The Disulfide Bond between Cys22 and Cys27 in the Protease Domain Modulate Clotting Activity of Coagulation Factor X

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Abstract

Keywords

- coagulation factor X
- ► Cys27
- disulfide bond
- oxidation/reduction balance
- molecular dynamic simulations

Introduction

Coagulation factor X (FX) is the conjunction of intrinsic and extrinsic coagulation pathway. It activates pro-thrombin into

received October 10, 2018 accepted after revision January 30, 2019 thrombin and is a key player in blood coagulation.¹ Decreased or elevated FX shifts the balance of haemostasis of blood and is associated with bleeding or thrombotic disorders. FX deficiency is a rare hereditary bleeding disorder, affecting 1 out of 1,000,000 general population, and is caused by genetic mutations of *F10* gene on chromosome $13.^{2,3}$ High-level FX in plasma, however, is a risk factor for venous thrombosis,

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The Cys22-Cys27 disulfide bond of factor X (FX) protease domain is not conserved among coagulation factors and its contribution to the physiological haemostasis and implication in the pathogenesis of haemostatic and thrombotic disorders remain to be elucidated. Mutation p.Cys27Ser was identified in a pedigree of congenital FX deficiency and fluorescence labelling study of transiently transfected HEK293 cells showed accumulation of FX p.Cys27Ser within cell, indicating incompetent secretion partially responsible for the FX deficiency. The clotting activity of FX p.Cys27Ser was decreased to about 90% of wild-type, while amidolytic and pro-thrombinase activities (kcat/Km) determined with recombinant FXa mutant were 1.33- and 4.77-fold lower. Molecular dynamic simulations revealed no major change in global structure between FXa p.Cys27Ser and wild-type FXa; however, without the Cys22-Cys27 disulfide bond, the insertion of newly formed N terminal of catalytic domain after the activation cleavage is hindered, perturbing the conformation transition from zymogen to

enzyme. The crystal structure of FXa shows that this disulfide bond is solvent

accessible, indicating that its stability might be subject to the oxidation/reduction balance. As demonstrated with FX p.Cys27Ser here, Cys22-Cys27 disulfide bond may

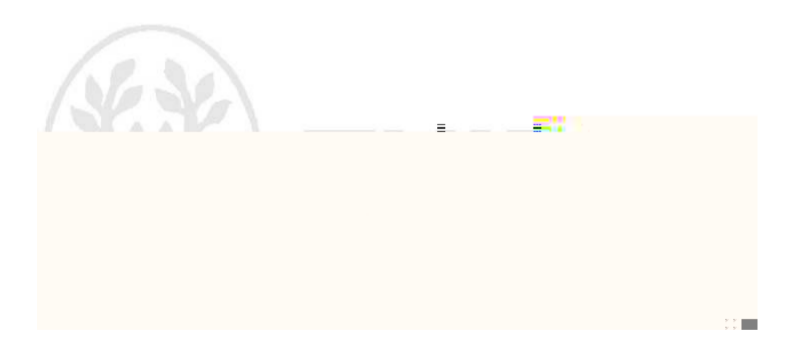
modulate FX clotting activity, with reduced FX pertaining less pro-coagulant activity.

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on a 10% polyacrylamide gel under non-reducing conditions to examine the homogeneity and purity of the isolated wild-type (WT) and mutant FX derivatives. The concentration of purified recombinant FX protein was determined by ELISA as described previously. The clotting activity of recombinant FX and its variant were determined using aPTT- and PT-based coagulation function tests following similar protocol mentioned above.

Determination of the FX Activation by the FVIIa–Tissue Factor Complex

The initial rate of activation of both recombinant wide-type and mutant FX (0–400 nM) by FVIIa (0.1 nM) in complex with tissue factor (TF) (2 nM) was monitored on phosphatidylcholine/phosphatidylserine (PC/PS) vesicles in TBS/Ca²⁺ for 3.5 minutes at room temperature in 30 μ L reactions as described.⁶ Activation reactions were terminated by addition of 30 μ L of ethylenediaminetetraacetic acid (EDTA) and subjected to amidolytic assay using S2765 as substrates. The concentration of FXa generated was determined by comparing its S2765 hydrolytic activity against standard curves prepared by serial diluted FXa (0–200 nM).

The Kinetic Analysis of FXa Amidolytic Activity

The amidolytic activity of FXa was characterized with steady-state kinetics analysis of hydrolysis of chromogenic substrates S2765. The FXa derivatives (2 nM) was incubated with various concentration of S2765 (25–1,600 μ M) in TBS/ Ca²⁺ buffer (0.02 M Tris-HCl, 0.1 M NaCl, 0.1 mg/mL bovine serum albumin, 0.1% polyethylene glycol 8000 and 5 mM Ca²⁺, pH 7.4) at room temperature for 15 minutes on a microplate reader (Molecular Devices, Menlo Park, California, United States). The rate of hydrolysis was recorded at 405 nm and the Km and kcat values were calculated with the Michaelis–Menten equation and the specificity of WT and mutant FX was determined using the ratio of kcat/Km.

Determination of the Activation of Pro-Thrombin by FXa The concentration dependence of pro-thrombin activation by FXa derivatives was determined in the absence and presence of FVa on PC/PS vesicles as described.⁷ Briefly, in the absence of FVa, pro-thrombin $(0-2 \mu M)$ was incubated with each FXa (10 nM) in TBS/Ca²⁺ on PC/PS vesicles (25 μ M) at room temperature for 30 to 90 minutes. EDTA was added to a final concentration of 20 mM and the initial rate of prothrombin activation was monitored from the rate of thrombin generation detected by an amidolytic activity assay using 450 µM S2238. The standard curve was prepared from the cleavage rate of S2238 by known concentrations of recombinant thrombin under the same conditions. In the presence of human FVa (20 nM), increasing concentration of prothrombin (30-2,000 nM) was incubated with each FXa derivative (0.1 nM) in TBS/Ca²⁺ on PC/PS vesicles (25 μ M) for 1.5 minutes at room temperature. After the termination

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GROMACS standard analysis tools and visual MD.¹³ Considering possible rotation of the carboxyl group of Asp194, here the distance between Ile16 and Asp194 was simply defined as the atomic distance between Ile16N_{ζ} and Asp194C_{γ}.

Results

Patient Data

Although the plasma FX:Ag of the proband only mildly decreased (69.7% of normal control), the clotting time was prolonged in both the aPTT and PT tests. The FX:C determined by all three means, including aPTT-, PT- and RVV-based methods, was consistently around 20% of normal of control. However, the amidolytic activity as determined by chromogenic assay using S2765 was only mildly compromised (around 68.3% of normal control). The coagulation function analysis of other family members showed that both aPTT and PT were within normal reference range (-Table 1), but all of them had their FX:C partially decreased. The FX:C of proband's father (I-1) was around 75% of normal, which was consistent

with his antigen level as well as chromogenic activity; the patient's mother (I-2) and son (III-1) had almost normal FX:Ag and chromogenic activity; however, their FX:C decreased to lower borderline of the reference range (\sim 50% of normal control). Genetic analysis of proband's F10 gene revealed two single nucleotide changes, c.736t > a and c.956a > g, which predicted missense mutations p.Cys27Ser and p. Tyr99Cys (Fig. 1B). The proband inherited p.Tyr99Cys from her mother (I-2) and had it passed onto her son (III-1). The discrepancy of FX:Ag and FX:C of both p.Tyr99Cys mutation carrier suggested that the mutant was a cross-reactive material positive mutation (CRM +) and the deficiency was mainly caused by impaired function of FX mutant. The other mutation p.Cys27Ser was traced back to proband's father (I-1) and its impact on function and expression of FX warranted further study to characterize (> Table 1).

Coagulation Activity of Recombinant FX Mutant The recombinant WT FX and FX p.Cys27Ser mutant showed similar migrant pattern on Western blot (Fig. 2A). The

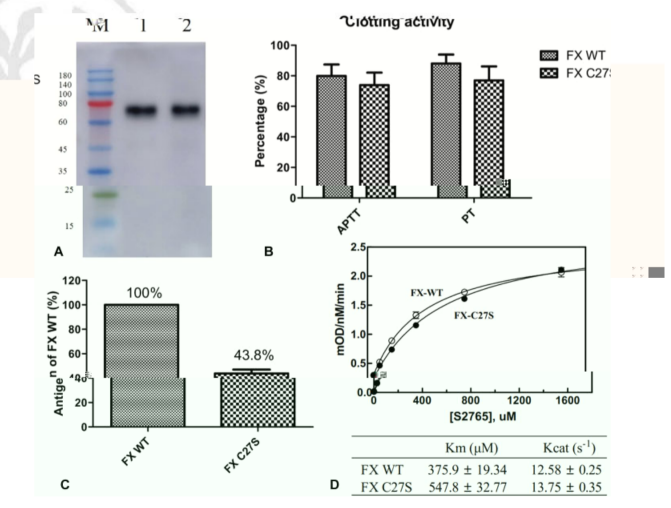


Fig. 2 In vitro expression and characterization of factor X (FX) mutant. (A) The migration pattern of recombinant FX mutant p.Cys27Ser is similar

clotting activity of FX mutant p.Cys27Ser determined by PTand aPTT-based clotting assays mildly decreased to around 90% of WT FX (Fig. 2B). The FX mutant p.Cys27Ser collected from supernatant of conditioned medium was much less than WT (43.8%), suggesting possible impediment of synthesis or secretion of the mutant (Fig. 2C).

Kinetic Analysis of FX p.Cys27Ser Cleavage of Chromogenic Substrate

The amino acid replacement increased the Km from 375 to 547 uM, but had little effect on the kcat (12.58 vs. 13.75 s⁻¹) (\succ Fig. 2D), suggesting the cleavage of chromogenic substrate S2765 by FX p.Cys27Ser was only marginally affected (1.33-fold change of kcat/Km).

FX Activation by FVIIa-TF Complex

The activation of FX by its physiological activator FVIIa–TF complex was slowed by the amino acid substitution p. Cys27Ser, with a 1.57-fold higher Km (from 90.44 to 142.4

nM) and 2.17-fold lower kcat (from 325.6 to 149.9 min⁻¹) (\succ Fig. 3A).

Activation of Pro-Thrombin by FXa and Pro-Thrombinase Complex

Similar to the amidolytic assay, the pro-thrombin activation activity by FX p.Cys27Ser mutant was impaired significantly, with a much smaller kcat (1.39 vs. 10.83 min⁻¹ of WT FXa) and greater Km (1031.0 vs. 365.2 nM of WT) (\succ Fig. 3C). Interestingly, the difference between WT and p.Cys27Ser mutant became minimal when FVa was present. The kcat of WT FXa and mutant was both increased dramatically (1,139 vs. 1,221 min⁻¹), and Km deceased to 139.5 and 713.6 nM, yielding 4.77-fold change of kcat/Km between WT FXa and mutant (8.16 vs. 1.71) (\succ Fig. 3D; \leftarrow Table 2).

Binding Af**fi**nity of FXa Mutant p.Cys27Ser and FVa The binding between FVa and FX was not affected significantly by the mutation p.Cys27Ser. The Kd determined was

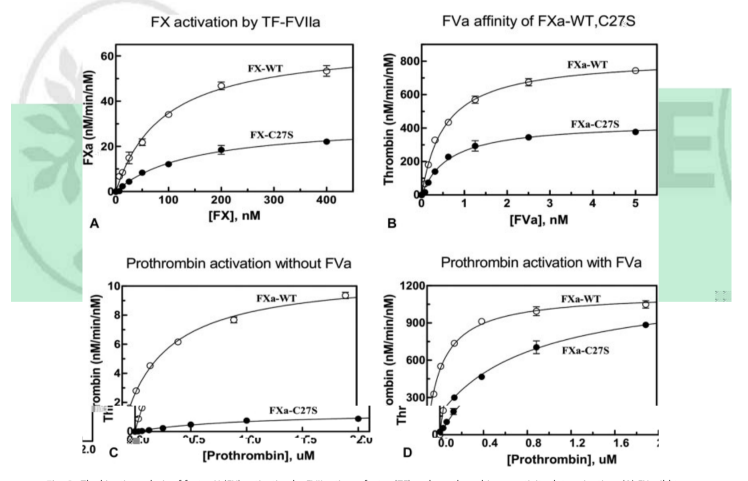


Fig. 3 The kinetic analysis of factor X (FX) activation by FVIIa-tissue factor (TF) and pro-thrombinase activity determination. (A) FX-wild-type (WT) (\circ) or FX-C27S (•) were activated by FVIIa (0.1 nM) in complex with TF (2 nM) on phosphatidylcholine/phosphatidylserine (PC/PS) vesicles in Tris-buffered saline (TBS)/Ca²⁺ for 3.5 minutes at room temperature. The reactions were terminated by 20 mM ethylenediaminetetraacetic acid (EDTA) and the FXa generation was determined from standard curves as described in the 'Materials and Method' section. (B) The activation of pro-thrombin (1 μ M) by FXa-WT (\circ) and FXa-C27S (•) (50 pM) was monitored in the presence of different concentrations of FVa (0–5 nM) on PC/PS vesicles (25 μ M) in TBS/Ca²⁺ for 30 seconds. Note that 20 mM EDTA was added to terminate reaction and the rate of thrombin generation was measured as described in the 'Materials and Method' section. (C) In the absence of FVa, various concentrations of pro-thrombin (0~2 μ M) were incubated with 10 nM of either FXa-WT (\circ) or FXa-C27S (•) on PC/PS vesicles in TBS/Ca²⁺. Following 30 to 90 minutes' incubation, EDTA was added to a final concentration of 20 mM and the initial rate of thrombin generation was measured from the cleavage rate of S2238 as described in the 'Materials and Method' section. (D) It is the sam1 T m 4 (n)-4.2/F117.3 (e) s-280.1F1 1 T f 4.2256 0 8T D 75 T c (C) T j /F3 1 T f 0.6274 0

0.545 nM for WT FXa and 0.652 nM for FXa p.Cys27Ser mutant (Fig. 3B).

Immuno**fl**uorescence Staining of FX Derivatives in Transient Transfected Cells

The immunofluorescence staining of FX (~Fig. 4A and C) or FX mutant (~Fig. 4B and D) intra-transiently transfected HEK-293 cell showed dramatically increased retention of FX molecule. Most of the recombinantly expressed FX Cys27Ser accumulated within the ER system (~Fig. 4B), and to a less extent, in the Golgi body (~Fig. 4D), which might be responsible for inefficient secretion of FX Cys27Ser mutant as observed previously. The fluorescence intensity of FX Cys27-Ser mutant was 1.5-fold higher than WT (~Fig. 4E).

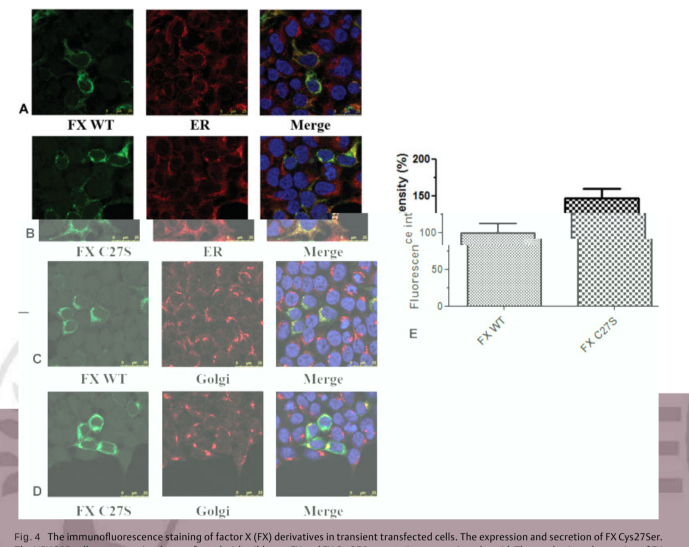
Molecular Dynamics Simulations of Cleaved FXa Models in the Activated Conformation

Based on the X-ray structure of the activated FXa (PDB ID: 2W26), molecular models were constructed for the WT, Cys27Ser mutant (MT) and a WT FXa with the Cys22-Cys27 DISB WT, as shown in \blacktriangleright Supplementary Table S2 (available in the online version). Starting from the initial models, all systems were simulated for more than 300 ns. After about 60 ns simulations, all the systems tended to be equilibrated. In this activated conformation, all the three systems have the N-terminal inserted into the activation pocket all the time, with lle16 relatively fixed to Asp194 by a salt bridge (\vdash Fig. 5). Comparison between the three systems

showed almost no difference in the fluctuations at the Nterminal segment (Fig. 5A), variations in the distance between Ile16 and Asp194 (Fig. 5B) or the local positions of the critical residues Ile16, Asp194 and Ser195 in the activation pocket and the catalytic pocket (Fig. 6). The hydrogen bonds network around the activation pocket and the backbone hydrogen bonds between Gln20 and Met156 to form an untypical β sheet are also observed. These results suggested that the structure of the catalytic site of the activated FXa might not be changed importantly in the Cys27Ser mutant. Also, considering that the relative position of the mutation site is far from the catalytic site, it is highly possible that the amino acid substitution does not change the amidolytic activity of FXa directly. Instead, it might hinder the early phase of conformation switch from inactive zymogen (FX) to enzyme with hydrolytic activity (FXa) after the scissile bond between heavy chain and light chain is cleaved, that is, the insertion of the N-terminal from the just cleaved position into the activation pocket.

Simulations of the Insertion of the N-Terminal into the Activation Pocket

In order to construct the models prior to the insertion, the Nterminal segments (residues 16–22) of the systems above were remodelled by Modeller in a possible extended conformation after the cleavage activation. In simulations of these systems, the N-terminal of the cleavage site was initially swinging around far from the activation pocket. Possibly because of



The HEK 293 cells were transiently transfected with wild-type FX and FX Cys27Ser mutation expressing plasmid. The synthesis and secretion of FX were traced with fluorescence-labelled FX-specific antibodies (red), the cell nucleus were stained with 4',6-diamidino-2-phenylindole (DAPI) (blue). The Golgi body and ER were also stained (red) to locate FX. (A) FX wild-type (WT) (green) + ER (red) + Merged; (B) FX Cys27Ser + ER + Merged; (C) FX WT + Golgi + Merged; and (D) FX Cys27Ser + Golgi + Merged. (E) The accumulation of synthesized FX and FX Cys27Ser mutant within cells was estimated by fluorescence intensity measurement.

the tethering of the Cys22-Cys27 disulfide bond, the swing in the WT system was relatively limited and the N-terminal segment was quickly anchored to the edge of the activation pocket, formed two strong backbone hydrogen bonds between Gln20 and Met157, which somewhat integrated the N-terminal segment into the core β sheet. This anchoring greatly further limited the fluctuation of the N-terminal segment. Although the end (Ile16 to Gly19) still wandered at the entry of the pocket for a period, it gradually entered into the activation pocket and formed the Ile16-Asp194 salt bridge spontaneously after 150 ns. On the other hand, the mutant and the WT systems with Cys22-Cys27 DISB both had the N-terminal segment wander around far away outside the pocket in the 300-ns simulations, meaning the insertion is importantly delayed (Fig. 7). Correspondingly, RMSF analyses showed more and more fluctuations in the length of the N-terminal segment and in the extent of fluctuations in the WT, DISB WT and MT systems, respectively (Fig. 5A). This difference is also obvious in comparison of the Ile16-Asp194 distances in

► Fig. 5B. After a short time of equilibration, the WT enzyme (in cyan) had the distance gradually shortened from 2 to 0.3 nm, which is consistent to a salt bridge; the DISB WT (in magenta) stayed nearly 2 nm; the mutant (in orange) stayed at 2 nm for about 200 ns and, after that, the N-terminal moved away back to about 3 nm. And clustering analyses also found more concentrated distribution in the WT system around the inserted conformation versus much more scattered distribution in the mutant (► Supplementary Figs. S2 and S3, available in the online version).

Discussion

The coagulation FX is a serine protease zymogen, belonging to the trypsin family just like other components of the coagulation cascade. Evolved by gene duplications from the same ancestry, FX has high homology with other coagulation factors, such as FIX, FVII, FXI and pro-thrombin.¹⁴ The strictly conserved residues shared among the coagulation factors are

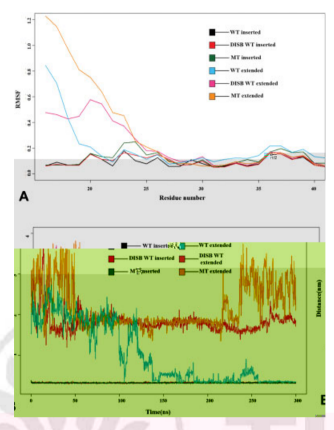


Fig. 5 Fluctuations of the N-terminal segments. (A) The root-meansquare fluctuations (RMSFs) of the N-terminal residues 16 to 51, showing the fluctuations of their backbone atoms. (B) Distances between the side chains of Ile16 to Asp194, which was defined as the atomic distance of Ile16N_c to Asp194C_v.

critical to the structure and function integrity,¹⁵ and mutations involving these sites lead either to null expression or complete loss of function.^{16–19} The non-conserved residues constitute significant portion of FX sequence; however, their significance in structure and function have not been fully revealed.

Naturally, FX exists as a mixture of molecules in zymogen or enzyme conformations.²⁰ Before cleavage and activation, the zymogen conformation is preferred. And once the scissile peptide bond between the heavy and the light chains is cleaved, the balance shift to the activated enzyme decisively. As other serine protease homologous to trypsin, this cleavage forms a new N-terminal of the catalytic domain, whose first eight amino acid residues take a dramatic movement and insert into the activation pocket around Asp194.²¹ Camire and colleagues showed that mutations at the N-terminal of the FX catalytic domain will maintain zymogenicity of FX and become resistant to its inhibitors.²² In the X-ray structure of FXa (PDB ID: 2W26), Asp194 and Ile16 form a featured strong salt bridge in the activation pocket. In the meanwhile, the activated conformation may be further stabilized by a network of hydrogen bonds around the pocket, such as between Asp194 carboxyl and backbone of Gly142 and Cys191, the amino group of Ile16 and backbone of Arg143, the amino group of Lys156 and backbone of Ile16 and Gly18 and the backbone of Val17 and Asp189. In addition, at the edge of the

activation pocket, the backbone hydrogen bonds between Gln20 and Met157 can form an untypical β sheet, which is also observed in other coagulation factors like FII, FVII, FIX and FXI.^{23–26} These interactions provide an extra anchor to fix the N-terminal segment at the entry of the activation pocket and facilitate the insertion. At last, Cys22 lies at the root of the swing N-terminal segment and forms a disulfide bond with Cys27. However, both the two cysteine residues and the disulfide bond between them are not totally conserved among the coagulation factors of the serine protease family. The contribution of the Cys22-Cys27 disulfide bond to the conformational transition of FX activation and enzymatic activity of FXa is still not clear. It is interesting to explore the effect of this disulfide bond on the enzyme conformation and its implication to the enzyme function.

The naturally occurring mutation p.Cys27Ser of FX provides important clues to the residues' contribution in physiological haemostasis and pathological thrombosis. F10 gene mutations either lead to insufficient expression or secretion of FX or encode FX with defective clotting activity. The FX deficiency is thus grossly sub-divided into two categories, CRM+ and CRM negative (CRM-), according to the ratio between residual FX:C and FX:Ag.²⁷ The p.Cys27Ser mutation was identified in patients with compound F10 mutations and the genotypic and phenotypic study of the patient and pedigree members showed that the mechanism underlying FX deficiency caused by mutation p. Cys27Ser cannot be attributed to either category exclusively. Both FX:Ag and FX:C decreased in heterozygous carrier of mutation p.Cys27Ser; however, unlike typical CRM- mutations, the FX:Ag and FX:C level are not consistent, with reduction of FXI:C more prominent than FXI:Ag. The in vitro expression study showed similar effect of mutation p.Cys27Ser on FX:C and FX:Ag, suggesting that amino acid substitution p.Cys27Ser not only impairs the expression and secretion of FX, but also compromises the zymogen activation or enzyme activity of FX.

The p.Cys27Ser replacement disrupts the disulfide bond, and as the results of most amino acid replacement, unorthodoxically folded FX mutant is thus accumulated in the ER system, leading to decreased FX:Ag in circulation. The FX p. Cys27Ser mutant, which passed quality control in the ER system and secreted into circulation, cannot be converted into its active form as efficient as its WT counterpart; furthermore, the enzymatic activity of activated FXa p.Cys27Ser mutant also becomes lower than WT FXa. The enzymatic kinetic analysis revealed higher Km and almost unaltered kcat towards synthetic chromogenic substrate, suggesting reduced binding affinity between the FXa enzyme and substrate. It is also interesting to note that although the impairment of FXa enzymatic activity by amino acid substitution p. Cys27Ser against its physiological substrate, pro-thrombin, is dramatic, the functional defect of the pro-thrombinase complex is relatively minor, which may attribute to the conformational change facilitated by co-factor FVa.²²

In order to explore the molecular mechanism of the differences between WT FX and FX mutant p. Cys27Ser, we compared the conformations and the dynamic motions of the Cys27Ser mutant and the WT FXa with the Cys22-Cys27 disulfide bond formed and broken, especially around the

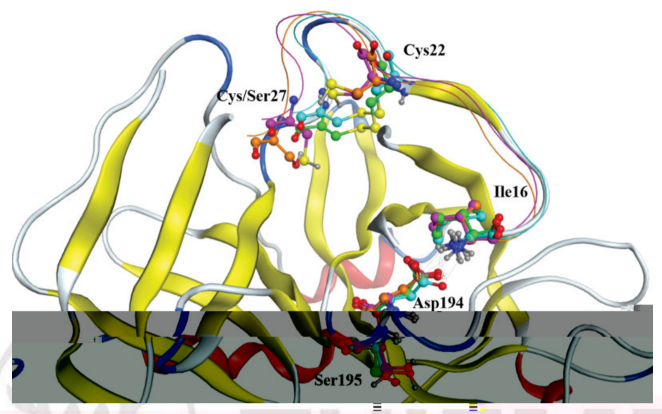


Fig. 6 Comparison of the activated conformations. The critical residues at the mutation site (Cys22 and Cys/Ser27), the activation pocket (Ile16, Asp194) and the catalytic site (Ser195) are visualized. Their oxygen atoms are coloured in red, nitrogen in blue, sulphur in yellow, hydrogen in grey, as well as their carbon atoms are coloured in cyan, magenta, orange and green for the wild-type (WT), disulfide broken (DISB) WT, mutated into serine (MT) systems and the template 2W26. The backbones of the N-terminal segment 16 to 27 of the three models (thin lines in cyan, magenta and orange in the same order) are also compared with the template 2W26, whose whole backbone are shown in cartoon and coloured by secondary structure, except that the loops of 70 to 79 and 142 to 153 are hidden for clarity.

activation pocket, by MD simulations.¹⁸ With an advantage in microscopic details, MD simulations have long been applied to studies on the molecular mechanism and drug design of FX coagulation factor, together with crystallographic analyses.^{28,29} However, most of those studies focused on the catalytic pocket or the specificity site, as far as we know, for the first time the activation pocket around Asp194 is looked as the target of MD simulations in this study. According to our MDs simulations, similar stable structures and interaction patterns were found in the inserted conformation of all the three systems, which may suggest similar final active form of these enzymes that resulted in similar kcat of WT and the Cys27Ser mutant. At the same time, the N-terminal segment in the conformation prior to the insertion is quite different. In the WT FXa with Cys22-Cys27 disulfide bond, the fluctuation of the N-terminal segment is quite limited and would be more easily transited into the inserted conformation. On the other hand, if the disulfide bond is broken by reduction or by Cys27Ser mutation, the N-terminal segment had higher possibility to overhang outside the activation pocket with great fluctuations. This difference in efficiency of insertion may partially contribute to the slower conversion of FX p. Cys27Ser zymogen to its active enzyme form. The hindrance of newly formed N-terminal insertion may also suggest higher portion of FX p.Cys27Ser mutant still maintain its

zymogen conformation even after the cleavage of the scissile bond between heavy chain and light chain, which might lead to lower binding affinity, that is, the higher Km of the mutant FXa with its substrates than the WT.

It should be noted that the Cys22-Cys27 disulfide bond is solvent accessible and its formation and disruption might be subjected to the influence of the oxidation/reduction balance in plasma. Therefore, the equilibration of the two species of FX, with or without the disulfide bond, could potentially shift, depending on the environmental oxidization pressures in the circulation. It has been shown that the formation and reduction of solvent accessible disulfide bond may work as a redox switch to regulate protein function. The intra-chain disulfide bond of Cys6-Cys11 of recombinant insulin was reported to affect the chain flexibility.³⁰ It has also been shown that the disulfide bond formation and disruption regulate the angiotensin release.³¹ The transition of angiotensinogen under oxidative environment to its more active form with sulphydryl-bridged plays an important role in pregnancy hypertension and pre-eclampsia. As shown in this study, the FX molecules with the Cys22-Cys27 disulfide bond is functionally advantageous over FX with the disulfide bond reduced. It has been shown that the oxidation pressure is significantly related to the thrombus formation. The simple oxidation/redox balance shift could significantly disturb coagulation reaction in vitro. The FX with Cys22-Cys27

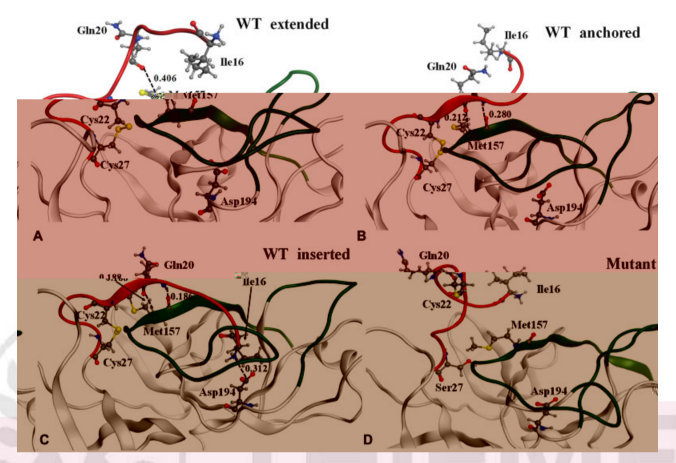


Fig. 7 Simulations of the just cleaved models of the wild-type (WT) and the mutant factor Xa (FXa). (A–C) The N-terminal segment of WT is spontaneously inserted into the activation pocket from the initially extended conformation (A), via an intermediate anchored conformation (B) and finally to the inserted conformation (C). On the other hand, the N-terminal segment of the mutated into serine (MT) system stays at the extended conformation (D) and swings around within the 300-ns simulation.

disulfide bond could be a more prominent form in milieu of vascular injury due to similar oxidation pressure and promote coagulation reactions and may thus contribute to the pathogenesis of thrombus upon atherosclerotic lesions.

Future investigations are warranted to explore the relationship between Cys22-Cys27 disulfide bond and oxidation/ redox balance in plasma and its implication in pathogenesis of thrombosis.

What is known about this topic?

• The conversion of coagulation factor X to its active form factor Xa upon cleavage between heavy and light chains is triggered by insertion of newly formed N terminal of catalytic domain into preformed "activation pocket."

What does this paper add?

• The disruption of non-conserved disulfide bond Cys22-Cys27 by mutation p.Cys27Ser not only hinders secretion of factor X, but also perturbs zymogen to enzyme conformation transition and impairs catalytic activity of factor Xa.

Authors' Contributions

F.L. performed molecular dynamic simulations and analysis; C.C. performed protein expression collection and purification and enzyme kinetical analysis; S.Q. and M.Z. contributed to discussion and manuscript revision of MD results; X.W. performed clinical data acquisition; X. X. and L.L. performed phenotype and genetic analysis; X. W. and Q.D. supervised studies conducted in Ruijin Hospital, Shanghai Jiao Tong University; D.W. supervised studies in School of Life Sciences and Biotechnology, Shanghai Jiao Tong university; and Q.X. and W.W. designed experiments, analysed the data and wrote the manuscript. All authors approved the final version of this manuscript.

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Conflict of Interest None declared.

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