TFPIα Interacts with FVa and FXa to Inhibit Prothrombinase During the Initiation of Coagulation

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TFPIα interacts with FVa and FXa to inhibit prothrombinase during the initiation of coagulation

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Key Points
- TFPIα inhibits prothrombinase through interactions with the FXa active site and B-domain acidic region and heavy chain of FXa-activated FVa.
- Leu252-Thr255 of the TFPIα basic region is necessary for inhibitory activity but not for binding the FXa-activated FVa acidic region.

Tissue factor pathway inhibitor α (TFPIα) inhibits prothrombinase, the thrombin-generating complex of factor Xa (FXa) and factor Va (FVa), during the initiation of coagulation. This inhibition requires binding of a conserved basic region within TFPIα to a conserved acidic region in FXa-activated and platelet-released FVa. In this study, the contribution of interactions between TFPIα and the FXa active site and FVa heavy chain to prothrombinase inhibition were examined to further define the inhibitory biochemistry. Removal of FXa active site binding by mutation or by deletion of the second Kunitz domain (K2) of TFPIα produced 17- or 34-fold weaker prothrombinase inhibition, respectively, establishing that K2 binding to the FXa active site is required for efficient inhibition. Substitution of the TFPIα basic region uncharged residues (Leu252, Ile253, Thr255) with Ala (TFPI-AAKA) produced 5.8-fold decreased inhibition. This finding was confirmed using a basic region peptide (Leu252-Lys261) and Ala substitution peptides, which established that the uncharged residues are required for prothrombinase inhibitory activity but not for binding the FVa acidic region. This suggests that the uncharged residues mediate a secondary interaction with FVa subsequent to acidic region binding. This secondary interaction seems to be with the FVa heavy chain, because the FV Leiden mutation weakened prothrombinase inhibition by TFPIα but did not alter TFPI-AAKA inhibitory activity. Thus, efficient inhibition of prothrombinase by TFPIα requires at least 3 intermolecular interactions: (1) the TFPIα basic region binds the FVa acidic region, (2) K2 binds the FXa active site, and (3) Leu252-Thr255 binds the FVa heavy chain.

Introduction

Prothrombinase, the complex of the serine protease factor Xa (FXa) and its cofactor factor Va (FVa), generates the thrombin needed to form a blood clot.1,2 FVa is generated through proteolytic activation of its precursor, FV, which is maintained as an inactive procofactor through interactions between basic and acidic regions within its activation peptide (B-domain).3-5 Removal of either the acidic or basic region activates FV to FVa.3,4 FVa generated through limited proteolysis by FXa6 and forms of FVa released from activated platelets7,8 retain the acidic region, which remains only during the initiation phase of coagulation, as thrombin catalyzes rapid removal of the entire B-domain.6,9
Forms of prothrombinase containing the FVa acidic region are inhibited by tissue factor pathway inhibitor α (TFPIα), a trivalent Kunitz-type serine protease inhibitor that regulates the initiation of coagulation.10-13

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However, the biochemical mechanism of this inhibition remains unclear. TFPIα contains a 9-residue basic sequence in its C-terminal region (LIKTKKRRK) that is nearly identical to the FV B-domain basic region (LIKTRKKKK). This sequence is conserved in both proteins across mammals. The TFPIβ region tightly binds to the FVa acidic region in a charge-dependent interaction that is blocked by negatively charged polymers, such as heparins and polyphosphates. TFPIβ does not effectively inhibit thrombin generation by prothrombinase if either the FVa acidic region or TFPIβ basic region is missing. Because a peptide mimicking the homologous FV basic region competes against FXa for FVa binding, it has been proposed that this peptide, and by extension TFPIβ, inhibits thrombin generation by displacing FXa from FVα, thereby preventing prothrombinase assembly.

The ability of the FV basic region peptide to compete against FXa for FVa binding suggests that inhibition by the peptide involves at least 2 binding sites: (1) a high-affinity interaction with the FVa acidic region, and (2) a low-affinity interaction with the FVa heavy chain, which does not occur in the absence of acidic region binding. Consistent with this supposition, the FV Leiden (FVL) mutation (Arg506→Gln in the FVa heavy chain) causes an approximately twofold decrease in the ability of either TFPIα or the TFPIβ basic region peptide to inhibit prothrombinase containing the FVa acidic region. This small change in inhibitory kinetics suggests that TFPIα may not bind directly to Arg506, but rather that its binding site is close enough to Arg506 that accessibility is affected by this mutation. In addition, poly-L-lysine does not inhibit prothrombinase activity, demonstrating that the basic residues, although required, are not sufficient for inhibition.

Even though the TFPIα and FV basic region peptides bind the FVa acidic region with low nanomolar affinity, inhibition of thrombin production requires 2 orders of magnitude higher concentrations of either peptide than TFPIα. This discrepancy suggests that the efficient inhibition achieved by TFPIα requires more than simply the interaction of the basic C-terminus with FVα. Because the second Kunitz domain (K2) of TFPIα is a well-described inhibitor of the FXa active site, we hypothesized that this interaction is also required for prothrombinase inhibition. In the present study, full-length forms of TFPIα with amino acid substitutions in K2 that prevent binding to the FXa active site or within the uncharged amino acids of the basic region, truncated forms of TFPIα, and basic region peptides were used in purified protein- and plasma-based assays to define the interactions required for prothrombinase inhibition.

Materials and methods

Proteins

Human FXa, thrombin, prothrombin, and FX were from Enzyme Research Laboratories (South Bend, IN). Human FV was from Paula B. Tracy or Kathleen M. Brummel-Ziedins (University of Pennsylvania, Philadelphia, PA). TF (Dade Innovin) was from Siemens (Washington, DC). Human FVIIa and a mouse antibody against the TFPI K2 domain (anti-K2) were obtained as described. A rabbit polyclonal antibody that recognizes the final 12 amino acids of TFPI (anti-CTP) was produced by Abcam (Burlingame, CA).

Goat anti-mouse IRDye680 and goat anti-rabbit IRDye800CW were from LI-COR Biosciences (Lincoln, NE).

Expression and purification of altered TFPIα proteins

The DNA sequences of TFPIα were synthesized with CD33 signal peptide and cloned into vector pJSV002 for mammalian expression. The genes of TFPIΔK2 (Arg107Ala) and TFPI-AAKA (Leu252Ala/Ile253Ala/Thr255Ala) were constructed based on wild-type TFPIα by site-directed mutagenesis. Plasmids encoding TFPI variants were transfected into HEK293-6E cells (1.0 × 10^6 cells per mL), with DNA molar ratio of 1:1. Transfection was performed following the FreeStyle 293 expression manual (Invitrogen). Cells were harvested for purification 5 days posttransfection.

For TFPIΔK2, the supernatant was applied to an anti-TFPI sepharose affinity column (anti-TFPI antibody coupled to the HiTrap NHS-activated HP column; GE Healthcare) and equilibrated in phosphate-buffered saline (PBS). The bound protein was eluted with 0.1 M of glycine-hydrochloride (HCl), pH 2.8. Fractions were collected and neutralized immediately with 1/20 volume 2 M of Tris-HCl, pH 9.0, for further purification. For TFPI-AAKA, the supernatant was applied to a Heparin Sepharose 6 Fast Flow affinity column (GE Healthcare), equilibrated in 20 mM of Tris-HCl, pH 7.5. The bound protein was eluted with a 0.2- to 1-M linear gradient of sodium chloride. Fractions containing the TFPI variants were collected. The pooled fractions were further purified by size-exclusion chromatography on a Superdex 75 prep grade column (GE Healthcare) in PBS.

His-thioredoxin (Trx)-3C-K3C (TFPIα residues 185-271) was synthesized and cloned into the pET32a vector. The plasmid was transformed into Escherichia coli BL21(DE3) host strain and cultured in terrific broth medium. Expression was induced when optical density measured at a wavelength of 600 nm reached 0.1, with 0.1 mM of isopropyl-β-D-thiogalactoside. Protein was expressed overnight at 18°C, and the bacterial pellets were then harvested for purification. The cell pellets were resuspended in lysis buffer (50 mM of Tris, pH 8.0, 0.1% Tween-20) and disrupted by French press. Cell debris was removed by centrifugation. Supernatant was applied to an Ni-NTA Superflow column (QIAGEN), equilibrated in 50 mM of Tris, pH 8.0; 4 M of urea; 300 mM of sodium chloride; and 20 mM of imidazole. The bound protein was eluted with a 20- to 400-mM linear gradient of imidazole. The pooled fractions were further purified by cation exchange chromatography on an SP-HP column (GE Healthcare). Eluted protein was digested by 3C protease to remove the His-Trx-3C tag. Then, the TFPI K3C fragment was further purified using an SP-HP column (GE Healthcare) and buffer exchanged to PBS.

All purified proteins were sterilized by filtration through a 0.2-μm filter unit (Sartorius). The purity was analyzed by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) and size-exclusion high-performance liquid chromatography. The protein identity was confirmed by mass spectrometry.

Peptides

Peptides were made by the Protein Chemistry Core Laboratory (BloodCenter of Wisconsin, Milwaukee, WI) or Peptide 2.0 (Chantilly, VA). Peptide concentrations were determined by measurement at 205 nm and comparison with a standard curve generated using a peptide of similar molecular weight and isoelectric point (GPYVKQNTLKLAT). The concentration of the peptide ran...
standard was measured at 280 nm (E1%, 10.4). The wild-type peptide (LIKTKRKRKK) was also synthesized with 2 N-terminal aminohexanoic acid spacers and 5(6)carboxy-X-hodamine (Life Technologies, Grand Island, NY) coupled to the N-terminus (Rhod-LIKTKRKRKK). The concentration of Rhod-LIKTKRKRKK was determined by measuring the absorbance at 575 nm (carboxy-X-hodamine; E1M, 82 000) and at 205 nm, which yielded similar results (190 and 215 μM, respectively).

**Phospholipid vesicles**

Phospholipid vesicles containing 20% phosphatidylserine, 20% phosphatidylethanolamine, and 60% phosphatidylcholine were prepared according to the method of Morrissey.26

**Prothrombinase activity assays**

FVακα, FVακα, FV810, or FV810 (0.5 nM) was incubated with phospholipid vesicles (20 μM), the thrombin inhibitor dansylarginine N-(3-ethyl-1,5-pentanediy)amide (DAPA; 3 μM; Haematologic Technologies, Essex Junction, VT), and varying concentrations of phospholipid vesicles (20 μM; Haematologic Technologies, Essex Junction, VT), and varying concentrations of TFPIα protein or peptide. DAPA was included in these reactions to limit thrombin-mediated removal of the FVa B-domain acidic region. Nesheim et al27 demonstrated that the inclusion of DAPA does not alter the rate of thrombin generation. Reactions were initiated by addition of prothrombin (1.4 μM) and FXa (5 nM). Aliquots were removed at timed intervals and quenched with EDTA (33 mM), diluting DAPA to a final concentration of 0.2 μM. Thrombin was measured using the chromogenic substrate Spectrozyme TH (0.32 mM; Sekisui Diagnostics, Lexington, MA). GraphPad Prism v.6 (GraphPad Software, La Jolla, CA) was used to determine 50% inhibition of FXa activity assays using TFPI-depleted plasma was initiated with FXa (0.1 nM) and collagen (15 μg/mL). Assays using PRP were initiated with FXa (0.1 nM) and collagen (15 μg/mL). Assays using TFPI-depleted plasma were initiated with FXa and phospholipid vesicles (4 μM).

**FXa activity assays**

FXa (0.2 nM) was incubated with varying concentrations of the altered TFPIα proteins, and cleavage of the chromogenic substrate Spectrozyme Xa (0.5 mM; Sekisui Diagnostics) was monitored at 405 nm.

**TF-FVIIa activity assays**

TF (6 pM) was incubated with phospholipid vesicles (20 μM), FVIIa (20 pM), and varying concentrations of the altered TFPIα proteins. Reactions were initiated by addition of FX (20 nM), and cleavage of Spectrozyme Xa was monitored at 405 nm.

**Fluorescence anisotropy**

Steady-state fluorescence anisotropy was measured in a QuantaMaster spectralfluorometer (Photon Technology International, Edison, NJ).12 Emission and excitation wavelengths were 580 and 605 nm, respectively. For direct binding measurements, Rhod-LIKTKRKRKK (30 nM) was incubated with increasing concentrations of FV810. For displacement experiments, Rhod-LIKTKRKRKK (30 nM) and FV810 (30 nM) were incubated with increasing concentrations of unlabeled peptide. Distribution coefficient (Kd) values were obtained from the direct binding32 and displacement experiments.31

**Western blotting**

Recombinant TFPIα proteins were diluted into sample preparation buffer (62.5 mM of Tris, pH 6.8; 10% glycerol; 1% SDS; 0.0005% Bromphenol Blue), separated by SDS-PAGE on a 4% to 20% acrylamide gel.32 and transferred to nitrocellulose.33 The blot was probed with mouse anti-K2 and rabbit anti-CTP (5 μg/mL each), followed by detection with goat anti-mouse IRDye680 and goat anti-rabbit IRDye800CW secondary antibodies (1:10 000 dilution each). The blot was developed using Odyssey Infrared Imager Running Image Studio v.4.0 (LI-COR Biosciences).

**Results**

**Characterization of recombinant TFPIα proteins**

The roles of the TFPIα K2 domain and the uncharged residues within the TFPIα basic region in prothrombinase inhibition were examined using altered recombinant TFPIα proteins (Figure 1A). These proteins included: (1) wild-type TFPIα; (2) 2 altered TFPIα proteins incapable of binding the FXa active site, TFPIαK2 (Arg107 of K2 mutated to Ala) and K3C, which lacks the K1 and K2 domains; and (3) TFPIα in which the uncharged residues of the basic region (Leu252, Ile253, Thr255) were substituted with Ala (TFPIα-AAKA). Because the TFPIα C-terminus is susceptible to proteolysis, the presence of an intact C-terminus was verified by western blotting with an antibody directed against the final 12 amino acids of TFPIα (Figure 1B). This antibody recognized all of the recombinant proteins used in these studies, confirming the presence of an intact C-terminus in each. A recombinant TFPIα containing only the K1 and K2 domains, previously shown to be a poor inhibitor of prothrombinase,12 was included as a negative control for the western blot. Similarly, an antibody against the K2 domain (anti-K2) recognized all of the recombinant proteins except K3C. TFPIα and TFPIα-AAKA directly inhibited FXa and also inhibited TF-FVIIa in assays using purified proteins, confirming the presence of active K1 and K2 domains (supplemental Figures 1 and 2). K3C did not inhibit TF-FVIIa or FXa, and TFPIαK2 only weakly inhibited FXa. The weak FXa inhibition by TFPIαK2 was likely caused by the functional K1 domain, as demonstrated by Petersen et al.34

**Binding of the FXa active site is required for efficient prothrombinase inhibition**

The role of the K2 domain in prothrombinase inhibition was evaluated using TFPIαK2 and K3C. K3C is incapable of inhibiting the FXa active site, and TFPIαK2 is a poor FXa inhibitor (supplemental Figure 1). In purified prothrombinase assays, TFPIαK2 was a 17-fold weaker inhibitor than wild-type TFPIα (IC50, 30.6 vs 1.8 nM), and K3C was even weaker than TFPIαK2 (IC50, 61.4 nM; Figure 2A). Consistent with the prothrombinase assays using purified proteins, TFPIα dose dependently inhibited FXa-initiated thrombin generation in TFPIα-depleted plasma, increasing the lag time by 79.9% ± 4.3% at 5 nM and 214.6% ± 20.6% at 10 nM (Figure 2B). TFPIαK2 had significantly less effect, only increasing the lag time by 45.5% ± 8.1% at 10 nM (P < .0001; Figure 2C). Because activated platelets
release forms of FVa that bind TFPI\(\alpha\) and provide a surface for prothrombinase assembly, the inhibitory activity of TFPI\(\alpha\) variants was also evaluated in PRP. Although this system is complicated by the presence of endogenous TFPI\(\alpha\), a similar inhibitory pattern was apparent. In PRP, 5 nM of TFPI\(\alpha\) inhibited FXa-initiated thrombin generation, delaying the lag time by 31% ± 10% (Figure 2D). In contrast, thrombin generation was not inhibited by 10 nM of TFPI\(\Delta\)K2 (Figure 2E) or K3C (Figure 2F). These results demonstrate that binding of TFPI\(\alpha\) to the FXa active site is required for efficient prothrombinase inhibition in assays performed using purified proteins and in assays performed using TFPI-depleted plasma or PRP.

**Uncharged residues of the TFPI\(\alpha\) basic region are required for efficient prothrombinase inhibition**

We previously showed that the basic residues of the TFPI\(\alpha\) basic region are required for prothrombinase inhibition, because this activity is blocked by negatively charged molecules, including heparins and polyphosphates.\(^{12,18}\) However, basic residues are not sufficient for inhibition, because poly-L-lysine does not block prothrombinase activity.\(^{21}\) Interestingly, the basic regions of both TFPI\(\alpha\) and FV contain 3 uncharged residues (Leu252, Ile253, and Thr255 in human TFPI\(\alpha\)). These uncharged residues are conserved in both proteins in mammals, birds, and reptiles (supplemental Figures 3-6). TFPI-AAKA, in which Leu252, Ile253, and Thr255 are substituted with Ala, was an intermediate inhibitor of purified prothrombinase (IC\(_{50}\) 10.4 nM) compared with TFPI\(\alpha\) (1.8 nM) and TFPI\(\Delta\)K2 (30.6 nM; Figure 3A). Consistent with the prothrombinase assays using purified proteins, TFPI-AAKA was a weaker inhibitor of FXa-initiated thrombin generation in TFPI-depleted plasma (Figure 3B) or PRP (Figure 3C) than TFPI\(\alpha\). TFPI-AAKA (10 nM) was similar to 5 nM of TFPI\(\alpha\) in delaying the lag time by 34% ± 4%, whereas 5 nM of TFPI-AAKA delayed the lag time by 17% ± 2%.

The importance of Leu252, Ile253, and Thr255 for prothrombinase inhibition was more apparent in studies using a TFPI\(\alpha\) basic region peptide (LIKTKRKRKK). Similar to a 26-residue C-terminal peptide previously characterized,\(^{12}\) LIKTKRKRKK inhibited prothrombinase assembled with FVaXa (IC\(_{50}\) 1.0 \(\mu\)M) but did not inhibit prothrombinase assembled with FVaIIa (Figure 4A; Table 1) and prolonged the lag time of FXa-initiated thrombin generation in PRP (Figure 4B). These results are consistent with inhibition requiring the FVa acidic region,\(^{12,16}\) which is present in FVaXa but absent from FVaIIa.\(^{6}\) Substitution of all 3 uncharged residues with Ala (AAAKKRKRKK) essentially abolished inhibitory activity (Figure 4C; Table 1). Similarly, AAKAKKRKRKK did not inhibit thrombin generation in PRP (Figure 4D). These results demonstrate that the uncharged Leu, Ile, and Thr amino acids within the TFPI\(\alpha\) basic region are required for efficient prothrombinase inhibition in assays performed using purified proteins and in assays performed using PRP.
Leu252, Ile253, and Thr255 each contribute to inhibitory activity

Single Ala substitution peptides were used to examine the individual contributions of Leu252, Ile253, and Thr255 to the inhibition of prothrombinase by the TFPIε basic region. Substitution of any of the 3 uncharged residues resulted in decreased prothrombinase inhibitory activity (Figure 5A). The Leu252 (AIKTKRKRKK) and Thr255 (LIKAKRKRKK) substitution peptides had the least inhibitory activity (IC50, 70 and 110 mM, respectively), whereas the Ile253 substitution peptide (LAKTKRKRKK) retained greater inhibitory activity (IC50, 17 mM). None of the single Ala peptides inhibited thrombin generation in PRP, at concentrations up to 20 mM (Figure 5B-D). Thus, each of these residues individually contributes to the inhibitory activity of the basic region peptide.

Conserved basic residues mediate FVa binding

Fluorescence anisotropy was used to measure peptide binding to FV810, a recombinant form of FVa that contains the B-domain acidic region and tightly binds TFPIε. The Kd of each peptide for binding FV810 was determined by measuring its ability to compete against Rhod-LIKTKRKRKK (Kd, 8.6 nM). All of the altered peptides had similar affinity for FV810, which was not different than that of LIKTKRKRKK (Figure 6A; Table 1). Thus, the uncharged residues do not contribute significantly to the binding affinity for FVa.

Because AAKAKRKRKK binds FV810 similarly to LIKTKRKRKK (Figure 6A) but does not inhibit prothrombinase activity (Figure 4D), we hypothesized that AAKAKRKRKK would compete against LIKTKRKRKK for binding to the FVa acidic region and prevent prothrombinase inhibition. Purified prothrombinase assays were performed in the presence of LIKTKRKRKK (3 mM) and increasing concentrations of AAKAKRKRKK, which reversed prothrombinase inhibition in a dose-dependent manner (IC50, 6.0 mM; Figure 6B).

Residues Leu252-Thr255 are disrupted by FVL

Because residues Leu252-Thr255 were not necessary for binding the FVa acidic region but were required for optimal inhibitory activity, we hypothesized that they may be involved in an interaction for binding FV810.
with the FVa heavy chain and mediate displacement of FXa. Consistent with this hypothesis, prothrombinase assembled with FVa Leiden (Arg506Gln) is somewhat less susceptible to inhibition by TFPI $\alpha$ than wild-type FVa.22 We compared the ability of TFPI-AAKA to inhibit prothrombinase assembled with either FV810 or FVL810. Consistent with the experiments using FXa-activated FVa, TFPI-AAKA was a weak inhibitor of prothrombinase containing FV810 (IC$_{50}$, 16.9 nM; 95% CI, 14.9-20.5 nM; Figure 7A). However, unlike TFPI$\alpha$, TFPI-AAKA inhibited prothrombinase containing FVL810 identically to prothrombinase containing FV810 (IC$_{50}$, 17.5 nM; 95% CI, 14.6-19.5 nM). In contrast, the interaction of TFPI$\alpha$ with FXa is unrelated to the effect of FVL, because TFPI$\Delta$K2 was a 1.6-fold weaker inhibitor of prothrombinase containing FVL810 (IC$_{50}$, 46.7 nM; 95% CI, 41.3-52.9 nM) compared with FV810 (IC$_{50}$, 27.9 nM; 95% CI, 21.7-35.9 nM), similar to the 1.7-fold shift observed with TFPI$\alpha$ (Figure 7B).22

**Discussion**

Inhibition of prothrombinase by TFPI$\alpha$ requires an exosite interaction between the TFPI$\alpha$ basic region and the FVa acidic region. Multiple lines of evidence support this assertion: (1) in vitro studies have shown that the TFPI$\alpha$ basic C-terminus binds to forms of FVa containing the acidic region with high affinity (K$_d$, ~90 pM) but does not bind to forms of FVa lacking the acidic region12,15; (2) 2 FV

---

**Figure 3.** The basic region uncharged residues enhance prothrombinase inhibition. (A) Prothrombinase activity was measured as in Figure 2A, in the presence of the indicated concentrations of TFPI-AAKA (filled square). The initial rate of thrombin generation is shown as a percentage of control (mean ± standard deviation; n ≥ 3). Lines represent best-fit inhibition curves. Inhibition of prothrombinase by TFPI$\alpha$ (open circle) and TFPI$\Delta$K2 (open square) is reproduced from Figure 2 for reference. (B-C) Thrombin generation was measured in TFPI-depleted plasma (B) or PRP (C) as in Figure 2, in the presence of the indicated concentrations of TFPI-AAKA. Shown are average thrombin generation curves from experiments performed in triplicate using TFPI-depleted plasma (B) or PRP from 4 donors (C).

---

**Figure 4.** The uncharged residues are required for prothrombinase inhibition by a peptide mimicking the TFPI$\alpha$ basic region. (A) Prothrombinase activity assays were performed as in Figure 2A using either FVa$\alpha$ (filled square) or FVaa (open circle) and the indicated concentrations of LIKTKRKRKK (LIKT). The initial rate of thrombin generation is shown as a percentage of control (mean ± standard deviation; n ≥ 3). Lines represent best-fit inhibition curves. (B,D) Thrombin generation was measured in platelet-rich plasma as in Figure 2D, in the presence of the indicated concentrations of LIKT (B) or AAKAKRKRKK (AAKA) (D). Shown are average thrombin generation curves from experiments using at least 3 donors. (C) Prothrombinase activity assays were performed using FVa$\alpha$ and the indicated concentrations of AAKA (open circle). Inhibition of FVa$\alpha$, prothrombinase by LIKT (filled square) is reproduced from panel A for reference.
because the basic region peptides bind the FVa acidic region with low nanomolar affinity, suggesting that structural features of TFPI K2 domains beyond its basic region mediate prothrombinase inhibition. FXa-activated FVa (used in the activity assays) and FV810 (used for binding measurements) are chemically dissimilar forms of FV/FVa. The structural differences between these forms were considered as an explanation for the discrepancy between the binding affinity and inhibitory activity observed. This does not seem likely. The major difference between FXa-activated FVa and FV810 is that the former contains the B-domain basic region peptide at 0.5 nM (equimolar concentration to the FVa) that is cleaved from FV when it is activated by FXa. However, micromolar concentrations of LKTKRRKK are required to inhibit prothrombinase containing FV810 similar to prothrombinase containing FXa-activated FVa, suggesting that nanomolar concentrations of the FV B-domain basic peptide have little effect on the inhibitory reaction. In this study, 2 additional interactions were identified that greatly increase the efficiency of prothrombinase inhibition, allowing for inhibition at low nanomolar concentrations of TFPI K2 (1) the TFPI K2 domain binds the FXa active site, and (2) residues Leu252-Thr255 of the K2 domain are required. This mechanism has since been extrapolated to full-length TFPI, which is being described as an inhibitor of prothrombinase assembly. However, basic region peptides inhibit prothrombinase only at micromolar concentrations, whereas TFPI K2 inhibits at low nanomolar concentrations. This is a striking discrepancy, particularly

Table 1. Prothrombinase inhibition by peptide variants

<table>
<thead>
<tr>
<th>Peptide</th>
<th>IC50, µM (95% CI)*</th>
<th>Kd (nM ± SEM)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>LKTKRRKK</td>
<td>1.0 (0.4-2.7)</td>
<td>6.0 ± 2.5</td>
</tr>
<tr>
<td>AAKAKRRKK</td>
<td>&gt;300</td>
<td>5.9 ± 2.4</td>
</tr>
<tr>
<td>AKTKRRKK</td>
<td>70 (60-82)</td>
<td>10.8 ± 3.9</td>
</tr>
<tr>
<td>LAKTKRRKK</td>
<td>17 (7.7-36)</td>
<td>16.2 ± 6.5</td>
</tr>
<tr>
<td>LIKAKRRKK</td>
<td>110 (89-130)</td>
<td>9.3 ± 3.6</td>
</tr>
</tbody>
</table>

CI, confidence interval; SEM, standard error of the mean.
*IC50 values determined by fitting the data presented in Figures 4 and 5 to a 4-parameter variable slope equation using GraphPad Prism v.6.
†Binding affinities determined by fitting the data presented in Figure 6A, as described by Marchand et al.4

Figure 5. Leu252, Ile253, and Thr255 each contribute to prothrombinase inhibition. (A) Prothrombinase activity was measured as in Figure 2A, in the presence of the indicated concentrations of AKTKRRKK (AKT; open diamond), LKTKRRKK (LAKT; open circle), or LIKAKRRKK (LIKA; filled triangle). The initial rate of thrombin generation is shown as a percentage of control (no peptide; mean ± standard deviation; n = 3). Lines represent best-fit inhibition curves. Inhibition of prothrombinase by LKTKRRKK (LKJT; filled square) is reproduced from Figure 3 for reference. (B-D) Thrombin generation was measured in PRP as in Figure 2D, using the indicated concentrations of AKT (B), LAKT (C), or LIKA (D). Shown are the average thrombin generation curves from experiments using 3 donors.
et al. Studies with FVL were used to further pursue this possibility. by blocking the binding of FXa to FVa, as proposed by Bunce. FVa heavy chain in a manner that blocks prothrombinase assembly inhibition suggested that these amino acids may interact with the uncharged residues for prothrombinase activity, because poly-L-lysine does not block prothrombinase assembly or function. Here, we establish that the uncharged residues of the TFPI basic region (Leu252, Ile253, Thr255) are required for efficient prothrombinase inhibitory activity. In experiments using recombinant TFPI with these amino acids substituted with Ala and several basic region peptides with the 3 amino acids either all or individually substituted with Ala. Interestingly, even though the uncharged residues are highly conserved, they did not contribute to the high-affinity interaction with the FVa acidic region, but they were required for prothrombinase inhibition.

The requirement of the uncharged amino acids for prothrombinase inhibition suggested that these amino acids may interact with the FVa heavy chain in a manner that blocks prothrombinase assembly by blocking the binding of FXa to FVa, as proposed by Bunce et al. Studies with FVL were used to further pursue this possibility. FVL is the most common genetic risk factor for thrombosis in whites and consists of the substitution of Arg506 in the heavy chain with Gin. FVL is resistant to proteolysis by activated protein C, providing 1 biochemical mechanism for the increased prevalence of thrombosis. Our laboratory recently found that TFPI is a 1.7-fold weaker inhibitor of prothrombinase containing FVL810 compared with FV810. Similarly, the prothrombinase inhibitory activity of LIKTKRKRKK was decreased 2.1-fold and that of TFPI-R107A was reduced 1.6-fold when using FVL810. These small shifts in inhibitory kinetics become physiologically meaningful under conditions of reduced TFPI concentration. For example, the combination of FVL and TFPI haploinsufficiency is lethal in mice, which die as a result of perinatal thrombosis. Because FXa binds FVa near Arg506, it was plausible that the uncharged amino acids within the TFPI basic region also bind near Arg506, perhaps explaining the decreased inhibitory activity toward FVL. Therefore, experiments were performed to examine the ability of the different altered forms of TFPI to inhibit prothrombinase assembled with FVL. Unlike TFPI, the inhibitory activity of TFPI-AAKA was not affected by FVL, a finding consistent with the uncharged amino acids interacting with the FVa heavy chain. The small shift in kinetics suggests that this interaction is near, but may not directly involve, Arg506.

Collectively, these findings can be described using a model where TFPI inhibits prothrombinase via at least 3 biochemical interactions:

The TFPI and FV basic regions contain conserved basic and uncharged residues. The importance of the basic residues is evident from the ability of negatively charged polymers, such as heparin, polyphosphate, and fucoidan, to block prothrombinase inhibition. However, the basic residues are not sufficient for inhibitory activity, because poly-L-lysine does not block prothrombinase assembly or function. Here, we establish that the uncharged residues of the TFPI basic region (Leu252, Ile253, Thr255) are required for efficient prothrombinase inhibitory activity in experiments using recombinant TFPI with these amino acids substituted with Ala and several basic region peptides with the 3 amino acids either all or individually substituted with Ala. Interestingly, even though the uncharged residues are highly conserved, they did not contribute to the high-affinity interaction with the FVa acidic region, but they were required for prothrombinase inhibition.

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Collectively, these findings can be described using a model where TFPI inhibits prothrombinase via at least 3 biochemical interactions:

Figure 6. The basic amino acids, but not the uncharged amino acids, mediate binding of the TFPI basic region to the FVa acidic region. (A) Rhod-LIKTKRKRKK (30 nM) was incubated with FV810 (30 nM) and the indicated concentrations of LIKTKRKRKK (LIKT; filled square), AAKAKRKRKK (AAKA; open square), AAKRKRKRKK (AIKT; filled circle), LAKTKRKRKK (LAKT; open circle), or LIKAKRKRKK (LIKA; filled triangle). Fluorescence anisotropy was measured and curve fits generated as described in Materials and methods. (B) FV810 (0.5 nM), phospholipid vesicles (20 µM), and the thrombin inhibitor DAPA (3 µM) were incubated with LIKTKRKRKK (LIKT; 3 µM) and varying concentrations of AAKA, and thrombin generation was measured as in Figure 2A. The initial rate of thrombin generation is shown as a percentage of control (no peptide; mean ± standard deviation; n = 3).

Figure 7. The Leu252-Thr254 region is responsible for the effect of FVL on prothrombinase inhibition by TFPI. FV810 (filled circle) or FVL810 (open circle; 0.5 nM) was incubated with phospholipid vesicles (20 µM), the thrombin inhibitor DAPA (3 µM), and the indicated concentrations of TFPI-AAKA (A) or TFPIΔLK2 (B), and thrombin generation was measured as in Figure 2A. The initial rate of thrombin generation is shown as a percentage of control (mean ± standard deviation; n = 3). Lines represent best-fit inhibition curves.
(1) the TFPIκ basic region binds the FVa acidic region in a charge-dependent interaction that is mediated by the respective basic and acidic residues; (2) amino acids Leu252-Thr255 interact with the FVa heavy chain in the vicinity of Arg506, either preventing FXa from binding FVa or causing it to bind in a manner that does not promote thrombin generation; and (3) the K2 domain of TFPIκ binds to the FXa active site. These interactions might occur in any order, although it seems most plausible that TFPIκ binds to the FVa acidic region before the heavy chain, because the latter is a low-affinity interaction. This model explains the discrepancy between the high affinity and low inhibitory activity of the TFPIκ and FV basic region peptides. Although these peptides are able to bind the FVa acidic region and heavy chain and block prothrombinase assembly, they do not block the FXa active site. High concentrations of peptide are required to push the equilibrium such that no FXa can bind and thus no activity can be measured. The model also explains why poly-L-lysine has no inhibitory activity,21 because it does not contain the Leu, Ile, and Thr residues and therefore would not interact with the FVa heavy chain and displace FXa.

This model is limited because it does not include other described mechanisms for how TFPIκ may alter prothrombinase activity. It was recently reported that TFPIκ, as well as a peptide mimicking the TFPIκ C-terminus, inhibits thrombin-mediated cleavage of FV at Arg1545.48 This inhibition is proposed to be the result of steric hindrance (ie, that the acidic region is close enough to Arg1545 such that when TFPIκ or the peptide is bound to the B-domain, thrombin cannot access the cleavage site). Thus, TFPIκ not only reduces thrombin generation through inhibition of prothrombinase assembly and activity, but also reduces the feedback activity of thrombin by blocking further FVa processing. Similarly, TFPIκ may alter cleavage of FVs at Arg506 by activated protein C49 or at Arg510 by factor Xla,50 because the TFPIκ basic region binds near these sites.

In summary, TFPIκ inhibits prothrombinase activity through the combination of disrupting prothrombinase assembly and blocking the FXa active site. This inhibition is mediated by the C-terminal basic region and the K2 domain. The basic and uncharged amino acids of the C-terminal region serve distinct roles in the inhibitory biochemistry. Each of these 3 interactions increases the efficiency of prothrombinase inhibition by TFPIκ. Therefore, all 3 are likely required for prothrombinase inhibition under physiological conditions.

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Authorship

Contribution: J.P.W. performed experiments, analyzed results, and wrote the manuscript; H.H.P., B.Y., X.W., and I.H. produced the TFPI variant proteins and wrote the manuscript; and A.E.M. designed the research, analyzed results, and wrote the manuscript.


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