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Proteolytic and Nonproteolytic Activation Mechanisms Result in Conformationally and Functionally Different Forms of Coagulation Factor XIII A

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Running title: Functionally different forms of Factor XIII A (44/50 characters)

Abbreviations: FXIII A, the A-subunit of coagulation Factor XIII; AP, Activation Peptide; ECM, extracellular matrix; FXIII A*, Factor XIII A activated by thrombin in the presence of 4 mM CaCl₂; FXIII A^{o,low}, Factor XIII A activated nonproteolytically in the presence of 4 mM CaCl₂; FXIII A^{o,high}, Factor XIII A activated nonproteolytically in the presence of ≥25 mM CaCl₂; PPACK, D-Phe-Pro-Arg chloromethyl ketone; AUC, analytical ultracentrifugation; DMSO, dimethyl sulfoxide; SEM, scanning electron microscopy; MDC, monodansylcadaverine; DMPDA, N,N-dimethyl-1,4-phenylenediamine.

Enzymes: Factor XIII A (EC 2.3.2.13).

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ABSTRACT

Factor XIIIa (FXIIIa) is a transglutaminase that crosslinks intra- and extracellular protein substrates. FXIIIa is expressed as an inactive zymogen, and during blood coagulation, it is activated by removal of an activation peptide by the protease thrombin. No such proteolytic FXIIIa activation is known to occur in other tissues or the intracellular form of FXIIIa. For those locations, FXIIIa is assumed instead to undergo activation by Ca^{2+} ions. Previously, we demonstrated a monomeric state for active FXIIIa. Current analytical ultracentrifugation and kinetic experiments revealed that thrombin-activated FXIIIa has a higher conformational flexibility and a stronger affinity toward glutamine substrate than does nonproteolytically activated FXIIIa. The proteolytic FXIIIa activation was further investigated in a context of fibrin clotting. **In a series of fibrin crosslinking assays and scanning electron microscopy studies of plasma clots, the activation rates of FXIIIa V34X variants was correlated with the extent of fibrin crosslinking and incorporation of nonfibrous protein into the clot.** Overall, the results suggest conformational and functional differences between active FXIIIa forms, thus expanding the understanding of FXIIIa function. Those differences may serve as a basis for developing therapeutic strategies to target FXIIIa in different physiological environments.

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INTRODUCTION

Factor XIII (FXIII) has been a subject of medically related research for almost a century. Perhaps the most studied is the role of FXIII in crosslinking the fibrin network, making it more mechanically stable and resistant to fibrinolysis. As physiological knowledge on FXIII expands, it becomes apparent that beyond serving as a key player of the blood coagulation system, FXIII functions in wound healing, bone tissue dynamics, signaling, and other areas [1-10].

FXIII is a transglutaminase that crosslinks protein substrates via an isopeptide bond in a Ca^{2+} -dependent manner. FXIII is expressed as an inactive zymogen and in plasma, it circulates in a heterotetrameric complex with carrier FXIIB (A_2B_2). During blood coagulation, plasma FXIII is activated by thrombin-mediated removal of N-terminal activation peptides (AP) from the A-subunits followed by binding of Ca^{2+} and dissociation of the B-subunits [11]. The active FXIII then introduces covalent crosslinks between polymerizing fibrin molecules and incorporates other proteins into the network, ultimately increasing mechanical stability of the resulting clot and its resistance to fibrinolysis [12].

Intracellularly, such as in platelets, monocytes, and macrophages, FXIII exists as an A_2 -homodimer and is thought to undergo slow nonproteolytic activation in the presence of low available Ca^{2+} concentrations [13-15]. In this physiological compartment, FXIII is present in the cytoplasm, associates with membrane [10, 16], and even appears in the nucleus [17]. Activation and translocation of FXIII from the platelet cytoplasm to the platelet membrane does not require thrombin dependent steps. The minor amount of transglutaminase activity observed in a subsequent fibrin-platelet environment is proposed to be due to non-proteolytic activation of FXIII A_2 [18]. Some other intracellular functions of FXIII include reorganization

of cytoskeletal proteins and chromatin remodeling [9]. FXIIIa is also secreted by osteoblasts into the extracellular matrix (ECM) [19], where it contributes to formation of the ECM itself and remodeling of the bone tissue [6, 16, 20]. Although a 37 kDa proteolytic fragment of FXIIIa was proposed to exist in the bone ECM [19, 21], it was later identified as transaldolase-1 that was immunoreactive with anti-FXIIIa antibody [22]. Thus, no evidence of proteolytic FXIIIa activation in bone ECM currently exists. However, with available ECM Ca^{2+} concentrations as high as 25 – 40 mM [23, 24], bone ECM FXIIIa may be activated by Ca^{2+} without proteolysis.

FXIIIa has also been implicated in pathological conditions such as thrombosis [25], inflammation, oncogenic events and diabetes [10], and arthritis [26]. Although FXIIIa is involved in an array of intra- and extracellular events, an in-depth understanding of FXIIIa activation and function in different physiological environments is lacking. With more knowledge, therapeutic strategies might be developed to target zymogen FXIII or activated FXIIIa under specific physiological or pathological conditions. Previously, we studied the oligomeric states of FXIIIa in different solution conditions [27]. We found that FXIII A_2 -homodimer dissociates into monomers during activation by thrombin. Cleavage of a single AP on the dimeric FXIII A_2 was sufficient to promote dissociation and full activity of the A-subunits. Nonproteolytic activation by high Ca^{2+} concentration also resulted in subunit dissociation, however, these species possessed lower activity [27].

In the current project, we hypothesized that in different physiological compartments such as blood plasma, intracellular, and bone ECM, activation of FXIIIa may result in conformationally different enzymatic species. We focused on probing functional implications of thrombin activation of FXIIIa in the presence of low mM Ca^{2+} (mimicking conditions in plasma, the resulting active species are henceforth denoted as FXIIIa*), nonproteolytic activation by low

mM Ca^{2+} (intracellular activation, $\text{FXIII}^{\text{o,low}}$), and activation in the presence of high (≥ 25 mM) Ca^{2+} (bone ECM, $\text{FXIII}^{\text{o,high}}$). We demonstrated an overall slower rate of FXIII nonproteolytic activation, as compared to the thrombin-mediated cleavage. Moreover, enzymatic activity assays revealed differences in the substrate affinities of FXIII^* and $\text{FXIII}^{\text{o,high}}$. Using a fibrin clotting model in a pure protein system as well as in plasma, we further examined how proteolytic activation of FXIII V34X variants can be used to control rate of fibrin crosslinking function and also influence fibrin clot structure. In summary, our results support proposed conformational differences between FXIII active forms in different physiological environments. Finally, we discuss how these differences may serve as a basis for differential therapeutic targeting of FXIII .

RESULTS

Solution properties of FXIII species in different activation conditions. Previously in a series of AUC experiments, we demonstrated that zymogenic FXIII A_2 homodimer completely dissociates into monomers upon proteolytic and nonproteolytic activation [27]. In the current work, we employed sedimentation velocity AUC to compare the solution dynamics of FXIII under different conditions. FXIII was activated in the presence of 100 mM Ca^{2+} ($\text{FXIII}^{\text{o,high}}$) or by thrombin in the presence of 4 mM Ca^{2+} (FXIII^*). The high 100 mM Ca^{2+} helped to quickly obtain uniform, monomeric $\text{FXIII}^{\text{o,high}}$ species [27]. As a control for a possible divalent cation-mediated ionic strength effect, additional samples of FXIII^* were supplemented with 100 mM Mg^{2+} . Unlike Ca^{2+} , this divalent cation does not efficiently support dissociation or activity of FXIII A-subunits [27]. To eliminate possible differences in sample handling between separate analytical runs, all sample conditions were subjected to AUC at the same time. The

resulting sedimentation coefficient distributions are presented in Fig. 1A. In the presence of 100 mM Ca^{2+} (FXIII^A^{o,high}, trace i), we observed a tall narrow peak, approximately 1 S unit in width. As reported previously [27], significant precipitation occurred in FXIII^A* samples during centrifugation, and the sedimentation distribution peak was much broader, approximately 2 S (trace ii). In the presence of 4 mM Ca^{2+} and 100 mM Mg^{2+} (trace iii), the FXIII^A* sedimentation peak was also broad (2 S).

Sedimentation coefficients represent the velocity of particle sedimentation (in the present case, FXIII^A molecules) in a centrifugal field [28]. Apart from the physical characteristics of a solution, which are accounted for in data analysis, the sedimentation coefficient ultimately depends on the weight and shape of those particles. Thus, in samples of monomeric FXIII^A, it is the diversity of shapes (conformations) that constitutes the sedimentation distribution. The broad sedimentation distribution peak indicates that FXIII^A* is more conformationally heterogeneous in solution than FXIII^A^{o,high}, and this heterogeneity is retained in the presence of 100 mM Mg^{2+} . This observation suggests that Ca^{2+} exerts a specific effect on FXIII^A conformation. The related divalent cation Mg^{2+} cannot mimic this property and, moreover, the conformational effect is not due to a simple increase in ionic strength.

To examine FXIII^A nonproteolytic activation at lower, more physiological Ca^{2+} concentrations, FXIII^A was incubated in the presence of 4 mM Ca^{2+} for 30 min – 96 h, followed by AUC. Unexpectedly, progressive precipitation was observed in these samples of FXIII^A^{o,low}, very similar to the solution behavior of the thrombin-activated FXIII^A*. In previous experiments, we found addition of 5% DMSO promoted solubility of FXIII^A* over the course of a few hours [27]. However, 4 mM Ca^{2+} -activation required longer incubation periods to obtain detectable results, and even addition of DMSO did not aid in full solubility of activated FXIII^A.

The actual amount of protein in solution was assessed via absorbance at the start of an AUC experiment (Fig. 1B, insert). The plot in Fig. 1B (open circles) then represents the amount of monomeric FXIII_A as a fraction of soluble protein in each sample. The most observed dissociation of the A-subunits, ~ 65%, was observed after 72 h incubation. Such nonproteolytic activation of FXIII_A may proceed intracellularly, although much slower than observed in the presence of 4 mM Ca²⁺, as intracellular Ca²⁺ is actually maintained at nanomolar to micromolar levels. Furthermore, the aggregation propensity of FXIII_A^{o,low} suggests its relatively short lifetime, in a good agreement with a study by Muszbek et al. [13], reporting a small, 6.5% of FXIII_A population within platelets being active.

In order to mimic activation conditions of bone tissue ECM, FXIII_A was incubated in the presence of higher, 25 mM Ca²⁺ for 30 min – 6 h (Fig. 1B, filled circles). The dissociation of the dimeric zymogen under these conditions occurred faster than at 4 mM Ca²⁺ (80% at 6 h of incubation) and, analogous to activation by 100 mM Ca²⁺, no precipitation of these FXIII_A^{o,high} species was observed (Fig. 1B, panel on the right). Thus, the current results indicate overall slower activation of FXIII_A by Ca²⁺ alone, as compared to the proteolysis by thrombin resulting in full dissociation of FXIII_A within minutes. High (≥ 25 mM) Ca²⁺ levels reduce conformational heterogeneity and aid in maintaining stability of activated FXIII_A species in solution.

A model system of FXIII_A proteolytic activation kinetics. To better understand FXIII_A function in different physiological contexts, the peculiarities of individual activation environments need to be considered. Due to highly regulated calcium homeostasis in the body, nonproteolytic activation pathways leading to FXIII_A^{o,low} and FXIII_A^{o,high} likely operate on an ongoing basis. There will thus be slow, constitutive FXIII activity occurring within cells and

ECM. Faster activation may then be achieved when higher Ca^{2+} concentrations are encountered as part of certain physiological or pathological processes. In blood plasma, where Ca^{2+} concentration is maintained at a low 1.5 mM, such constitutive generation of FXIII^{a,low} is prevented by association of FXIII^a with the inhibitory FXIII^b [15]. Under exigent cases such as bleeding, the proteolytic removal of AP facilitates fast FXIII^a activation. The concomitant use of the serine protease thrombin to activate FXIII^a and also convert fibrinogen to polymerizing fibrin adds complexity to this clotting mechanism. [29]. As a result, the kinetics of FXIII^{a*} generation has a significant impact on the functional outcome of the clotting process.

To explore this outcome, we generated and expressed FXIII^a variants with different amino acid residues at position 34 of the FXIII activation peptide. These variants differ in their rate of cleavage by thrombin [30] (Fig. 2A). The current studies provided the first testing of AP residues F34 and W34 on FXIII^a proteolytic activation and resulting transglutaminase function. Using an extended range of FXIII activation rates, comparisons could be made with the naturally occurring V34 and L34. The FXIII^a AP variants were combined with human FXIII-free fibrinogen and clotting was initiated by addition of thrombin and Ca^{2+} . As expected, the rate of FXIII^a proteolytic activation determined development of FXIII^{a*} crosslinking activity (Fig. 2B). The fastest cleaved FXIII^a L34 generated both γ - γ and higher molecular weight (HMW) fibrin species already at an early stage (5 min) of the experiment. Slow cleavage of FXIII^a W34 by thrombin resulted in the slowest appearance of γ - γ and very little HMW crosslinks. Interestingly, F34 FXIII^a is cleaved by thrombin slightly slower than V34 variant (Fig. 2A); however, both variants crosslinked fibrin to a similar extent. This effect may be explained by the fact that the cleavage of a single AP on dimeric FXIII^a zymogen results in dissociation and promotes full activity of both cleaved and noncleaved A-subunits [27, 31]. In addition, a

competition of fibrinogen and FXIIIa for thrombin may have further masked the difference in V34 and F34 cleavage rate observed in the absence of fibrinogen. The four full-length FXIII V34X variants could thus be used to control the timing of appearance of FXIII activity without diminishing thrombin-mediated conversion of fibrinogen to fibrin or altering the clot gelation dynamics. Moreover, the extent of crosslinking before initiation of fibrinolysis could be documented.

To further probe the functional impact of FXIIIa activation rate, three AP variants that demonstrated different degrees of fibrin crosslinking (L34, V34, and W34) were introduced into FXIIIa-deficient murine plasma, and clotting was initiated by addition of thrombin and Ca^{2+} . The murine plasma system was already well established for introducing and screening FXIII V34, V34L, and/or G33A based clotting effects [32-34]. The resulting clots were imaged by SEM and representative photographs are shown in Fig. 3. Clots formed in the presence of FXIIIa better retained their three-dimensional structure due to FXIIIa-mediated covalent crosslinking of the fibrin fibers. Interestingly, we observed nonfibrous protein networks embedded to different degrees within the crosslinked fibrin clots. The least amount occurred with the FXIII AP- variant W34 (slower proteolytic activation) and the most with the L34 (faster activation). Such networks were uniformly distributed and must be covalently crosslinked within the clot, since they remained even after extensive washing during SEM sample preparation. The nonfibrous material may represent unpolymerized fibrin or other plasma proteins and is expected to contribute to overall clot penetrability and mechanical and fibrinolytic stability.

Catalytic comparisons of FXIIIa* and FXIIIa^{o,high} Previously, when using a standard coupled ammonia release assay [35], we observed lower catalytic activity for FXIIIa^{o,high}, as

compared to FXIII^A* [27]. In the current work, we attempted to probe this difference in more detail. The transglutaminase activities of FXIII^A activated by thrombin in the presence of 4 mM Ca²⁺ (FXIII^A*) or by 100 mM Ca²⁺ (FXIII^{o,high}) were compared in a series of enzymatic evaluations. Since it was impossible to isolate stable, uniformly monomeric species of FXIII^{o,low} on a reasonable time scale, this FXIII^A form was not included in catalytic comparisons.

To assess the catalytic differences between FXIII^A* and FXIII^{o,high}, we employed a spectrophotometric assay using K9 peptide (¹LGPGQSKVIG¹⁰) as a glutamine substrate and chromogenic DMPDA as second, amine substrate. Unlike the ammonia release assay, the transglutaminase reaction was now monitored directly at its second step, formation of a cross-linked product. An apparent K_m for K9 peptide was almost two-fold lower for FXIII^A*, compared to FXIII^{o,high}. This lower K_m indicated better interaction of the FXIII^A* enzymatic form with the glutamine substrate (Fig. 4A). On the other hand, FXIII^{o,high} demonstrated a slightly stronger V_{max} value suggesting that once saturated with the glutamine donor K9, FXIII^{o,high} provides faster substrate turnover than FXIII^A*. Thus, the observed individual kinetic parameters further corroborate proposed conformational/dynamic differences between FXIII^A* and FXIII^{o,high}.

We next conducted a series of SDS-PAGE assays to monitor FXIII^A-catalyzed crosslinking of a lysine mimic monodansylcadaverine (MDC) to a glutamine donor fibrinogen α C (233-425). The dansyl moiety of MDC provides a fluorescent tag on the protein and once the α C band is resolved on SDS-PAGE, the crosslinking reaction can be monitored by fluorescence of that band. α C represents a physiological FXIII^A protein substrate and contains three reactive glutamines [36]. While four lysine residues are present in the α C (233-425) sequence, no competing α C- α C conjugation, that would result in appearance of species of ≥ 40 kDa, was

detected even at high α C concentrations (Fig. 4B, panel iv). At 5 μ M α C concentration, the extent of MDC incorporation catalyzed by FXIII^{A*} was greater than by FXIII^{A^o,high} (Fig. 4B, panel i). Interestingly, increased monovalent ionic strength has been shown to promote higher FXIII^A activity [13, 15, 37, 38]. To rule out the possibility of an inhibitory effect of divalent ionic strength, the Ca^{2+} concentration in the crosslinking reaction was lowered from 100 mM down to 25 mM Ca^{2+} (Fig. 4B, panel ii). MDC crosslinking to α C by FXIII^{A^o,high} was essentially the same regardless of Ca^{2+} concentration in the crosslinking reaction mix, and in both cases, the FXIII^{A^o,high} activity was lower than that catalyzed by FXIII^{A*}. When α C concentration was raised to 40 μ M, the difference between crosslinking activity of FXIII^{A*} and FXIII^{A^o,high} became indistinguishable (Fig. 4B, panel iii). Overall, the results obtained with the short K9 peptide and a larger, more physiological α C substrate, suggest that FXIII^{A*} interacts with the glutamine substrate more readily than does FXIII^{A^o,high}.

DISCUSSION

Factor XIII activity has been reported in a variety of intra- and extracellular physiological environments and has been implicated in numerous pathological events. Being the final player of the coagulation cascade, FXIII^A has been recognized as a promising target for novel anticoagulant agents, with a prospect of fewer side effects [25, 39-41]. When implementing new coagulation management regimes, it may be desirable not to affect FXIII function in nonhemostatic events. On the other hand, those nonhemostatic functions may themselves present therapeutic interest. It is important to keep in mind that FXIII is activated either proteolytically or non-proteolytically depending on the physiological environment. To support future

pharmaceutical intervention approaches, critical new knowledge on these two FXIII activation strategies and their subsequent functional properties is needed.

Using AUC in previous work, we demonstrated dissociation of the FXIII A₂ dimer upon activation and that it is the monomeric FXIII^A that is catalytically competent [27]. In the current project, we investigated and compared dynamic and functional implications of proteolytic and nonproteolytic FXIII^A activation pathways. While thrombin-mediated proteolysis resulted in full dissociation of the A-subunits within minutes, activation by Ca²⁺ alone at physiological concentrations proceeded slower and took hours. Due to highly regulated Ca homeostasis, however, this activation pathway may operate constantly, thus providing constitutive FXIII activity within cells and ECM.

In contrast to FXIII^{A*} and FXIII^{A^o,low}, nonproteolytic activation in the presence of high (≥ 25 mM) Ca²⁺ did not result in aggregation of FXIII^{A^o,high} species. Increased Ca²⁺ concentration was shown before to stabilize active FXIII^A in solution [31], and in current AUC experiments, we correlated higher Ca²⁺ levels with reduced conformational heterogeneity of FXIII^{A^o,high}, as compared to FXIII^{A*}. The poor solubility of FXIII^{A^o,low} may suggest partial conformational similarity with FXIII^{A*}. However, the slow appearance of this form has limited its isolation and characterization in the current study.

In our enzymatic evaluations, FXIII^{A*} had an almost two fold lower K_m for the glutamine substrate than had FXIII^{A^o,high}, thus suggesting stronger interaction of FXIII^{A*} with that substrate. On the other hand, FXIII^{A^o,high} provided faster reaction turnover upon saturation with the glutamine donor. It is well known that the transglutaminase reaction proceeds through a mechanism involving addition of Q (glutamine) and K (amine or lysine) substrates. That is, the K-substrate enters the reaction after the enzyme forms a complex with the Q-substrate [42].

Therefore, this faster turnover by FXIII^{o,high} may be due to better interaction with the K-substrate and faster release of the crosslinked product.

In early work, Lewis with coworkers [43] reported that each A-subunit of plasma zymogen FXIII A₂B₂ binds a Ca²⁺ ion with 100 μM K_D. Hornyak et al. [44] later demonstrated that another Ca²⁺ ion is required by each A-subunit to dissociate from the B-subunits and to expose catalytic cysteines. Stieler with coworkers [40] recently crystallized FXIII^{o,high} with a covalently bound Q-mimic inhibitor, providing the first molecular structure of active FXIII^{o,high}. Besides the previously observed first, zymogenic Ca²⁺ site [45], the Stieler research group elegantly demonstrated the functional significance of two additional sites. In particular, development of the 2nd Ca²⁺ site brought about conformational changes in the FXIII^{o,high} molecule to initiate formation of a hydrophobic pocket for the K-substrate entrance. This site is the same as that presumed by Hornyak [44] to promote exposure of the reactive thiol on A-subunits. Coordination of the 3rd Ca²⁺ was proposed by Stieler et al. [40] to conclude formation of the hydrophobic pocket and to facilitate binding of the K-substrate.

Since the addition of Q and K is sequential and the crystal structure presented by Stieler et al. [40] depicts a covalent FXIII^{o,high}–Q-substrate intermediate rather than a free enzyme, we entertain an idea that the 3rd site for Ca²⁺ is filled upon binding of the Q-substrate. Whether binding of Ca²⁺ at the 3rd site precedes binding of the K-substrate or vice versa, is open to speculation. It seems plausible that at relatively low Ca²⁺ levels coordination of this 3rd site and completion of the hydrophobic pocket is dependent on the availability of the K-substrate. Such a possibility is consistent with the fact that in the absence of K-substrate, a water molecule acts as an acyl acceptor resulting in net deamidation of the glutamine to glutamate [46]. Intriguingly, the Ca²⁺ requirement for deamidation catalyzed by guinea pig liver transglutaminase was

demonstrated to be much lower than that for transamidation (crosslinking) [47], further supporting this possibility for a homologous FXIII_A.

In conditions where Ca concentrations are relatively high, such as in bone ECM, the 3rd FXIII_A Ca²⁺ site may already be saturated even in a free enzyme, prior to the Q-substrate binding. Our kinetic results suggest that this saturation facilitates a conformation (FXIII_A^{o,high}) that has a weaker affinity for the Q-substrate (indicated by higher K_m in the current study) but provides faster reaction turnover, than FXIII_A* (higher observed V_{max}). In addition to its stabilizing role observed in our AUC experiments, the FXIII_A^{o,high} conformation may also favor transamidation of the protein substrates over deamidation in acidic conditions of the bone ECM.

Thus far, the FXIII_A* form has been elusive from crystallographic efforts and also a challenging subject of solution experiments. The current studies effectively demonstrate that conformational differences do exist between FXIII_A* and FXIII_A^{o,high}. As mentioned before, the crystal structures of non-proteolytically activated FXIII_A^{o,high} in complexes with inhibitors are available from the Stieler group (PDB 4KTY, 5MHM, 5MHN, 5MHO). It would thus be feasible to screen for pharmaceutical candidates that confer specificity to FXIII_A^{o,high} and eliminate those affecting the FXIII_A*. Should this orthosteric design fail, a search for allosteric effectors may be more successful [41]. Previous studies in our group involving chemical modification and hydrogen-deuterium exchange suggested better solvent accessibility of the β-sandwich region of FXIII_A*, while β-barrel 1 was more solvent-exposed in FXIII_A^{o,high} [48, 49]. These local structural differences may serve as a starting point for allosteric targeting of FXIII_A.

Prior studies have shown that increasing levels of FXIII_A* affect the rate and extent of fibrin crosslinking [12, 33, 50]. Taking this property into consideration, the fibrin clot network could be further manipulated by regulating proteolytic activation of FXIII_A. In the current work,

we showed that the FXIII V34X variants (L34, F34, and W34) influenced such FXIII^{A*} generation rates and in response affected fibrin crosslinking. Faster proteolytic activation (and hence, higher amount of FXIII^{A*}) during blood coagulation resulted in a greater extent of fibrin crosslinking and incorporation of other proteins into the clot (FXIII L34>V34,F34>W34). The current results effectively complement FXIII activation peptide models examined by Duval and coworkers. Their studies **focused on A34, L34, M34** along with surrounding FXIII AP residues [32]. Our focus has been on fast activating L34 versus aromatic V34X substitutions that are well tolerated by the anticoagulant thrombin W215A species [30]. Thus, therapeutic strategies to specifically limit thrombin's ability to activate FXIII^A and/or fibrinogen, may be promising. The tunable FXIII-based ³⁴XVPR↓G³⁸ amino acid sequence might also be employed as a prodrug linker. Thrombin-dependent cleavage of the linker sequence would release the pharmaceutical agent at a desirable rate and specifically at the clotting site. This approach would also ensure that neither FXIII^{A^{o,low}} nor FXIII^{A^{o,high}} are affected.

Overall, the presented research expands the understanding of the functional outcomes of FXIII activation pathways. Based on our previous and current results, we offer the following model: nonproteolytic FXIII^A activation proceeds relatively slow, but constantly. High mM Ca²⁺ levels, such as in bone ECM, stabilize the FXIII^{A^{o,high}} form and provide its constitutive activity as part of the tissue remodeling processes. By contrast, thrombin-mediated AP-removal facilitates binding of Ca²⁺ at its low level in plasma, thus promoting fast buildup of FXIII crosslinking activity. The resultant FXIII^{A*} species have higher affinity towards glutamine substrates and respond even to low concentrations of these substrates, ensuring clot-stabilizing function during blood coagulation. At the same time, the tendency of FXIII^{A*} to aggregation may determine its relatively short life time in plasma. Different activation conditions also result

in conformationally different FXIIIa forms, thus providing a possible basis for developing therapeutic strategies for compartment-specific management of FXIIIa function.

EXPERIMENTAL PROCEDURES

Materials – Recombinant yeast expressed FXIIIa was a gift from the late Dr. Paul Bishop (Zymogenetics, Seattle, WA, USA). Recombinant human thrombin was generously provided by Dr. Enrico Di Cera and Dr. Leslie Pelc (St. Louis University, MO, USA). Bovine thrombin was purchased from Sigma-Aldrich (USA). Human FXIII-free peak 1 fibrinogen was purchased from Enzyme Research Laboratories (South Bend, IN, USA). Murine plasma was obtained from mice lacking the FXIII catalytic A subunit (FXIII A^{-/-}) [26, 51]. **Such FXIII-deficient mice were engineered to be missing the FXIII A subunits but still express the FXIII B subunits.** These mice were maintained at the Cincinnati Children’s Hospital Medical Center animal care facility. All studies involving the use of animals were approved by the Cincinnati Children’s Hospital Medical Center Institutional Animal Care and Use Committee. Cincinnati Children’s Hospital is an AAALAC accredited institution. Thrombin inhibitor D-phenylalanyl-prolyl-arginyl chloromethyl ketone (PPACK) was purchased from Haematologic Technologies (Essex Junction, VT, USA). A transglutaminase glutamine substrate peptide K9 (¹LGPGQSKVIG¹⁰) was synthesized by New England Peptide (Gardner, MA, USA). All other reagents were of the highest purity available.

AUC studies of FXIIIa – Molar concentration in this work refers to A-subunits as opposed to A₂-dimers of FXIIIa. 2 μM zymogen FXIIIa was incubated in borate buffer (20 mM boric acid, pH 7.8, 150 mM NaCl) in the presence of 25 – 100 mM CaCl₂ for 30 min – 6 h at 37 °C. FXIIIa was also incubated in the borate buffer at a lower 4 mM CaCl₂ for 30 min – 96 h. To achieve better solubility of FXIIIa under these conditions, the 4 mM Ca²⁺-samples were supplemented with 5% DMSO. For proteolytic activation, 2 μM FXIIIa was incubated in the borate buffer with bovine thrombin (3.5 NIH units/ml) and 4 mM CaCl₂ for 30 min at 37 °C, followed by inhibition of thrombin with 760 nM PPACK. In addition to 4 mM CaCl₂, some samples of the proteolytically activated FXIIIa were supplemented with 100 mM MgCl₂.

A detailed description of AUC experimental parameters can be found in [27]. Briefly, samples of activated FXIIIa were subjected to sedimentation velocity AUC in an Optima XL-A ultracentrifuge (Beckman Coulter) at 20 °C and 50,000 rpm. Data were analyzed using the program SEDFIT (www.analyticalultracentrifugation.com), and experimental sedimentation coefficients were corrected based on the measured density and viscosity of the buffer, thus allowing direct comparison of results obtained with different experimental conditions. Analytical runs were performed with two independent samples for each condition examined. **Control studies repeated in the current project yielded Sedimentation Coefficient values that were highly comparable.**

Expression and purification of FXIIIa V34X variants and Fibrinogen αC (233-425) – pGEX plasmid vectors encoding GST-tagged FXIIIa V34 and GST-tagged fibrinogen αC (233-425) were employed [52]. Single amino acid V34X substitutions (L, F, or W) were introduced into the GST-FXIIIa plasmid using the QuikChange II Mutagenesis kit (Agilent Technologies).

The expression vectors were transformed into *E. coli* BL21 (DE3) Gold cells and proteins were expressed using auto-inducing media [53]. Using an AKTA Prime FPLC, GST-tagged proteins were purified by affinity chromatography on a GST-trap column (GE Healthcare, USA) system. Target proteins were cleaved from the GST-tag by in-column digestion with PreScission protease and eluted with tris buffered saline (TBS, Tris Acetate 50 mM, pH 7.4, NaCl 150 mM). The purity was assessed using SDS-PAGE, and molar concentration was determined from absorbance readings at 280 nm using $\epsilon = 125710 \text{ M}^{-1}\text{cm}^{-1}$ for V34, F34, L34 FXIIIa variants and $131210 \text{ M}^{-1}\text{cm}^{-1}$ for W34 FXIIIa. An $\epsilon = 41480 \text{ M}^{-1}\text{cm}^{-1}$ was applied for fibrinogen αC (233-425). These extinction coefficients were calculated using ProtParam tool (www.expasy.org).

Fibrin crosslinking in the presence of FXIIIa V34X variants – Human FXIII-free fibrinogen (1 mg/ml), a recombinant FXIII AP variant (V34, L34, F34, or W34, 50 nM), and recombinant human thrombin (12 nM) were incubated in a 20 μL TBS containing 2.5 mM CaCl_2 at 37 °C. At each time point (5 – 60 min), the reaction was quenched by addition of 10 μL of reducing sample loading buffer followed by boiling for 5 min, resulting in full solubilization of formed clots. Samples were then subjected to SDS-PAGE on an 8% gel. These fibrin crosslinking reactions were performed three independent times, with essentially the same results.

SEM analysis of plasma clots formed in the presence of FXIIIa V34X variants – A recombinant FXIII AP variant (L34, V34, or W34, 100 nM final concentration) was administered into citrated plasma from FXIIIa-deficient mice. Our FXIII concentration (100 nM) is within the physiological range of 86-173 nM [54]. After 5 min incubation at 37 °C, clotting was initiated by addition of bovine thrombin (to a final concentration of 2.1 NIH units/ml) and CaCl_2 (final

13.5 mM, to exceed the citrate in plasma) in the borate buffer, resulting in a 1:4 dilution of the stock plasma. Clots were incubated for 2 h at 37 °C in closed microfuge tubes, rinsed with deionized H₂O, and dehydrated in the presence of increasing (10 – 100%) concentrations of ethanol and dried with hexamethyldisilazane. The samples were next sputter-coated with Au-Pd and imaged using a Zeiss Supra 35 electron microscope.

Since we aimed to study FXIIIa-mediated crosslinking of the fibrin clots, no glutaraldehyde (a chemical crosslinking agent) was used to stabilize the samples. Undiluted plasma clots suffered great deformation during drying and were too dense. A 1:4 dilution of the plasma (and hence, endogenous fibrinogen) resulted in thinner, more suitable SEM samples that allowed for adequate comparisons between exogenous FXIIIa variants. These experiments were performed in duplicate.

MDC crosslinking SDS-PAGE based activity assay – In order to compare activities of proteolytically and nonproteolytically activated FXIIIa, a series of MDC crosslinking assays was performed. For proteolytic activation, 1 μM FXIIIa was incubated in TBS with 30 nM recombinant human thrombin and 4 mM CaCl₂ for 30 min at 37 °C, followed by addition of 760 nM thrombin inhibitor PPACK. Nonproteolytic activation was achieved by 30 min incubation of 1 μM FXIIIa in TBS in the presence of 100 mM CaCl₂.

Recombinantly expressed fibrinogen αC (233-425) was used as a glutamine donor (Q-substrate). αC (5 – 40 μM final) was preincubated for 5 min at 37 °C with a lysine mimic MDC (K-substrate, 1 mM final, from a 20 mM stock solution in methanol) in TBS containing CaCl₂ (4 – 100 mM final concentration). The crosslinking was initiated by addition of activated FXIIIa (final concentration 100 nM). Reaction aliquots were taken at 1 – 7 min and quenched by

addition of reducing sample loading buffer, followed by 3 min of boiling. Samples were resolved using SDS-PAGE on 15% gels. Prior to Coomassie blue staining, the gels were photographed under UV light. The different MDC assay series were performed three independent times with comparable results obtained each time.

Continuous spectrophotometric kinetic assay – In this assay adapted from de Macedo et al. [55], FXIIIa incorporated chromogenic K-substrate N,N-dimethyl-1,4-phenylenediamine (DMPDA) into a Q-substrate K9-peptide. The reaction progress was monitored by an increase in absorbance at 278 nm resulting from an anilide functionality of the crosslinked product. Nonproteolytic and proteolytic activation were performed as described for the MDC assay, except that FXIIIa was 4 μM in the activation mix. K9-peptide (46 – 1386 μM final) was preincubated with DMPDA (700 μM final, from a 100 mM stock solution in methanol) in the borate buffer with 4 or 100 mM of CaCl_2 for 5 min at 37 °C. The reaction was initiated by addition of activated FXIIIa (final concentration 1 μM). Reaction velocities were determined over the initial part of the absorbance curve. An $\epsilon = 8940 \text{ M}^{-1} \text{ cm}^{-1}$ [55] was applied to convert absorbance to μM of crosslinked product. Less than 15% of both Q and K substrates reacted in the initial linear region of the absorbance curves. Velocities in $\mu\text{M min}^{-1}$ were fitted to the Michaelis-Menten equation as a function of the K9 concentrations using Kaleidagraph software (Synergy). The resultant K_m and V_{max} parameters represent apparent values and serve for comparison of nonproteolytically and proteolytically activated FXIIIa. The kinetic assays were performed in triplicate (three independent trials) for each activation condition. Results are presented as mean \pm SD (N = 3).

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Author contributions. BA and MM designed the research. HP, KS, and RA constructed the expression vectors for GST tagged FXIIIa V34 and Fbg α C (233-425) and also contributed to research planning. MF supplied the FXIIIa-deficient murine plasma (FXIII A^{-/-}) and guidance on its use. BA and WD carried out the experiments. BA and MM wrote the manuscript. All authors reviewed the data, and both critiqued and approved the manuscript.

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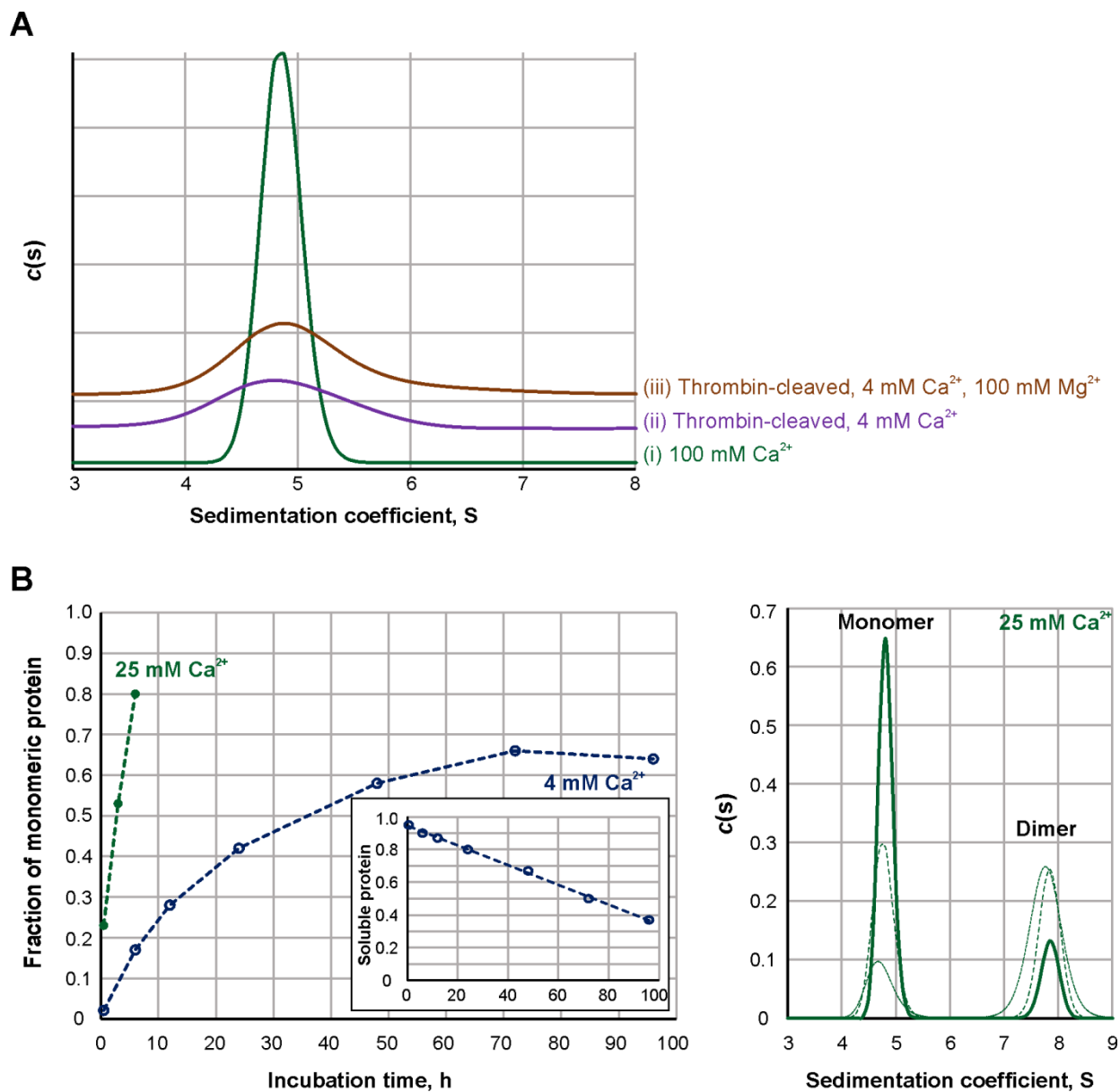


Fig. 1. Sedimentation properties and activation rate of FXIIIa under different conditions. A – Sedimentation profiles of FXIIIa studied by AUC: 2 μM FXIIIa was activated at 37 $^{\circ}\text{C}$ for 30 min nonproteolytically by 100 mM CaCl_2 (trace I, green) or proteolytically by 3.5 NIH units/ml bovine thrombin in the presence of 4 mM CaCl_2 (trace ii, purple). An additional sample of thrombin-activated FXIIIa contained 4 mM CaCl_2 and 100 mM MgCl_2 (trace iii, orange). Two independent samples were analyzed for each condition, with the same results.

B – Dissociation progress of 2 μM FXIIIa in the presence of 4 (open blue circles) and 25 mM CaCl_2 (filled green circles). Graph on the left presents quantitative analysis of sedimentation velocity AUC data. The insert demonstrates fraction of soluble protein (estimated from absorbance at 280 nm) in samples of 4 mM Ca^{2+} -activated FXIIIa as a function of time. The panel on the right depicts AUC sedimentation profiles for the FXIIIa samples incubated in the presence of 25 mM CaCl_2 for 30 min (dotted green line), 3 h (dashed green line) and 6 h (solid green line).

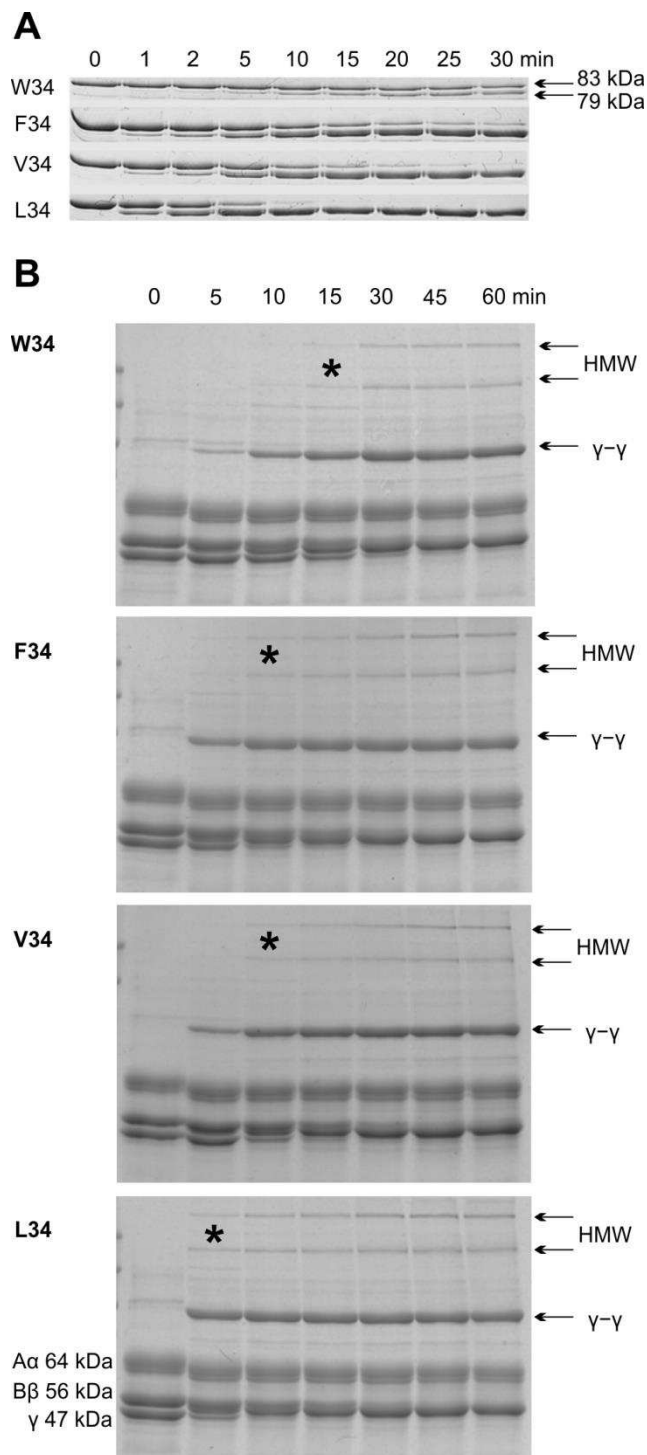


Fig. 2. Effect of the rate of FXIII A proteolytic activation on fibrin crosslinking.

A – 1 μ M recombinant FXIII A AP variants were incubated with 30 nM recombinant human thrombin at 37 °C. Aliquots were withdrawn at denoted time points and thrombin was inhibited with 760 nM PPACK. The samples were then subjected to SDS-PAGE on 8% gels. FXIII A AP cleavage resulting in a 4 kDa molecular weight loss could then be followed by monitoring the appearance of a 79 kDa band.

B – Fibrin crosslinking by FXIII A AP variants. 1 mg/ml human FXIII-free fibrinogen was combined with a recombinant FXIII A AP-variant (50 nM) at 37 °C, and crosslinking was initiated by addition of 12 nM recombinant human thrombin and 2.5 mM CaCl₂. At each denoted time point, the reaction was stopped by addition of reducing sample loading buffer and boiling. The samples were resolved via SDS-PAGE (8% gel). Fibrinogen chains A α , B β , γ , HMW (high molecular weight crosslinks), and γ - γ crosslinks are annotated. * symbol denotes the earliest detection of HMW species during the course of the crosslinking reaction.

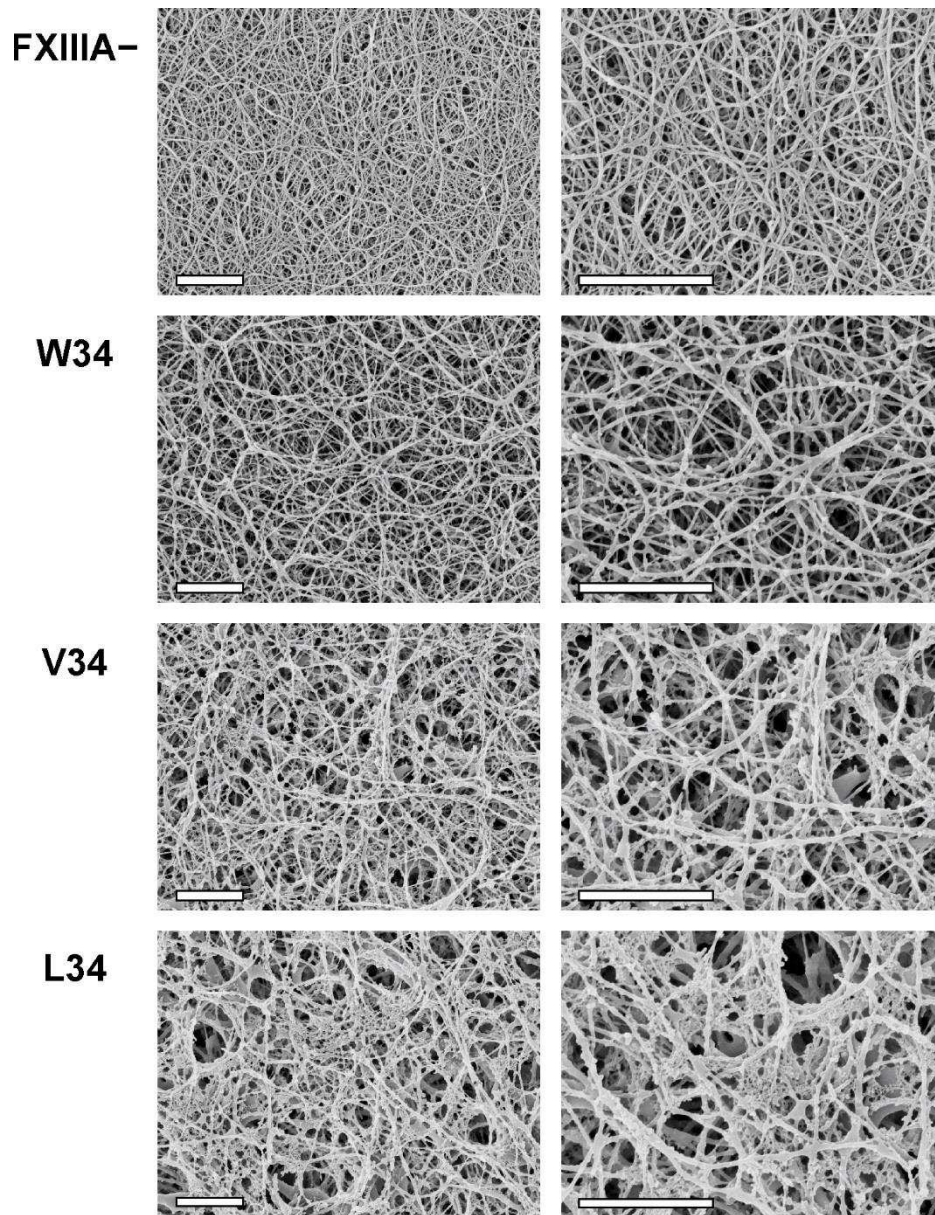


Fig. 3. Scanning electron microscopy of fibrin clots in the presence of FXIII A P variants.

100 nM FXIII A P variants were combined with plasma from FXIII A-deficient mice (final dilution of plasma was 1:4). Control samples were made without FXIII A (FXIII A⁻). Clotting was initiated by addition of 2.1 NIH units/ml bovine thrombin and 13.5 mM CaCl₂. Clots were formed for 2 h at 37 °C and prepared for SEM as described in Materials and Methods. For each FXIII A P variant, two clots were studied, with essentially the same results. Each set of clots (FXIII A⁻, W34, V34, and L34) was formed using plasma obtained from the same mouse. Shown are representative SEM photographs at two magnifications: 10,000x (left) and 20,000x (right), scale bars are 2 μm. Note that during the SEM sample preparation process, the fixative agent glutaraldehyde was not included to avoid artificial protein crosslinking. As a result, the FXIII A⁻ clots were significantly flattened during the dehydration process. In response, the resultant FXIII A⁻ fibrin fibers on the SEM images appeared denser packed.

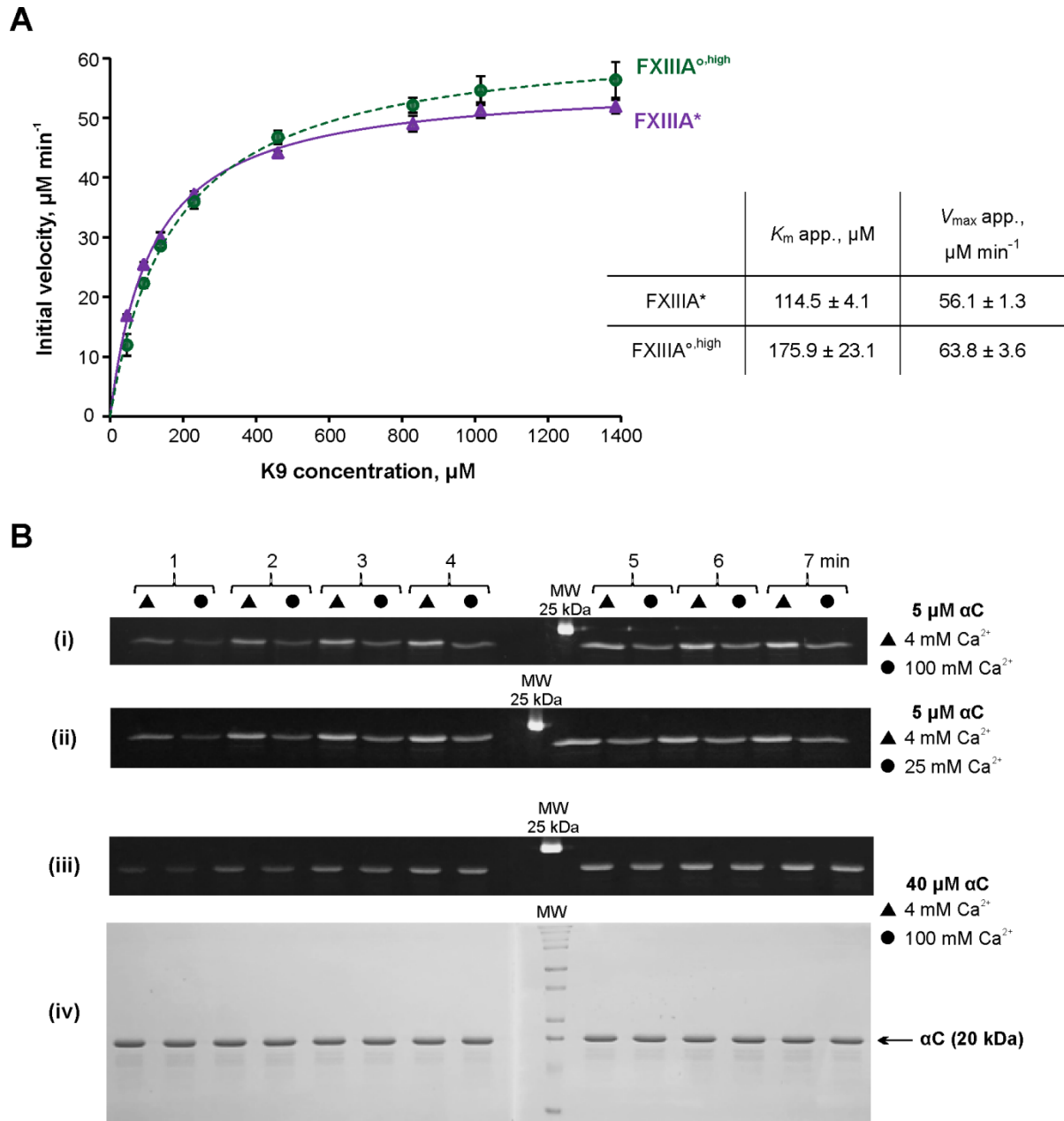


Fig. 4. Transglutaminase activity of FXIII A under different conditions.

A – Spectrophotometric kinetic assay. Glutamine donor K9 peptide was preincubated with the lysine mimic DMPDA at 37 °C for 5 min, and the crosslinking reaction was initiated by addition of 1 μM FXIII A^{o,high} (green circle) or FXIII A* (purple circle). The reaction was followed by an increase in absorbance at 278 nm due to formation of the anilide crosslinked product. The plot represents Michaelis-Menten fits of the initial reaction velocities as function of the K9 concentration. Apparent K_m and V_{max} resulting from the fits are shown in the table insert. Values are presented as mean \pm SD (N=3).

B – MDC crosslinking SDS-PAGE based assay. Recombinant fibrinogen αC (233-425) was preincubated with 1 mM MDC at 37 °C for 5 min. The crosslinking reaction was initiated by addition of 100 nM FXIII A^{o,high} or FXIII A*. Reaction aliquots were withdrawn at 1 – 7 min and quenched by addition of reducing sample buffer and boiling. The time point samples were loaded on 15% SDS-PAGE side by side for FXIII A* (triangles) and FXIII A^{o,high} (circles). Two gels were run (1 – 4 and 5 – 7 min time points). The gels were aligned and photographed under UV light (panels i – iii). Panel iv – Coomassie Blue-

stained gel pair from panel iii demonstrating absence of α C- α C conjugation. FXIII^{A*} was always preactivated in the presence of 4 mM CaCl₂, and FXIII^{A^ohigh} – in the presence of 100 mM CaCl₂. Concentrations of α C and CaCl₂ in the crosslinking reaction mix are annotated on the right.