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The Effects of the Contact Activation System on Hemorrhage

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The contact activation system (CAS) exerts effects on coagulation via multiple mechanisms, which modulate both the intrinsic and extrinsic coagulation cascades as well as fibrinolysis and platelet activation. While the effects of the CAS on blood coagulation measured as activated partial thromboplastin time shortening are well documented, genetic mutations that result in deficiencies in the expression of either plasma prekallikrein (PPK) or factor XII (FXII) are not associated with spontaneous bleeding or increased bleeding risk during surgery. Deficiencies in these proteins are often undiagnosed for decades and detected later in life during routine coagulation assays without an apparent clinical phenotype. Increased interest in the CAS as a potentially safe target for antithrombotic therapies has emerged, in large part, from studies on animal models with provoked thrombosis, which have shown that deficiencies in PPK or FXII can reduce thrombus formation without increasing bleeding. Gene targeting and pharmacological studies in healthy animals have confirmed that PPK and FXII blockade does not cause coagulopathies. These findings support the conclusion that CAS is not required for hemostasis. However, while deficiencies in FXII and PPK do not significantly affect bleeding associated with peripheral wounds, recent reports have demonstrated that these proteins can promote hemorrhage in the retina and brain. Intravitreal injection of plasma kallikrein (PKal) induces retinal hemorrhage and intracerebral injection of PKal increases intracranial bleeding. PPK deficiency and PKal inhibition ameliorates hematoma formation following cerebrovascular injury in diabetic animals. Moreover, both PPK and FXII deficiency are protective against intracerebral hemorrhage caused by tissue plasminogen activator-mediated thrombolytic therapy in mice with thrombotic middle cerebral artery occlusion. Thus, while the CAS is not required for hemostasis, its inhibition may provide an opportunity to reduce hemorrhage in the retina and brain. Characterization of the mechanisms and potential clinical implications associated with the effects of the CAS on hemorrhage requires further consideration of the effects of PPK and FXII on hemorrhage beyond their putative effects on coagulation cascades. Here, we review the experimental and clinical evidence on the effects of the CAS on bleeding and hemostatic mechanisms.

Keywords: kallikrein–kinin system, hemorrhage, hemostasis, factor XII, plasma kallikrein, contact activation system, coagulation

INTRODUCTION

The contact activation system (CAS) represents a group of plasma proteins, including factor XII (FXII), plasma prekallikrein (PPK), and high molecular weight kininogen (HK) that promotes inflammation and coagulation upon contact of blood with an activating surface or protease (1, 2). CAS activation is initiated by interactions of FXII with a negatively charged surface or a protease.
Contact System in Hemorrhage

that induces a conformational change in FXII leading to its proteolytic cleavage and the generation of the serine protease FXIIa (3, 4). The two primary substrates for FXIIa are PPK and factor XI (FXI). FXIIa-mediated cleavage of PPK results in its zymogen activation to the serine protease plasma kallikrein (PKal), which converts FXII to FXIIa and thereby provides positive feedback amplification of the CAS. The effects of the CAS on inflammation, vascular permeability, and edema are primarily attributed to the Kallikrein–Kinin System. This system involves PKal-mediated cleavage of HK to generate the nonapeptide hormone bradykinin (BK), which activates B2 receptors (B2R) that are expressed on a variety of vascular, neuronal, and immune cell types. Binding of BK to B2R activates proinflammatory signaling pathways that dilate vessels, induce chemotaxis of neutrophils, and increase vascular permeability (5, 6). C1-inhibitor (C1-INH) is the primary physiological inhibitor of both FXIIa and PKal, and C1-INH deficiency facilities CAS activation and BK-mediated angioedema (7). Both PKal and the B2R are clinically significant mediators of hereditary angioedema (HAE) (8, 9). In addition to its effects on inflammation and vascular permeability, BK has been shown to induce expression of both tissue factor (TF) and tissue plasminogen activator (tPA), which activates the extrinsic coagulation pathway and fibrinolysis, respectively (10–13) (Figure 1).

The acute effects of the CAS on coagulation have been primarily attributed to FXIIa-mediated activation of FXI and thereby the intrinsic coagulation pathway leading to fibrin generation. Although the biochemical roles of PKal and FXIIa as upstream triggers for the intrinsic coagulation system were recognized over 50 years ago (14), the clinical significance of these factors on thrombosis and hemorrhage is not fully understood. It is well documented that CAS blockade results in prolonged coagulation times in the activated partial thromboplastin time (aPTT) assay (15). The absence of thrombotic and hemostatic abnormalities in individuals with genetic PPK or FXII deficiency has suggested that the CAS plays a minimal role in physiological coagulation. However, information from rare case studies of individuals with deficiencies in individual components of the CAS has provided limited insight into the potential clinical significance of PKal and FXIIa in coagulation in specific diseases. This is further complicated by the interpretation of acquired and mild CAS component deficiencies, which may reflect the activation of the system rather than its suppression. Key observation from studies of humans with genetic FXII and PPK deficiencies is the absence of spontaneous bleeding disorders and increased bleeding risk during surgery, which are observed in individuals with genetic mutations in downstream in the intrinsic pathway, such as FIX (hemophilia B) and FXI (hemophilia C). The main evidence suggesting that CAS inhibition can reduce pathological thrombosis without increasing bleeding has emerged from studies using PPK and FXII-deficient animals with acute and artificially provoked coagulation (16–20). These clinical and experimental findings have generated increased interest in the potential therapeutic opportunities of targeting the CAS in thrombosis. Indeed the observations of the blockade of the CAS results in reduced thrombosis without a concomitant increase in bleeding in preclinical studies have suggested that therapies that inhibit the CAS may offer novel strategies for safe and effective antithrombotics. Moreover, recent studies suggest that blockade of the CAS can improve hemostasis in the retina and brain. In this review, we discuss emerging understanding of the role of CAS on bleeding and hemostasis and the clinical implications of this research.

HEMOSTASIS

Hemostasis is the process that prevents blood loss following vascular injury. The mechanisms of hemostasis involve both serial and parallel events, mediated by the interactions between circulating factors in blood with components of activated or injured vascular tissue. Given the critical nature of hemostasis, the system has evolved both redundant and reinforcing mechanisms that facilitate both rapid and stable cessation of blood loss. The sequence of events and the relative contributions and requirements of individual coagulation components may vary according to the affected tissue and type of injury, and possibly also the animal species. While thrombosis and hemostasis share a host of coagulation factors, the etiology of these processes has important distinctions, which may create therapeutic opportunities to selectively target thrombosis without affecting hemostasis.

Although animal models do not fully recapitulate the complexity of human physiology and disease, studies on animals with targeted deficiencies in coagulation factors have provided insight into contributions of individual components of coagulation in
hemostatic mechanisms. Multiple approaches are used to quantify hemostatic responses in rodents, including tail tip bleeding, cuticle bleeding, and tail vein transection assays. Deficiencies in FXII and PPK in healthy animals exert little or no effect on tail bleeding time despite aPTT prolongation (Table 1), in contrast to prothrombin deficiency, which markedly increases blood loss in tail wound models (21).

Deficiencies in downstream factors in the intrinsic cascade (such as FIX and FXI) result in bleeding disorders, demonstrating their role in physiological hemostasis. Paradoxically, deficiencies in the CAS factors FXII, PPK, or HK, which trigger coagulation in vitro and contribute to thrombosis in animal models, are not associated with a bleeding diathesis. However, while surgical interventions provide a general assessment of hemostasis, it does not fully model the potential contributions of the CAS in all hemorrhagic conditions. Indeed, recent studies have suggested that the CAS may play an important role in hemorrhage in the retina and brain in certain diseases (17, 20, 30).

### Effects of Kallikrein–Kinin System on Platelet Activation

The initial steps in hemostasis involve vasoconstriction and the local aggregation of platelets to rapidly restrict bleeding at the site of endothelium disruption. Vascular spasm and vasoconstriction, which reduces local blood flow, is triggered by factors released from vascular smooth muscle, endothelial cells, and platelets, and reflexes initiated by the local sympathetic nervous system. Platelets adhere via glycoprotein VI (GPVI) receptors to exposed basement membrane collagen at the site disrupted endothelium to initiate platelet aggregation. This process leads to tethering and activation of platelets, additional platelet receptor-collagen interactions, and platelet plug formation. GPVI play a critical role in normal hemostasis, and its deficiency results in abnormal platelet-responses to collagen and mild bleeding tendency (31).

A report by Liu et al. (17) has shown that PKal inhibits collagen-induced platelet aggregation. PKal also interferes with platelet aggregation induced by collagen-related peptide, a GPVI agonist, but not by convulxin; a snake venom toxin that is a collagen-independent GPVI agonist. Moreover, PKal does not alter either thrombin’s or ADP’s effects on platelet aggregation. These findings suggest that PKal’s effects on platelet activation were specific to GPVI interactions with collagen. Interestingly, the inhibitory effects of PKal on collagen-induced platelet activation were not mimicked with PPK, suggesting that the active form of this protein is required for its inhibitory effect. These findings suggest that PKal binding to collagen in the basement membrane at the site of endothelium disruption interferes with GPVI-mediated platelet activation during initial stages of hemostasis. PKal binding to collagen and its effects on platelet aggregation are increased at elevated glucose concentrations, suggesting that hyperglycemia enhances PKal inhibition of collagen-induced platelet aggregation and thereby may contribute to hematoma expansion in diabetes. Although the mechanism by which glucose increases the binding of collagen to PKal has not been identified, previous reports have suggested that an exposure of collagen to hyperglycemia can alter collagen conformation and facilitate protein binding (32). Interestingly, hyperosmolar concentrations of mannitol, a treatment commonly used to reduce brain edema following stroke, increased hematoma expansion in non-diabetic rats and enhanced PKal’s inhibitory effects on collagen-induced platelet aggregation in vitro. These results suggest that plasma osmolality may alter PKal’s effects on coagulation. The mechanism of PKal interference of collagen-platelet GPVI binding appears analogous to the effects of anopheline antplatelet protein in the saliva of the malaria vector mosquito, which also interferes with collagen-induced platelet activation and hemostasis (33). Further studies are needed to map the domain on PKal that binds to collagen and the factors that modulate this binding. In addition to altering platelet interactions, PKal binding to collagen may retain this enzyme at the site of vascular injury to facilitate its local inflammatory effects. In contrast to the inhibitory effects of PKal on hemostasis, FXII activation by collagen and laminin in the subendothelial matrix can lead to thrombin generation, which may promote secondary mechanisms of platelet activation at the site endothelium disruption (34, 35). Although these findings suggest that the CAS can exert both positive and negative effects on platelet activation during hemostasis, gene targeting and pharmacological studies indicated that blockade of CAS reduces cerebral hemorrhage, which is a primary area of concern regarding bleeding in patients on antithrombotic therapy.

### TABLE 1 | Effects of contact activation system inhibition and deficiencies in experimental models.

| Condition | aPTT | Tail bleeding | Spontaneous bleeding | Species | Reference |
|-----------|------|---------------|----------------------|---------|-----------|
| FXII−/−   | Prolonged | No changes | Not reported | Mice | Renne et al. (16) |
| FXII−/−   | Prolonged | No changes | Not reported | Mice | Iwaki et al. (22) |
| Anti-FXII (9A2 and 15H8) | Prolonged | No changes | No | Mice | Matafonov et al. (23) |
| FXII ASO | Not reported | No changes | No | Mice | Merkulov et al. (28) |
| FXII−/− or infestin-4 | Prolonged | No changes | No | Mice | Liu et al. (17) |
| Infestin-4 | Prolonged | No changes | No | Rat | Cai et al. (27) |
| FXII−/− or FXII sRNAi | Prolonged | No changes | No | Mice | Revenko et al. (24) |
| PPK ASO | Not reported | No changes | No | Mice | Järnkron et al. (29) |
| Ktkb1−/− | Prolonged | No changes | Not reported | Mice | Liu et al. (17) |
| PPK ASO | Not reported | No changes | No | Mice | Bird et al. (18) |
| Kng1−/− | Prolonged | No changes | Not reported | Mice | Merkulov et al. (28) |
| Bdkrb2−/− | Prolonged | Prolonged | Not reported | Mice | Shariati-Madar et al. (29) |

ASO, antisense oligonucleotide; aPTT, activated partial thromboplastin time; infestin-4 is an FXIIa inhibitor; 9A2 and 15H8 are anti-FXII inhibitory antibodies; FXII, factor XII; PPK, plasma prekallikrein.
Effects of Kallikrein–Kinin System on Fibrin Formation

Stabilization of the nascent thrombus involves additional coagulation factors that generate and crosslink fibrin, which stabilizes nascent platelet plugs. FXIIa-mediated activation of FXI results in the activation of the intrinsic coagulation cascade, leading to thrombin activation and fibrinogen cleavage and the stabilization and expansion of clots. Although FXIIa can serve as a trigger to activate the intrinsic pathway from the CAS, positive feedback activation of FXI by thrombin can also activate this pathway following extrinsic activation (36–39). Thus, while FXII appears to contribute to arterial thrombus formation and growth, at least in animal models, FXII does not appear to be required for clot formation during hemostasis. Activated platelets actively contribute to this secondary hemostatic process by generating surface-bound polyphosphate nanoparticles (40), which activates FXII and thereby promotes local fibrin generation by the clot. Although FXII, PPK, or HK deficiency is not associated with bleeding, FXI deficiency in patients is associated with mild bleeding, suggesting that the role of FXI in hemostasis is independent of the CAS. The upstream activation of FXI by thrombin may explain why FXII is not required for hemostasis in contrast to FXI, where its deficiency is associated with hemophilia C.

Clinical trials and epidemiologic data indicate that FXI contributes to thromboembolic diseases. Studies with animal models also suggest that the activation of FXI by FXIIa promotes physiological thrombus formation (41). Furthermore, “exposed” collagen also initiates the contact phase of coagulation by binding to FXII and enhances coagulation in vitro (35). The conversion into active FXIIa is dependent on repetitive negative charge exposed by collagen fibrils (42). van der Meijden and colleagues (43) showed that collagen potentiates thrombus formation via binding to FXII, leading to its activation and subsequent FXI activation. This report suggests a dual role for collagen in thrombin generation by stimulating platelet activation via GPVI and by direct effects on FXII. In addition, FXIa-mediated thrombin activation may contribute to clot stability thrombin-activatable fibrinolysis inhibitor-mediated removal of lysine residues on fibrin, which increases its resistance to fibrinolysis (44). Moreover, FXIa proteolytically degrades and neutralizes TF pathway inhibitor, which can promote thrombin generation via the extrinsic pathway (45). Recently, Stavrous and colleagues (13) demonstrated that PPK deficiency in mice increase prostacyclin through Sirt1 and KLF4, which reduced vascular TF expression and thrombosis. Thus, while CAS’s effects on both the intrinsic and extrinsic pathways contribute to clot stabilization, mechanistic redundancy contributing to fibrin generation minimizes the requirements of the CAS in this process.

**EFFECTS OF PPK, FXII, AND C1-INH DEFICIENCY ON COAGULATION IN HUMANS**

Most reports have indicated that individuals with severe deficiencies in either PPK or FXII do not display abnormalities in either thrombosis or bleeding (Table 2), and often go undiagnosed for decades without apparent clinical phenotype. The absence of a bleeding diathesis in FXII and PPK deficiencies appears to contradict the key role of these proteins in coagulation via the intrinsic pathway in vitro, and this has left the physiological significance of the CAS in coagulation and thrombosis unclear. However, CAS could be a trigger for thrombosis associated with extracorporeal circuits (46) and cardiopulmonary bypass (47) by exposing blood to artificial surfaces, non-physiological shear stress, and osmotic forces. Administration of an antibody that neutralizes FXIIa (3F7) has been shown to have a similar effect as heparin in preventing thrombosis during extracorporeal circuits (48).

Activation of FXII by kaolin (negatively charged aluminum silicate particulate) provides the basis for the aPTT, a clinical clotting assay, which is widely used to assess the integrity of the intrinsic pathway and to monitor anticoagulation with heparin. Deficiencies in FXII, PPK, and HK results in markedly prolonged aPTT and no increases in spontaneous bleeding or impairment in hemostasis (Table 2). Normalization of a severely increased aPTT (>120 s) after prolonged preincubation with aPTT reagent occurred in plasma deficient in PPK but not in plasma deficient in FXII, HK, FXI, FIX, FVIII, and Passovoy trait plasma deficiency characterized by abnormal coagulation affecting the intrinsic coagulation system or plasma containing lupus anticoagulant. Autoactivation of FXII in PPK-deficient plasma in the presence of kaolin paralleled the normalization of aPTT. The addition of OT-2, a monoclonal antibody neutralizing FXII, prevents the normalization of aPTT (66). These results suggest that autoactivation of FXII is responsible for normalization of a severely prolonged aPTT upon increased preincubation time in PPK-deficient plasma.

In humans, PPK deficiency is not associated with hemorrhagic disorders, indicating PKal is not required for hemostasis. Although some case reports have associated FXII deficiency with prothrombotic events, a clinical study failed to demonstrate a significant correlation between increased risk of thrombosis and FXII deficiency (78). Girolami and colleagues (79) re-evaluated case reports on FXII deficiency and thrombosis. They showed that, in most cases, FXII deficiency was associated with other congenital or acquired prothrombotic risk factors. While these studies indicate that the CAS is not essential for either thrombosis or hemostasis, individuals with genetic deficiencies in the CAS are rare and it is not possible to ascertain from the anecdotal clinical information whether PKal and FXIIa could have significant roles in thrombosis or hemostasis associated with specific clinical indications.

C1-inhibitor is the primary endogenous inhibitor of both PKal and FXIIa. HAE is a rare genetic disease caused by deficiencies in C1-INH concentration (Type I) or C1-INH activity (Type II) (80). HAE has a prevalence of approximately 1:50,000 and is characterized by episodes of vasogenic edema, which vary in severity and affected tissue, and can become life threatening when affecting the larynx. The primary mechanism for this disease is the uncontrolled activity of PKal, which results in increased cleavage of HK, and increased production of BK. While the molecular and physiological triggers for the onset of attacks are not fully understood, increased levels of FXIIa is observed...
tally induced thrombosis (16, 24, 26, 28), indicating the potential
or HK genes have been shown to protect mice from experimen-
Pharmacological inhibition of FXIIa and deficiency in either FXII
or requirements of additional factors not activated by the CAS to
apparent clinical coagulopathies in HAE during attacks and/
subjects with HAE during an attack (81, 83, 84). The absence of
thrombin-anti-thrombin complex, and FVIIa are elevated in
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TABLE 1 | Case reports of deficiencies in contact activation system proteins.

| Deficiency | aPTT | Thrombosis and clinical presentation | Spontaneous bleeding | Reference  
|------------|------|---------------------------------|---------------------|----------- |
| FXII       | Prolonged | Bilateral femoral vein thrombosis | No           | Cei et al. (49) |
| FXII (mild)| Slightly prolonged | Venous thrombosis | No           | Lessiari et al. (50) |
| FXII       | Prolonged | Coronary artery bypass grafting | No           | Conaglen et al. (51) |
| FXII       | Prolonged | Arterial and venous thrombosis | No           | Hellsten et al. (52) |
| FXII       | Prolonged | Coronary artery bypass grafting | No           | Moorman et al. (53) |
| FXII (mild)| Prolonged | Occlusive thrombus in the circumflex and anterior descending arteries | No           | Penny et al. (54) |
| FXII       | Prolonged | Coronary artery bypass grafting | No           | Rygal and Kuc (55) |
| FXII       | Prolonged | Coronary artery disease | No           | Cronbaugh et al. (56) |
| FXII       | Prolonged | Coronary artery bypass grafting | No           | Salmender et al. (57) |
| FXII       | Prolonged | No thrombosis | No           | van Veen et al. (58) |
| FXII       | Prolonged | Coronary artery disease | No           | Wood (59) |
| FXII (mild)| Prolonged | Bilateral lower limb deep vein thrombosis | No           | Vergnes et al. (60) |
| FXI        | Prolonged | Retinal venous thrombosis | No           | Borrego-Sanz et al. (61) |
| FXII (mild)| Prolonged | Deep vein thrombosis, after abdominal surgery | No           | Cornuckella et al. (62) |
| FXII       | Prolonged | Cardiopulmonary bypass | No           | Gerhardt et al. (63) |
| PPK        | Prolonged | No thrombosis | No           | van Veen et al. (58) |
| PPK        | Prolonged | Coronary artery disease | No           | Oram et al. (64) |
| PPK        | Prolonged | No thrombosis | No           | Cankovic et al. (65) |
| PPK        | Prolonged | No thrombosis | No           | Asmis et al. (66) |
| PPK        | Prolonged | No thrombosis | No           | Maak et al. (67) |
| PPK        | Prolonged | No thrombosis | No           | Dela Cadena (68) |
| PPK        | Prolonged | Ischemic stroke | No           | Francois et al. (69) |
| PPK        | Prolonged | No thrombosis | No           | Nakao et al. (70) |
| PPK        | Prolonged | No thrombosis | No           | Lombardi et al. (71) |
| PPK        | Prolonged | No thrombosis | No           | Wynne Jones et al. (72) |
| PPK        | Prolonged | No thrombosis | No           | Poon et al. (73) |
| HK         | No thrombosis | No thrombosis | No           | Cankovic et al. (65) |
| HK         | No thrombosis | Cardiopulmonary bypass | No           | Davidson et al. (74) |
| HK         | No thrombosis | No thrombosis | No           | Lefere et al. (75) |
| HK         | No thrombosis | Vertebral-basilar artery thrombosis following trauma | No           | Stormorken et al. (76) |

aPTT, activated partial thromboplastin time; FXII, factor XII; PPK, plasma prekallikrein; HK, high molecular weight kininogen; NA, not available.

In this disease (81) which suggest that attacks are mediated by
the CAS. Since HAE is not associated with coagulopathies, this
disorder provides an opportunity to evaluate the effects of the
CAS on downstream coagulation pathways, independent of
confounding input associated with comorbidities of thrombosis
and hemostasis. Indeed, plasma obtained from HAE subjects
during edematous attacks display shortened aPTT, compared
to plasma collected from these subjects during remission (82,
83). In addition, FXIa activity, prothrombin fragment F1 + 2,
thrombin-anti-thrombin complex, and FVIIa are elevated in
subjects with HAE during an attack (81, 83, 84). The absence of
apparent clinical coagulopathies in HAE during attacks could be
due to compensatory mechanisms that occur during attacks and/
or requirements of additional factors not activated by the CAS to
initiate or promote thrombosis formation.

**EFFECTS OF PPK AND FXII DEFICIENCY ON COAGULATION IN RODENTS**

Consistent with PPK and FXII deficiencies in humans, PPK,
and FXII-deficient mice display prolonged aPTT (Table 1).
Pharmacological inhibition of FXIIa and deficiency in either FXII
or HK genes have been shown to protect mice from experiment-
tally induced thrombosis (16, 24, 26, 28), indicating the potential
role of the CAS in thrombotic disease. Cheng and colleagues
(41) demonstrated that a neutralizing antibody to FXI (14E11)
had a comparable effect to FXI deficiency in a FeCl3 model of
thrombosis. Antibody 14E11 binds FXI and interferes with FXI
activation by FXIIa, suggesting that thrombus formation in this
model requires FXI activation by FXIIa. These data have gener-
ated interest in developing strategies to therapeutically inhibit
FXIIa and contact activation to treat or prevent thromboembolic
disorders. Interestingly, thrombosis protection in mice was
greater with FXII deficiency than with FXI deficiency, suggesting
that FXII may exert effects on coagulation that do not require
FXI (41).

Factor XII-deficient mice showed defective thrombus forma-
tion (16) and protective effect against experimental ischemic
stroke (85) and pulmonary embolism (86). Intravital microscopy
showed that the initial adhesion of platelets at the site of injury is
not affected by FXII deficiency, although the formation and sta-
bilization of three-dimensional thrombi is impaired (16). Similar
defective thrombus formation was observed in FXI−/− mice (87, 88).
Furthermore, FXII deficiency or pharmacological inhibitor of
FXIIa showed decreased infarct volume without the presence of
intracerebral hemorrhage (ICH) (85). Reconstitution of FXII-
deficient mice with human FXII protein restored susceptibility
for ischemic stroke. A recent report by Nickel and colleagues
(25) has shown that prostasomes released from prostate cancer activated FXII and that both FXI- and FXII-deficient mice are protected against cancer-associated thrombosis (25). However, the observation that PPK-deficient mice were not protected in the prostasome-induced thrombosis model is surprising and appears different than other CAS-associated thrombosis models, which require PPK. Taken together, these findings suggest that coagulation mediated by intrinsic pathway through FXII is critical for pathological thrombus formation in certain conditions without being relevant for hemostasis.

In addition, the CAS mediates effects on thrombosis via the kallikrein–kinin system. Genetic ablation of Kng1 in mice showed a delayed time to carotid artery occlusion in a laser injury model (28). BKB2R−/− mice have prolonged bleeding time and delayed carotid artery occlusion times in the rose Bengal thrombosis model and these effects were attributed to increased expression of angiotensin receptor 2 and elevated nitric oxide and prostacyclin levels (29). These findings suggest that the BK system contributes to the effects of the CAS on coagulation.

**POSSIBLE ROLE OF THE KALLIKREIN–KININ SYSTEM IN HEMORRHAGE**

Components of the CAS are normally restricted from contact with the vascular basement membrane by the endothelium. Vascular hyperpermeability enables components of the CAS to leak into the subendothelial space and gain contact with the basement membrane. The potential functions of the CAS in the subendothelial space have received relatively little attention. PKal binds to collagen (89), which may contribute to the retention of its catalytic activity in the extracellular matrix (ECM) at the site of injury and thereby enable local amplification of its inflammatory and edematous actions. Moreover, PKal in this subendothelial space may have additional substrates and functions, which are not apparent in the plasma.

**Diabetic Retinopathy**

In diabetic retinopathy, retinal vascular hyperpermeability can lead to diabetic macular edema (DME); the leading cause of vision loss in working-age adults in most developed countries (90). Proteomic analysis of vitreous fluid from patients with DME has revealed increased concentrations of PPK, HK, and FXII in this fluid compared with people without DME (91–93). These increases have been attributed to the increased diffusion of these CAS components from the blood into the neuroretinal interstitial fluid and vitreous humor. Recent reports have shown that intravitreal injection of PKal alters both retinal vascular function and ultrastructure. A single injection of PKal in the vitreous of rats induced retinal vascular hyperpermeability and retinal layer thickening (94), whereas two injections of purified PKal into the vitreous caused retinal bleeding that appeared similar to retinal microhemorrhages that occur in diabetic retinopathy (30). While injections of BK mimicked PKal’s effect on retinal permeability and thickening at 24 h repeated injections of BK did not cause retinal bleeding at 48 h. These findings suggest that PKal in the subendothelial space may contribute to retinal hemorrhages, which are a hallmark of advanced stages of diabetic retinopathy. Proteomics has been used to investigate the effects of PKal that may contribute to this disruption of the blood–retinal barrier. Analyses of conditioned media from both cultured pericytes and astrocytes incubated with PKal have revealed increases in proteolytic fragments of ECM proteins, including multiple collagen isoforms 1–6, laminin β1 and γ1, nidogen 1 and 2, and fibronectin, compared with conditioned media from cells that were not exposed to PKal (30, 95). In a purified system, PKal was shown to cleave COL4A, suggesting that PKal can proteolytically cleave collagen (30). Although the causes of retinal bleeding in diabetic retinopathy are not yet known, these studies suggest a mechanism by which vascular permeability, via the extravasation of PKal, may contribute to retinal hemorrhage, in part, mediated by the proteolytic degradation of the vascular basement membrane.

**Cerebral Hematoma Formation**

Diabetes and hyperglycemia are associated with increased ICH and worse clinical outcomes following a cerebrovascular accident (96–98). Diabetes and acute hyperglycemia in non-diabetic rodents increase hematoma formation in an experimental model of ICH (17). In this report by Liu et al., intracerebral injection of autologous blood induced hematoma expansion, which was ameliorated by PPK deficiency and PKal inhibition. Moreover, intracerebral injection of purified PKal mimicked the effects of autologous blood injection on hematoma expansion in diabetic animals. PKal has been reported to mediate plasminogen activation (99, 100), which could potentially increase fibrinolysis in the ICH model. However, intracerebral injection of neither tPA nor plasmin mimicked the effects of PKal on hematoma expansion in diabetic animals (17). Moreover, covalently deactivated PKal also induced hemorrhage in this model, suggesting that PKal’s effects in this model were not mediated by its catalytic activity. The effects of hyperglycemia and PKal on ICH are rapid, within 30 min, suggesting an effect on an early step in hemostasis. Since in vitro studies have shown that PKal binding to collagen interferes with its effects on GPVI-mediated platelet activation, these findings suggest that PKal may decrease platelet plug formation, which plays an early step in hemostasis.

**Thrombolysis**

Tissue plasminogen activator is the only approved treatment for thrombotic stroke (101). While its prompt administration following stroke onset has been shown to improve clinical outcomes, its use is limited due to an increased risk of intracranial hemorrhage when tPA is used after the recommended 3 h therapeutic window, which can negate the potential benefits of vascular recanalization. Although the mechanisms that mediate the increase in hemorrhage induced by tPA are not fully understood, we have shown that both PPK and FXII deficiency markedly reduce tPA-induced hemorrhagic conversion in mice with a thrombotic middle cerebral artery occlusion (20). Studies using purified proteins, as well as plasma, revealed that plasmin activates FXII and cleaves it into a fragment with a molecular weight slightly higher than the expected size of FXIIa light chain generated with PKal (20, 102–105). Plasmin cleaves
FXII at Arg353 leads to zymogen activation and the generation of FXIIa (104). Plasmin’s effects on FXIIa activity is markedly increased in the presence of dextran sulfate (104) and inhibited by e-aminocaproic acid and a FXIIa inhibitory mAb (105). These reports have suggested that plasmin-mediated cleavage of FXII increases FXIIa-like activity. The consequences of plasmin-mediated cleavage of FXII at Lys346 on FXIIa-like catalytic activity are not yet available. Plasmin-mediated cleavage of FXII could facilitate its autoactivation and/or cleavage by PKal in plasma, which would thereby activate the kallikrein–kinin system. These findings have suggested that the CAS may either worsen vascular damage or impair hemostasis in the cerebral vasculature following tPA administration. Studies using human plasma have shown that tPAs effects on the CAS are mediated by plasmin (20), which has been identified as a physiological activator of the CAS (104, 105). Although the clinical significance of this pathway in hemorrhagic conversion in ischemic stroke is not yet available, two lines of evidence support the conclusion that tPA therapy activates the CAS in stroke patients. First, a recent report by Marcos-Contreras and colleagues has shown that intravenous infusion of tPA in patients result in plasma HK cleavage (106), suggesting that the tPA activates the circulating CAS. Second, tPA therapy can cause orolingual angioedema that has been attributed to increased BK action (107, 108, 109), suggesting that tPA increases CAS activity. In addition to reducing hemorrhage, PPK, and FXII deficiency also reduced infarct volume and cerebral edema in mice with stroke treated with tPA. These findings are consistent with previous reports showing that blockade of the CAS has neuroprotective effects in mice with filament-mediated middle cerebral artery occlusion (85, 110). Although it is tempting to speculate that inhibition of either PKal or FXIIa during the administration of tPA may ameliorate hemorrhagic transformation and provide neuroprotection during thrombolytic therapy, additional information on the role of the CAS on stroke outcomes in animal models is needed.

**THERAPEUTIC IMPLICATIONS AND CONCLUSION**

Currently available anticoagulants used for prevention or treatment of thromboembolic events [heparins, vitamin K antagonists (for example, warfarin), and inhibitors of thrombin or factor Xa] all target enzymes of the coagulation cascade that are essential for the formation of fibrin, a protein necessary for controlling injury-related blood loss. As a result, currently used anticoagulants increase the risk of bleeding and are associated with an increase risk in potentially life-threatening hemorrhage (111). Bleeding is the primary complication of anticoagulation therapy and a significant risk of all currently used anticoagulants, even when maintained within their therapeutic ranges (112). The CAS exerts effects on coagulation at multiple levels. CAS activation of PPK has been implicated in promoting spontaneous microvascular bleeding and the impairment of collagen-induced platelet activation. In addition, both PPK and FXII, via activation of the intrinsic coagulation cascade promotes clot stabilization and growth. Feedback activation of FXI by thrombin contributes to clot stabilization and may explain the lack of dependence of hemostasis on FXII and PPK. PKal has been implicated in promoting spontaneous microvascular bleeding and the impairment of collagen-induced platelet activation. Pharmacological blockade of PKal has been shown to provide beneficial effects on cerebral hemostasis in animal models. Although inhibition of the CAS reduces provoked arterial and venous thrombosis in animal models, the clinical significance of the CAS in thrombosis is not yet available. The clinical indications for targeting the CAS for thrombosis will require the identification of thrombotic processes that are dependent on the CAS, which potentially include interactions with artificial surfaces and disease processes that generate factors that activate FXII. In addition, inhibition of the CAS may provide an opportunity to reduce cerebral hemorrhage, which is one of the primary concerns of increased bleeding risk associated with current antithrombotics.

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FS and EF performed literature searches and wrote the manuscript.

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