of VTE were 27% in the 200 mg dose regimen proved non-inferior and the 300 mg dose levels and replaced by the 300 mg dose regimen. In this study, the 200 mg dose regimen proved non-inferior and the 300 mg dose regimen superior compared to enoxaparin in preventing VTE. In all knee arthroplasty studies, VTE was the primary endpoint. In a randomized fashion. The main goal was to investigate pharmacokinetics, -dynamics, and adverse events. Notably, no pharmacokinetic differences were observed between injections pre and post hemodialysis. Further, no drug accumulation and no drug-related serious adverse events were noted. Importantly, the study was not powered to estimate efficacy and safety, but predefined comparisons of the clot burden of the hemodialysis circuit suggested a substantial reduction (up to -29%) in severe clotting events after treatment onset of IONIS-FXI. Clinically relevant bleeding occurred in one patient in the placebo and in one patient in the IONIS-FXI group, but minor bleeding was numerically more frequent in patients with the study drug (23 vs. 8%). Another subsequent study in end-stage kidney disease patients has already been completed but not yet published (NCT03358030). Further studies with the ligand-conjugated version of IONIS-FXI, termed fesomersen or IONIS-FXI-L are ongoing (NCT04534114). Osocimab, is a long-acting, fully human monoclonal antibody with a time to maximum plasma concentration of 1–4 h and a half-life of 30–44 days after intravenous application. It binds adjacent to the active site of FXIa and prevents it from activating FIX. In this first trial on FXIa inhibition, efficacy and safety of 4 different doses (0.3, 0.6, 1.2, 1.8 mg/kg) administered pre- or postoperatively via a single, 60-min, intravenous infusion were compared to 40 mg enoxaparin once daily or 2.5 mg apixaban twice daily in 813 patients who underwent total knee replacement (86). Initially, three dose levels of osocimab administered postoperatively were evaluated, but during the ongoing study and without any safety concerns being raised, a higher postoperative dose (1.8 mg/kg) and two preoperative doses (0.3, 1.8 mg/kg) were initiated as prespecified. In the per-protocol analysis, which consisted of only 74% of randomized patients due to missing data from venography, non-inferiority to enoxaparin was shown for the 3 higher doses of osocimab (0.6, 1.2, 1.8 mg/kg). Superiority was shown for the highest dose given preoperatively, while non-inferiority could not be attributed to 0.3 mg/kg osocimab. VTE rates in the treatment groups ranged from 23.7 to 15.7% with postoperative osocimab and 29.9–11.3% with preoperative osocimab compared to 26.3% in the enoxaparin group and 14.5% in the apixaban group. Only one major bleeding event occurred in the study in a patient with a preoperative dose of 1.8 mg/kg. In summary, bleeding rates

| Drug | Type | Mechanism | Administration route | Studies (NCT) | Population (N) | Comparator | Status |
|------|------|-----------|----------------------|--------------|---------------|------------|--------|
| IONIS-FXI | Antisense oligonucleotide of FXI | Inhibits FXI messenger RNA | Subcutaneous (weekly) | NCT01713361, NCT02553889, NCT03558030 | ESKD (300), ESKD (49), ESKD (200) | Placebo, Placebo, Placebo | Published, Published, Published |
| Osocimab | Monoclonal antibody to FXI | Binds and inhibits FXIa | Intravenous, subcutaneous (monthly) | NCT03276143, NCT04523220 | ESKD (813), ESKD (886) | Placebo | Published, Ongoing |
| Abelacimab | Monoclonal antibody to FXI/FXI | Binds and inhibits FXI and FXIa | Subcutaneous (monthly) | EudrACT 2019-003756-37, NCT04753283, NCT05171049, NCT05171075 | TKA (412), AF (1,200), CAT (1,655), CAT (1,020) | Placebo, Rivaroxaban, Apixaban, Dalteparin | Published, Published, Published, Published |
| Milvexian | Small molecule inhibitor of FXI | Binds and inhibits FXI | Oral (daily) | NCT03891524, NCT03766581 | TKA (1,242), Stroke (2,366) | Placebo | Published, Published |
| Xisomab 3G3 | Monoclonal antibody to FXI | Binds FXI and blocks activation by FXIa | Intravenous (single dose) | NCT03612856, NCT04465780 | ESKD (24), CRT (50) | Placebo | None, Published |
| Fesomersen | Antisense oligonucleotide of FXI | Inhibits FXI messenger RNA | Subcutaneous (weekly) | NCT04534114 | ESKD (305) | Placebo | Ongoing |
| Asundexian | Small molecule inhibitor of FXI | Binds and inhibits FXI | Oral (daily) | NCT04218266, NCT04304508, NCT04304534 | AF (753), AMI (1,592), Stroke (1,780) | Placebo | Published, Published, Published |

AF, atrial fibrillation; CAT, cancer-associated thrombosis; CRT, catheter-related thrombosis in cancer patients; ESKD, end-stage kidney disease; TKA, total knee arthroplasty.

TABLE 1 | Overview of factor XI inhibitors in clinical trials.
in the osocimab treatment groups were lower compared to the control group and ranged between 1% with 0.3 mg/kg and 4.7% with 1.8 mg/kg osocimab compared to 5.9% in the enoxaparin arm. In the apixaban group, only 2% of patients experienced clinically relevant bleeding. Serious adverse events were similarly distributed across groups, but higher rates of hypersensitivity and infusion-related reactions were reported for osocimab. However, none resulted in discontinuation of the study drug. A study with anticipated 600 patients with end-stage kidney disease on hemodialysis is under way (NCT04523220).

**Abelacimab** is a monoclonal antibody that binds to the catalytic domain of FXI and locks it in itszymogen (inactive precursor) conformation, thereby preventing its activation by FXIIa or thrombin (FIIa) (91). Three different dosing regimens of abelacimab (30, 75, 150 mg) were tested and randomized against 40 mg of enoxaparin in patients undergoing knee replacement (87). Given the immediate method of action, the monoclonal antibody was administered in a single, intravenous infusion 4–8 h after surgery. After 412 from the anticipated 600 patients were included, the trial was stopped early. This decision was driven by slowed recruitment because of the Covid-19 pandemic. Nevertheless, power was sufficient to show non-inferiority to enoxaparin for all three regimens of abelacimab, while the 75 and 150 mg doses even met criteria for superiority. Rates for the primary efficacy endpoint VTE were 13, 5, 4, and 22% for 30, 75, 150 mg of abelacimab, and 40 mg of enoxaparin, respectively. No clinically relevant bleeding occurred in the enoxaparin arm and the abelacimab treatment group with the highest dose. In the two remaining groups, 2% experienced clinically relevant non-major bleeding. In total, 5 serious adverse events were reported in the abelacimab groups including wound infections, ileus, and ovarian cyst torsion. A randomized study with rivaroxaban as comparator evaluating safety but not efficacy of abelacimab is currently ongoing in 1,200 patients with atrial fibrillation (NCT04755283), and two phase III trials on cancer-associated thrombosis with the aim to enroll 1,655 and 1,020 patients, respectively, have recently been initiated (NCT03766581).

**Xisomab 3G3/AB023** is a recombinant antibody that binds to FXI and selectively reduces its activation by FXIIa without attenuating the feedback mechanism by thrombin (FIIa) on FXI. Due to this unique mechanism, it rather acts as a FXIIa inhibitor and might be categorized as such. Interestingly, the elimination half-life of xisomab 3G3 increases with dose. At a dose of 0.1 mg/kg, a half-life of 1.3 h was measured; at administration of 5 mg/kg, the half-life increased to 121 h (92). This phenomenon of a short half-life at lower doses is likely due to the rapid binding of a large proportion of free xisomab 3G3 to FXI (93). Xisomab 3G3 was tested in a randomized controlled trial of 24 patients with end-stage kidney disease in need of heparin-free hemodialysis (90). Patients were randomized to 0.25, 0.5 mg/kg xisomab 3G3, or placebo injected as a single bolus into the dialysis line proximal to the dialyzer at the start of hemodialysis. The study’s primary aim was to assure safety with regard to bleeding events in this high-risk population. There were no clinically relevant bleeding events while patients were on the study drug, and the time to hemostasis at vascular access sites (i.e., the duration of pressure applied to the access site until bleeding stopped) was unchanged after xisomab 3G3 administration compared to before. One major bleeding event occurred 32 days after drug discontinuation but was deemed unrelated to the study drug by the study investigator. Results on efficacy were underpowered and exploratory but suggested a reduction of occlusive events requiring circuit exchange and lower levels of C-reactive protein and thrombin-antithrombin complexes in patients receiving xisomab 3G3. A subsequent study for the prevention of catheter-associated thrombosis in patients with cancer (NCT04465760) is currently ongoing to further investigate a proof of concept in patients exposed to artificial surfaces.

**Asundexian** is a small molecule direct inhibitor of FXIa with oral bioavailability and a half-life of approximately 12 h, is a selective, reversible, direct inhibitor of FXIa. It has been studied in the largest phase II study so far, which included 1,242 patients undergoing knee arthroplasty to compare 7 different postoperative regimens of milvexian to enoxaparin for the efficacy of preventing VTE and the safety with regard to bleeding events and serious adverse events (88). Both predefined proof of efficacy criteria (i.e., a significant dose-response relationship trend and a VTE incidence that was significantly lower than 30% with the pooled twice-daily milvexian regimens) were met. Furthermore, 4 of the 7 dose regimens (50, 100, 200 mg twice daily, and 200 mg once daily) showed superiority compared to 40 mg of enoxaparin. Serious adverse events were rare and equally distributed across all treatment groups and no major bleeding event occurred in the milvexian treatment groups. Clinically relevant non-major bleeding was below or at 1% with no dose-response relationship seen across the seven different regimens. Similar to the previous studies mentioned, milvexian increased the aPTT ratio in a dose-dependent manner, while enoxaparin did not. A second study in the patients with ischemic stroke is still ongoing (NCT03766581).

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**Asundexian** is a small molecule direct inhibitor of FXIa with oral bioavailability and a half-life of approximately 17 h. Asundexian is renally eliminated to less than 15% and has no clinically relevant interaction with cytochrome P450 (CYP) 3A4 (94). The safety of 2 different dose regimens (20 and 50 mg once daily) was studied in a randomized, double-blind, phase II study in 753 patients with atrial fibrillation and moderate-to-high risk for stroke and bleeding (95). Apixaban, the DOAC believed to have the lowest bleeding risk, was used as a control. Patients were treated for a period of 12 weeks and followed up for safety outcomes. The primary endpoint was the composite of major bleeding and clinically relevant non-major bleeding as defined by the International Society on Thrombosis and Hemostasis (ISTH). After 4 weeks of treatment, asundexian 20 and 50 mg at trough levels resulted in an 81 and 92% reduction in FXIa activity levels, respectively. In total, 10 clinically relevant non-major bleeding and no major bleeding occurred during the study period. The bleeding rates were 1.2, 0.4, and 2.4% in the asundexian 20 mg, asundexian 50 mg, and apixaban treatment groups, respectively, which showed a significant reduction in bleeding rates at a prespecified 90% confidence interval for the 50 mg regimen and in pooled analysis of both asundexian treatment groups. The rates of any adverse event and the rates of adverse events
leading to drug discontinuation were similar across treatment groups, but 2 and 1 patients experienced ischemic stroke in the asundexian 20 and 50 mg arm, respectively, while none suffered from stroke in the apixaban group. In summary, asundexian was safer than apixaban with regard to bleeding events, but efficacy needs to be determined in phase III trials. A second phase II trial in 1,592 patients with recent myocardial infarction has already been completed but not published yet (NCT04304534) and one more study in patients with recent non-cardioembolic stroke is currently ongoing (NCT04304508).

Meta-Analysis of Phase II Studies With Factor XI Inhibitors
To explore the efficacy and safety of FXI inhibitors in completed clinical trials, we performed an exploratory meta-analysis of all phase II studies investigating inhibitors of FXI in patients undergoing total knee arthroplasty. The specified endpoints were confirmed VTE for the efficacy analysis and clinically relevant bleeding (the composite of major bleeding and clinically relevant non-major bleeding) for the safety analysis. Figure 2 displays forest plots summarizing the occurrence of VTE and clinically relevant bleeding events in the 4 phase II studies, respectively, with treatment arms of each study pooled and compared against the enoxaparin comparator group. In a random-effects meta-analysis, treatment with a FXI inhibitor, when compared to 40 mg of enoxaparin, was associated with a decreased risk of VTE by a risk ratio of 0.59 (95% CI, 0.37–0.94; \( p = 0.038 \)). Strikingly, the number of bleeding events in patients treated with FXI inhibitors was also reduced: the risk ratio to develop clinically relevant bleeding in patients treated with a FXI inhibitor compared to enoxaparin was 0.41 (95% CI, 0.19–0.92; \( p = 0.039 \)).

CONCLUSION AND POTENTIAL INDICATIONS
Patients with an elevated risk of bleeding and patients whose blood is exposed to artificial surfaces require novel therapeutic concepts to attenuate the risk of thrombosis without increasing the risk of bleeding. Currently, all anticoagulant strategies target factors of the common pathway of the coagulation cascade, while a more upstream inhibition could be a promising treatment approach in the desire to prevent thrombosis without impairing hemostasis. It is of note that hemostasis strongly depends on the extrinsic pathway, whereas the intrinsic pathway seems less relevant for physiological processes but contributes to thrombosis. As a result, research is currently focused on inhibiting the intrinsic pathway, with FXI emerging as the most promising candidate. Six different inhibitors of FXI successfully completed phase II trials in patients undergoing total knee arthroplasty, patients suffering from end-stage kidney disease, and patients with AF. A new term summarizing these new anticoagulants targeting the intrinsic pathway has not been defined yet. Taken together, all inhibitors under investigation in knee arthroplasty studies proved to be at least non-inferior to LMWH in preventing VTE and highlighted the role of FXI in the pathogenesis of post-surgical thrombosis. This was somewhat surprising as knee arthroplasty is considered a prime example of extrinsic coagulation cascade activation due to a strong TF trigger. Pooled study results suggested a 41% reduction of VTE events with FXI inhibitors compared to LMWH (Figure 2). Of note, the meta-analysis on the safety outcome (i.e., clinically relevant bleeding) also suggested a 59% reduction of risk of bleeding in patients treated with a FXI inhibitor compared...
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