Antithrombotic Effect of a Protein-Type I Class Snake Venom Metalloproteinase, Kistomin, Is Mediated by Affecting Glycoprotein Ib-von Willebrand Factor Interaction

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ABSTRACT

Binding of von Willebrand factor (vWF) to platelet glycoprotein (GP) Ib-IX-V mediates platelet activation in the early stage of thrombus formation. Kistomin, a snake venom metalloproteinase (SVMP) purified from venom of Calloselasma rhodostoma, has been shown to inhibit vWF-induced platelet aggregation. However, its action mechanism, structure-function relationship, and in vivo antithrombotic effects are still largely unknown. In the present study, cDNA encoding kistomin precursor was cloned and revealed that kistomin is a P-I class SVMP with only a protease domain. Further analysis indicated that kistomin specifically inhibited vWF-induced platelet aggregation through binding and cleavage of platelet GPIbα and vWF. Cleavage of platelet GPIbα by kistomin resulted in release of 45- and 130-kDa soluble fragments, indicating that kistomin cleaves GPIbα at two distinct sites. In parallel, cleavage of vWF by kistomin also resulted in the formation of low-molecular-mass multimers of vWF. In ex vivo and in vivo studies, kistomin cleaved platelet GPIbα in whole blood. Moreover, GPIbα agonist-induced platelet aggregation ex vivo was inhibited, and tail-bleeding time was prolonged in mice administered kistomin intravenously. Kistomin’s in vivo antithrombotic effect was also evidenced by prolonging the occlusion time in mesenteric microvessels of mice. In conclusion, kistomin, a P-I class metalloproteinase, has a relative specificity for GPIbα and vWF and its proteolytic activity on GPIbα-vWF is responsible for its antithrombotic activity both in vitro and in vivo. Kistomin can be useful as a tool for studying metalloproteinase-substrate interactions and has potential being developed as an antithrombotic agent.

Platelets play a key role in hemostasis and thrombosis. Exposure of subendothelial von Willebrand factor (vWF) is the first step to form thrombi to arrest blood loss at the sites of trauma, but abnormal embolism may also cause ischemia in pathogenic condition (Andrews and Berndt, 2004). The glycoprotein (GP) Ib complex (one of the major adhesive receptors expressed on platelets) that interacts with vWF is composed of GPIbα, GPIbβ, GPIIX, and GPV. GPIbα consists of N-terminal flank, leucine-rich repeat, anionic sulfated tyrosine sequence, macroglycopeptide domain, transmembrane region, and cytoplasmic tail (Andrews et al., 2003). Plasma vWF circulates primarily as the dimer form, and the multimeric forms of vWF are existed in the subendothelial matrix (Canobbio et al., 2004). It has been reported that Bernard-Soulier syndrome and platelet-type von Willebrand disease are inherited bleeding disorders caused by mutations in GPIb complex and vWF gene, respectively, suggesting that GPIbα-vWF interaction is very important for hemostasis. Therefore, modulation of the GPIbα-vWF interactions during thrombotic complications could be beneficial (Bonnefoy et al., 2003). However, in contrast to extensive application of αIIbβ3 antagonists during acute coronary diseases, no GPIbα-vWF axis inhibitor is commercially available, although some GPIbα or vWF antagonists are being developed.

Snake venom proteases are invaluable tools for studying coagulation and hemostasis (Marsh, 2001). For examples, fibrinogen and antithrombin III can be assayed by using snake venom thrombin-like enzymes. Among these snake-derived proteases, snake venom metalloproteinases (SVMPs), which are abundant in venoms from Viperidae and Crotalinae, are key enzymes responsible for local hemorrhage and are metal ion-dependent for their full function (Matsui et al., 2000; Kamiguti, 2005). The protein structural classification of SVMPs is presented as protein-type I (P-I;
having only metalloproteinase domain), P-II (having metalloproteinase and disintegrin domain), P-III (having metalloproteinase, disintegrin-like and cysteine-rich domain), and P-IV (having P-III structure plus lectin-like domains connected by disulfide bonds) (Fox and Serrano, 2005). It has been suggested that the additional disintegrin-like and the cysteine-rich regions domains may direct SVMP to its targets (Fox and Serrano, 2005). However, its structure-activity relationship remains unclear.

Kistomin, a 25-kDa SVMP purified from Calloselasma rhodostoma venom in our laboratory, has been shown to degrade fibrinogen and inhibit ristocetin-induced platelet agglutination, suggesting that it is a GPIbcleaving protease (Huang et al., 1993). However, its action mechanism, structure-function relationship, and in vivo antithrombotic effects are still largely unknown. In this study, cDNA-encoding kistomin was cloned and kistomin’s cleaving and binding specificities for vWF and GP Ibα were demonstrated. More importantly, kistomin’s antithrombotic effect was examined in an in vivo animal model.

Materials and Methods

Materials. Anti-GP Ibα mAb M45 and SZ2, directed to the sulfated tyrosine residues of GP Ibα and inhibited ristocetin-dependent binding of vWF to GP Ibα, were obtained from CLB Immunoreagents (Amsterdam, The Netherlands) and Immunotech (Marseillel, France), respectively. The murine mAb against α2β1, 6F1, was kindly provided by Dr. Barry S. Coller (Mount Sinai School of Medicine, New York, NY). FITC-conjugated goat anti-mouse IgG was from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA). Heparin was kindly provided by Dr. Barry S. Coller (Mount Sinai School of Medicine). PCR was performed by using Taq DNA polymerase (Ab Peptides, St. Louis, MO) and the primers with a hot start at 94°C for 10 min and 30 cycles of denaturation (1 min, at 94°C), annealing (1 min, at 50°C), and extension (2 min, at 72°C). The murine mAb against α2β1, 6F1, was kindly provided by Dr. Barry S. Coller (Mount Sinai School of Medicine, New York, NY). FITC-conjugated goat anti-mouse IgG was from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA). Heparin was kindly provided by Dr. Barry S. Coller (Mount Sinai School of Medicine). Primer-annealing temperature was set at 45°C. The final PCR products were analyzed by 1% (w/v) agarose-gel electrophoresis and purified by electroelution. The purified DNA fragments were ligated to pEHI-TOPO vector with a Thermus aquaticus cloning kit (Invitrogen, Carlsbad, CA), and sequence analysis was performed. The specificity of the second amplification was confirmed from the overlapping sequences that were distinguishable from the other SVMPs. Sequences were assembled with the GCG program (Wisconsin Package version 10.1; Genetics Computer Group).

Flow Cytometric and Western Blot Analysis of Platelet Receptor Expression. Washed human platelets were prepared as described above. PS (3.8 × 10^10 platelets/ml) containing 2 μM PGE1, was incubated with kistomin 20 μg/ml at 37°C for 10 min. After an extensive wash, platelets were labeled with mAb against GP Ibα (SZ2), αIbβ3 (TE3), or α2β1 integrin (6F1) at room temperature (RT) for 30 min. Labeled cells were washed with Tyrode’s solution and then incubated with secondary FITC-conjugated goat anti-mouse IgG (CALTAG Lab, Burlingame, CA) RT for 30 min with a continuous shaking. After incubation, cells were washed, resuspended in phosphate-buffered saline (PBS), and analyzed immediately by FACSCalibur (BD Biosciences, San Jose, CA).

Western Blot Analysis of Platelet GP Ibα. Platelet suspension was centrifuged at 200 g for 5 min. After removing the supernatant, pellets were lysed by 1% Triton X-100 buffer (in PBS). Aliquots of cell lysates and supernatants were resolved on 10% SDS-PAGE under reducing conditions and electrotransferred to Immobilon-PVDF membrane (Millipore). After blocking in a 0.5% BSA in Tris-buffered saline 1 h at 4°C, the blots were probed with anti-GP Ibα mAb (1:1000) for overnight at 4°C and followed by the horseradish peroxidase-goat anti-mouse IgG. The protein was visualized by adding enhanced chemiluminescence (ECL) solution (Pierce, Rockford, IL).

Flow Cytometric Analysis of Platelet Receptor Expression. Washed human platelets were prepared as described above. PS (3.8 × 10^10 platelets/ml) containing 2 μM PGE1, was incubated with kistomin 20 μg/ml at 37°C for 10 min. After an extensive wash, platelets were labeled with mAb against GP Ibα (SZ2), αIbβ3 (TE3), or α2β1 integrin (6F1) at room temperature (RT) for 30 min. Labeled cells were washed with Tyrode’s solution and then incubated with secondary FITC-conjugated goat anti-mouse IgG (CALTAG Lab, Burlingame, CA) RT for 30 min with a continuous shaking. After incubation, cells were washed, resuspended in phosphate-buffered saline (PBS), and analyzed immediately by FACSCalibur (BD Biosciences, San Jose, CA).

Protein Sequencing of the Fragmented Kistomin and cDNA Cloning of Kistomin Precursor. Protein sequencing of the fragmented kistomin was performed as described previously (Wu et al., 2001a). In brief, fragmented kistomin was obtained by alklylation with vinylpyridine and followed by incubation with CNBr. The fragments were applied to high-performance liquid chromatography, and the major fraction was subjected to protein sequencing. For sequencing of the autoproteolytic fragment of kistomin, kistomin was auto-proteolyzed in 1% SDS and 0.5 M Tris-HCl solution, transblotted onto PVDF membrane, and then analyzed by sequencing.

Total RNA was isolated from C. rhodostoma venom glands with a BlueExtract kit (LTK BioLaboratories Co., Ltd, Linko, Taipei, Taiwan), cDNA was synthesized from 1.5 μg of the total RNA and used to construct a cDNA library in the Uni-ZAP XR vector (Stratagene, La Jolla, CA).

To obtain cDNA of putative SVMP, recombinant λ DNA species were prepared from the cDNA library and used as templates in PCR. Primers for the first screening of cDNA library were 5′-TCCATGGAAGTCCGAAAGTGC/CTA/TGTGCTGGA/AA-3′ and 5′-TCAGGTGGGTCT/TT/ GAAGCAGGG-3′. Amplification was performed by using 72°C DNA polymerase (Ab Peptides, St. Louis, MO) and the primers with a hot start at 94°C for 10 min and 30 cycles of denaturation (1 min, at 94°C), annealing (1 min, at 50°C), and extension (2 min, at 72°C).
Cleavage of vWF by Kistomin. Purified human vWF (1.5 μg) incubated with or without kistomin were analyzed as described previously (Wu et al., 2001b) with a minor modification. In brief, aliquots of the mixture were analyzed by SDS-1% agarose gel electrophoresis (~2 mm) in a Mupid-2 Mini-Gel system (Cosmo Bio Co., Ltd, Tokyo, Japan) and electroblotted onto PVDF membrane. Immuno-blotts were developed with peroxidase-conjugated antihuman vWF antibody (DakoPatts, Glostrup, Denmark).

Fluorescent Dye-induced Platelet Thrombus Formation in Mesenteric Microvessels of Mice. Fluorescent dye-induced platelet thrombus formation in mesenteric microvessels of mice was performed as described previously (Chang and Huang, 1994) with some modifications. In brief, after male ICR mice (12~15 g) were anesthetized with sodium pentobarbital (50 μg/g, i.p.), fluorescein sodium (12.5 μg/g) was i.v. injected. A segment of small intestine attached to its mesentery was loosely exteriorized for microscopic observation. Venules with diameters of 30 to 40 μm were selected to produce a microthrombus. In the epi-illumination system, the area of irradiation (wavelength above 520 nm) was approximately 50 μm².

To further determine the cDNA sequence of kistomin precursor, cDNA cloning was performed. To obtain the remaining cDNA sequence of prokistomin contains the characteristic zinc-chelating sequence HEIGHNLGMEHD (Fig. 1, catalytic site), which is similar to that of other SVMPs, such as fibrolase (Randolph et al., 1992) and jararhagin (Paine et al., 1992).

Kistomin Inhibited Ristocetin-induced Platelet Agglutination and Aggregation. To re-examine kistomin’s activity in inhibiting platelet function, PS agglutination and PRP aggregation were performed. As shown in Fig. 2, kistomin concentration-dependently inhibited ristocetin-induced platelet agglutination and aggregation with a half-maximal inhibition concentration (IC50) at 2.04 μg/ml (0.079 μM) and 8.25 μg/ml (0.321 μM), respectively. However, this inhibition was abolished by the treatment of kistomin with EDTA or o-phenanthroline (data not shown), indicating the involvement of an enzymatic reaction.

Kistomin Cleaved Platelet GPIbα. To further elucidate the possible action mechanism of kistomin in inhibiting ristocetin-induced platelet aggregation, GPIbα expression on platelets was analyzed by flow cytometry and Western blotting. As depicted in immunofluorescence staining with anti-GPIbα mAb SZ2, kistomin treatment rapidly reduced the level of GPIbα expression on platelets, whereas the expression of the other two important platelet receptors, namely αIIbβ3 and α2β1 integrins, was not affected (Fig. 3A).

To characterize the proteolytic properties of kistomin on platelet GPIbα, Western blotting was performed. It was found that platelet intact GPIbα (~140 kDa) was cleaved by kistomin in a time-dependent manner, which could be abolished by EDTA (Fig. 3B). One intact GPIbα and two fragments migrated at molecular masses of ~140, ~130, and ~45 kDa, respectively, were detected by anti-GPIbα mAb SZ2 in total platelet lysate (arrows). We were surprised to find that only two fragments (~130 and ~45 kDa) were detected in supernatant (Fig. 3C), indicating that kistomin cleaves platelet GPIbα at two distinct sites to generate two soluble fragments, which can be recognized by the SZ2 mAb.

Binding of Kistomin to GPIbα. Because platelet GPIbα was cleaved by kistomin, we next investigated whether kistomin bound to GPIbα. To stop kistomin’s enzymatic activity, the experiment was performed at 4°C. Under this condition, kistomin bound to GPIbα and replaced anti-GPIbα M45 mAb binding to platelets in a concentration-dependent manner (Fig. 4A). Binding of kistomin to GPIbα concentration-dependently increased and reached saturation at the concentrations more than 20 μg/ml (Fig. 4B). This result was confirmed by the observation that immobilized kistomin directly interacted with GPIbα in platelet lysate. It is noteworthy that this binding was not affected in the presence of EDTA (Fig. 4C), suggesting that bivalent cations are not required in this interaction. A similar binding ability was also found in immobilized agglucetin, a tetrameric GPIbα-binding protein from A. acutus (Wang and Huang, 2001), in which it migrated as two distinct bands at 16.2 and 14.5 kDa (Fig. 4, lane 3). In contrast, rhodostomin, an RGD-containing disintegrin purified from C. rhodostoma venom (Huang et al., 1987), failed to bind platelet GPIbα (data not shown).

Effect of Kistomin on the Multimeric Structure of vWF. We have shown that kistomin could bind and cleave platelet GPIbα (Fig. 3 and 4). To further examine whether kistomin affected multimeric structure of vWF, human vWF preincubated with or without kistomin was added to platelet suspension and ristocetin-induced platelet aggregation was measured. After the operation (15 min), the mouse was immediately immersed vertically in normal saline at 37°C. Bleeding time was recorded from the time bleeding started until it completely stopped.

Results

Protein Sequencing and cDNA Cloning of Kistomin. To have an insight on kistomin’s primary structure, protein sequencing was performed. A 14-kDa autoproteolytic fragment from kistomin was sequenced, and its N-terminal was revealed as LSKRKPHNDAQFLTKDFDG (fragment 1). Moreover, two peptide sequences, VDKHNGNIKKIE (fragment 2) and APEVNNPTKFFSDC (fragment 3), were obtained from CNBr digestion of kistomin.

To further determine the cDNA sequence of kistomin precursor, cDNA cloning was performed. Two primers in accordance with the conserved PKMCGV sequence of SVMP and the autoproteolytic fragment were used in the first amplification of kistomin cDNA. After screening of the C. rhodostoma cDNA library, a clone with approximately ~900 bp was obtained, representing the kistomin precursor sequence containing the partial metalloproteinase domain of kistomin (data not shown). To obtain the remaining cDNA sequence of kistomin, a pair of primers according to the sequences from kistomin-CNBr-digested fragment (VDKHNNGNIKKIE) and vector was used. Figure 1 showed the assembled cDNA sequence and the deduced amino acid sequence of the kistomin precursor. Three partial sequences obtained from direct protein sequencing were found in the deduced amino acid sequence with 100% identity (Fig. 1, underlined sequences), indicating that it is a kistomin precursor. The precursor, designated prokistomin, consists of a prosequence, presequence, and a metalloproteinase domain and belongs to a P-I SVMP. A putative start site was indicated, and mature kistomin predicted from this site was 227 residues (Fig. 1), which was estimated to be 25.7 kDa. The deduced amino acid sequence of prokistomin contains the characteristic zinc-chelating sequence HEIGHNLGMEHD (Fig. 1, catalytic site), which is similar to that of other SVMPs, such as fibrolase (Randolph et al., 1992) and jararhagin (Paine et al., 1992).
measured. Figure 5A showed that ristocetin-induced agglutination was time-dependently reduced under this condition but almost fully restored by re-adding a new intact vWF. Further analysis revealed that high-molecular-mass multimers of vWF obviously decreased and the low-molecular-mass multimers concomitantly increased in the presence of kistomin (Fig. 5B, lanes 1 and 2). Again, the cleavage of vWF by kistomin was abolished in the presence of EDTA or o-phenanthroline (Fig. 5B, lanes 3 and 4). Taken together, our results indicate that kistomin can bind and cleave platelet GPIbα and vWF and subsequently inhibits vWF-induced platelet agglutination and aggregation.

Kistomin Affected Thrombosis and Hemostasis in Vivo. We next investigated whether kistomin exerted anti-thrombotic effect in vivo. It was shown that kistomin potently decreased antibody binding to platelet GPIbα in human whole blood, whereas crotalin and triflamp, two P-I SVMPs purified from venom of C. atrox (Wu et al., 2001b) and T. flavoviridis (Tseng et al., 2004a), respectively, were less effective at the same concentration in cleaving GPIbα (Fig. 6). We therefore hypothesized that kistomin could be an active protease in both in vitro and in vivo conditions. To confirm this hypothesis, we measured ex vivo platelet aggregation in PRP or PS prepared from kistomin-pretreated mouse. Be-

Fig. 1. cDNA sequence and deduced amino acid sequence of the putative protein precursor of kistomin. The sequence begins with the presequence, and the translation stop codon is indicated by an asterisk (*). Three fragments from CNBr-digested and autoproteolytic kistomin are indicated by underlines.
cause ristocetin is ineffective in causing platelet aggregation in mouse PRP, a snake venom-derived GPIIbα agonist, gramicidin, was used in this assay (Wu et al., 2001a). As shown in Fig. 7A, gramicidin-induced platelet agglutination was inhibited by the GPIIbα antagonist, agkistin, suggesting that a GPIIbα-mediated pathway was involved in this agglutination. The agglutination was suppressed in PRP prepared from mouse treated with kistomin (Fig. 7B); however, collagen-, ADP-, convulxin- and thrombin-induced platelet aggregation was not significantly affected (Fig. 7, C–F). Thus, kistomin

![Figure 2](image-url)

**Fig. 2.** Effect of kistomin on ristocetin-induced platelet agglutination and aggregation. Washed PS (A) and PRP (B) were pretreated with various concentrations of kistomin before the addition of ristocetin (1 mg/ml, arrow). Platelet agglutination and aggregation were monitored by turbidimetry in aggregometer. C and D, quantitative analysis of the data from A and B and similar experiments were performed. Data are presented as percentage of control and are mean ± S.E.M. (n = 3).

![Figure 3](image-url)

**Fig. 3.** Effect of kistomin on platelet GPIIbα. A, flow cytometric analysis of GPIIbα expression on platelets. Washed platelets treated with PBS (gray area) or kistomin (5 µg/ml, open area) at 37°C for 10 min were incubated with anti-GPIIbα (SZ2), anti-αIIbβ3 (7E3) or anti-α2β1 (6F1) mAbs and subjected to be analyzed by flow cytometry. B and C, Western blot analysis of GPIIbα expression on platelets. Washed platelets were treated with kistomin (20 µg/ml) at 37°C for (B) different duration as indicated or (C) for 30 min in the absence or presence of EDTA. Total cell lysates, cells pellets (P), and supernatant (S) were obtained as described under Materials and Methods and analyzed by Western blotting. An arrowhead indicates an intact GPIIbα expression on platelet. Note that 130-kDa (open arrowheads) and 45-kDa fragments (arrows) were observed in total cell lysates (B) and in the supernatant (C) of kistomin-treated platelets. This experiment is representative of at least three similar experiments.
impaired mouse platelet function specifically through affecting GPIbα in vivo.

We then determined kistomin’s antithrombotic effects in vivo. In fluorescent dye-treated mice, thrombus formation was observed in irradiated mesenteric venules of mice. The occlusion time of irradiated vessels was $134.3 \pm 6.2$ s in control mice ($n = 12$) but was prolonged to $173.2 \pm 26.7$ s ($n = 13$) and $301.9 \pm 43.5$ s ($n = 10$) by aspirin at doses of 150 and 250 $\mu$g/g, respectively. We were surprised to find that compared with aspirin, kistomin also exerted potent antithrombotic effect in vivo, prolonging the occlusion time to 194.9 $\pm$ 12.5 s ($n = 10$) and 273.2 $\pm$ 16.2 s ($n = 10$) at doses of 1.5 and 7 $\mu$g/g, respectively (Table 1). Kistomin was approximately 4000-fold more potent than aspirin in prolonging microvessel occlusion time on a molar basis. In parallel, the tail bleeding time of mice (control, 90.8 $\pm$ 5.0 s, $n = 17$) was prolonged to $172.1 \pm 26.3$ s and more than 1800 s by kistomin at the given doses of 1.5 and 7 $\mu$g/g, respectively. In contrast, aspirin at 150 and 250 $\mu$g/g also increased bleeding time to 559 $\pm$ 48.9 s ($n = 13$) and more than 1800 s ($n = 10$), respectively (Table 2).

**Discussion**

Platelet GPIbα-vWF interaction has been identified as an important target for therapeutics to prevent ischemic cardio-

vascular events (Jackson and Schoenwaelder, 2003). We have found that kistomin inhibited ristocetin-induced platelet agglutination in platelet suspension (Huang et al., 1993); however, its action mechanism, antithrombotic activity, and structure-activity relationship are still largely unknown. In this study, we demonstrated that kistomin is capable of binding to platelet GPIbα and cleaves GPIbα and vWF, exhibiting potent antplatelet and antithrombotic activities in vitro and in vivo. More importantly, cDNA encoding kistomin precursor was cloned and revealed that mature kistomin is a P-I SVMP with only a metalloproteinase domain. The sequence of kistomin is shown with 51% identity to the P-I SVMP fibrolase, a fibrinolytic enzyme from *Agkistrodon contortrix contortrix* venom (Randolph et al., 1992), and with 40% identity to the P-III SVMP jararhagin, an $\alpha_2$β1- and vWF-cleavage protease from *Bothrops jararaca* venom (Paine et al., 1992). Moreover, kistomin shares 29% identity with the proteinase domain of human ADAMTS13 (a disintegrin and metalloproteinase with thrombospondin motifs 13), an endogenous metalloproteinase specifically cleaving between...

**Fig. 4.** Kistomin binds to platelet GP Ibα. A, kistomin competitively replaced anti-GPIbα mAb binding to platelets. Human washed platelets were incubated with PBS (gray area) or various concentrations of kistomin (open areas, 15, 30, and 60 $\mu$g/ml) at 4°C for 30 min, followed by incubation with FITC-conjugated anti-GPIbα mAb, M45, and then analyzed by flow cytometry. The histogram was representative from three similar experiments. B, quantitative analysis of binding assay data from A and similar experiments were performed. Data were expressed as mean fluorescence and were mean $\pm$ S.E. ($n = 3$) C, platelet GPIbα bound to immobilized kistomin. Kistomin (Kis, 20 $\mu$g) pretreated with vehicle or EDTA (10 mM) and agglutinin (agg, 15 $\mu$g) were applied to SDS-PAGE and transferred to PVDF membrane. The membrane was incubated with platelet total lysate and followed by Western blot analysis using anti-GPIbα mAb, SZ2. This experiment is a representative one of at least three similar experiments.

**Fig. 5.** Effect of kistomin on the multimeric structure of vWF. A, pre-treatment of vWF with kistomin compromised vWF-induced platelet aggregation. Washed human platelets were incubated with kistomin (3 $\mu$g/ml)-pretreated vWF (1 $\mu$g/ml) at 37°C for the indicated times and ristocetin (ris, 1 mg/ml) was added to induce platelet aggregation. In the case of prior coincubation of vWF and kistomin for 20 min, intact vWF (10 $\mu$g/ml, arrow) was readded 3 min after addition of ristocetin. B, kistomin cleaved the multimeric structure of vWF. Human vWF (0.5 $\mu$g) was incubated at 37°C for 30 min with PBS (lane 1), kistomin (15 $\mu$g/ml, lane 2), EDTA (10 mM)-treated kistomin (lane 3), or $\alpha$-phenanthrolene (10 mM)-treated kistomin (lane 4). Aliquots of each reaction mixture were subjected to SDS-1% agarose electrophoresis and vWF multimer were detected by peroxidase-conjugated anti-vWF antibody after blotting to a PVDF membrane. This experiment is representative of at least three similar experiments.
Kistomin inhibits vWF-induced platelet agglutination and aggregation through acting on platelet GPIbα and vWF. Several lines of evidence indicate that the platelet GPIbα and vWF were cleaved by kistomin during their coinoculation. First, a significantly reduced binding signal in flow cytometry was observed after platelets were treated with kistomin (Fig. 3A). Second, the outer membrane portion of GPIbα was cleaved from platelet membrane into supernatant in the presence of kistomin and generated two soluble fragments, which migrated at the molecular masses of 45 and 130 kDa (Fig. 3, B and C). Third, kistomin competitively inhibited anti-GPIbα mAb interaction with platelet and directly bound to platelet GPIbα, as determined by flow cytometry and Western blotting (Fig. 4). Fourth, vWF-induced platelet agglutination was compromised by a preincubation of vWF with kistomin but reversed by adding an intact vWF (Fig. 5A). Fifth, upon kistomin incubation, high molecular mass multimers of vWF generated low molecular mass multimers (Fig. 5B). Last, EDTA pretreatment can abolish these activities of kistomin (Figs. 3, 5, and data not shown), indicating that kistomin is a typical SVMP, mediating antiplatelet activities through cleaving GPIbα and vWF.

Regarding specificity, the surface marker analysis showed that kistomin failed to affect the binding of anti-α2β1 (6F1) and αIIbβ3 (7E3) mAbs to platelets. In contrast, kistomin specifically inhibited the binding of anti-GPIbα mAbs to platelets, including AP1, 6D1, and SZ2 (Fig. 3A and data not shown). Moreover, platelets prepared from kistomin-treated mice were unable to agglutinate in response to GPIbα-agonist induction but were able to aggregate in response to other agonists (Fig. 7), suggesting its relative specificity toward GPIbα both in vitro and in vivo. The GPIbα-binding epitopes for 6D1 and AP1 have been demonstrated to be located at the amino acid residues 104 to 128 and 201 to 268, respectively (Coller et al., 1983). SZ2 mAb has been shown to recognize anionic sulfated tyrosine residues 269 to 282 of GPIbα (Ward et al., 1996). Therefore, failed binding of these Abs to platelets in the presence of kistomin indicates that kistomin may cleave GPIbα, downstreaming the anionic tyrosine sulfated region. In Fig. 3, we found that outer membrane portion of GPIbα was cleaved by kistomin to generate a ~130-kDa soluble fragment. Because it was a soluble fragment found in total cell lysate and in the supernatant (Fig. 3C), the possibility of cleavage of GPIbα at the site near N terminus was excluded. Therefore, the first cleavage site was hypothesized to be located near C terminus of GPIbα. The cytoplasmic tail of GPIbα contains 96 amino acid residues (Berndt et al., 2001) and is estimated to have a molecular mass of approximately 10 kDa. Therefore, the first cleavage site on GPIbα...
may be located near the outer membrane of platelet. Second, a ~45-kDa soluble fragment increased gradually accompanying with a decrease of the ~130-kDa fragment (Fig. 3), suggesting that kistomin’s second cleavage site is located within the ~130 kDa GPIbα fragment. This is evidenced by the observations that inactivated kistomin competitively replaced the binding of anti-GPIbα M45 mAb, which recognizes anionic sulfated tyrosine residues of GPIbα (Fig. 4). Kistomin seems to act like mocarhagin, a P-III SVMP, by cleaving GPIbα at a single site between Glu282 and Asp283 to generate a ~40-kDa fragment. However, this ~45-kDa fragment could be recognized by SZ22 on Western blotting analysis (Fig. 3, B and C), indicating that the binding epitope of SZ2, anionic sulfated tyrosine residues of GPIbα, still remained on the fragment. According to the size of the second fragment (~45kDa) and the binding epitopes of anti-GPIbα mAbs (SZ2 and M45), we postulated that the second kistomin-cleavage site on GPIbα is near downstream of the anionic sulfated tyrosine region. Taken together, we suggest that kistomin cleaves GPIbα at two distinct sites, one of which is located at the region near the outer membrane and another locates near anionic sulfated tyrosine. Kistomin’s exact cleavage sites on GPIbα are still under investigation in our laboratory.

Our study also demonstrated that kistomin potentially cleaved platelet GPIbα in human whole blood, compared with crotalin and triflamp, two P-I SVMPs (Fig. 6). It has been shown that human α2-macroglobulin and mouse macroglobulin are abundant in serum and capable of inhibiting and neutralizing the proteolytic activity of most proteinases, including some SVMPs (Tseng et al., 2004b). However, in this report, kistomin was demonstrated to elicit its antiplatelet and antithrombotic in vivo (Fig. 7 and Tables 1 and 2). These suggest that kistomin is less susceptible to be neutralized by globulins in serum and possibly can be developed as an antithrombotic agent. This is supported by the data shown in

### TABLE 1

Effect of kistomin on fluorescent dye-induced thrombus formation in mesenteric venous mice of rats

<table>
<thead>
<tr>
<th>Values are presented as means ± S.E.M. of experimental number (n) indicated.</th>
<th>Occlusion Time n</th>
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<tbody>
<tr>
<td>Control (PBS)</td>
<td>134.3 ± 6.2</td>
</tr>
<tr>
<td>Aspirin 150 μg/g</td>
<td>172.6 ± 6.3***</td>
</tr>
<tr>
<td>250 μg/g</td>
<td>287.4 ± 15.2***</td>
</tr>
<tr>
<td>Kistomin 1.5 μg/g</td>
<td>194.9 ± 12.3**</td>
</tr>
<tr>
<td>7 μg/g</td>
<td>273.2 ± 16.2***</td>
</tr>
</tbody>
</table>

*** P < 0.001 compared with control.

### TABLE 2

Effect of kistomin on the tail bleeding time of mice

<table>
<thead>
<tr>
<th>Values are presented as means ± S.E.M. of experimental number (n) indicated.</th>
<th>Tail Bleeding Time n</th>
</tr>
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<tbody>
<tr>
<td>Control (PBS)</td>
<td>90.8 ± 5.1</td>
</tr>
<tr>
<td>Aspirin 150 μg/g</td>
<td>559.0 ± 48.5***</td>
</tr>
<tr>
<td>&gt;1800.0***</td>
<td></td>
</tr>
<tr>
<td>Kistomin 1.5 μg/g</td>
<td>172.1 ± 26.3**</td>
</tr>
<tr>
<td>&gt;1800.0***</td>
<td></td>
</tr>
</tbody>
</table>

*** P < 0.001 compared with control.

### References


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