The disulfide isomerase ERp57 mediates platelet aggregation, hemostasis, and thrombosis

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A close homologue to protein disulfide isomerase (PDI) called ERp57 forms disulfide bonds in glycoproteins in the endoplasmic reticulum and is expressed on the platelet surface. We generated 2 rabbit Abs to ERp57. One Ab strongly inhibited ERp57 in a functional assay and strongly inhibited platelet aggregation. There was minimal cross-reactivity of this Ab with PDI by Western blot or in the functional assay. This Ab substantially inhibited activation of the αIIbβ3 fibrinogen receptor and P-selectin expression. Furthermore, adding ERp57 to platelets potentiated aggregation. In contrast, adding a catalytically inactive ERp57 inhibited platelet aggregation. When infused into mice the inactive ERp57 prolonged the tail bleeding times. We generated 2 IgG2a mAbs that reacted with ERp57 by immunoblot. One of these Abs inhibited both ERp57 activity and platelet aggregation. The other Ab did not inhibit ERp57 activity or platelet aggregation. The inhibitory Ab inhibited activation of αIIbβ3 and P-selectin expression, prolonged tail bleeding times, and inhibited FeCl3-induced thrombosis in mice. Finally, we found that a commonly used mAb to PDI also inhibited ERp57 activity. We conclude that a glycoprotein-specific member of the PDI family, ERp57, is required for platelet aggregation, hemostasis, and thrombosis. (Blood. 2012;119(7):1737-1746)

Introduction

The prototypic disulfide isomerase (PDI) was discovered approximately 50 years ago as an enzyme that forms disulfide bonds in nascent proteins in the endoplasmic reticulum (ER). Extracellular PDI is now known to mediate platelet aggregation and thrombus formation in vivo. ERp5 is another member of the PDI family that has a role in platelet aggregation. ERp57 is a member of the PDI family of enzymes that is also found in platelets. On platelet activation, ERp57 is secreted by platelets and recruited to the platelet surface. ER substrates of ERp57 are mostly heavily glycosylated disulfide-bonded proteins that include integrin subunits (α2, α3, α6, β1, β3). The role of ERp57 in platelet function is unknown.

ERp57 interacts cotranslationally with the ER lectins calnexin and calreticulin in the folding of glycoproteins and has been studied for its role in the assembly of the MHC class 1 molecules. While it was thought that ERp57 requires the chaperones calnexin and calreticulin to gain access to its substrates, not all ERp57-dependent disulfide bond formation in the ER requires calnexin and calreticulin.

ERp57 is of similar size and domain structure to PDI with 33% identity in its amino acid sequences. ERp57 has 505 amino acids while PDI has 508 with the catalytic a and a’ domains sharing 50% amino acid identity. Like PDI it contains 2 CGHC active-site sequences and catalyzes the reversible oxidation of thiols to disulfides and the isomerization of disulfide bonds. The substrate binding b and b’ domains of PDI and ERp57 domains share only approximately 20% amino acid identity and the small C-terminal region of ERp57 contains multiple lysine residues whereas in PDI, the region is highly acidic.

In the current studies, we raised both rabbit and mAbs to ERp57 to study the role of this enzyme in platelet function and hemostasis. Because of the high homology between ERp57 and PDI, we examined the ability of these Abs to inhibit PDI. Using Abs that preferentially inhibited ERp57, we document a role for ERp57 in platelet aggregation, hemostasis, and thrombosis.

Methods

Materials

Glutathione (GSH), glutathione disulfide (GSSG), DTT, Sephadex G-25, normal rabbit IgG, and a monoclonal mouse IgG2a (raised against pristane) were purchased from Sigma-Aldrich. The IgG2a mouse monoclonal anti-PDI RB.90 was purchased from Affinity BioReagents. The thrombin receptor agonist peptide SFLLRN was synthesized as previously described and was also synthesized by Enzo Life Sciences.

Ab generation

The cDNA for ERp57 was provided by Dr Peppi Koivunen (University of Oulu, Oulu, Finland). The cDNA was subcloned into a pTRIEX-4 Neo vector (Novagen) containing an N-terminal histidine tag. ERp57 was expressed in Escherichia coli strain BL21 (DE3)pLysS (Promega) and purified on a Ni Sepharose High-Performance column (GE Healthcare). The cDNAs for PDI and ERp57 were obtained from Dr M. Kikuchi (Ritsumeikan University, Kyoto, Japan), respectively. PDI and ERp57 were prepared using identical methods as for ERp57. Polyclonal Abs to ERp57 were raised in 2 New Zealand white rabbits by the GenScript Company, and affinity purified against an ERp57 column and a protein A column. Abs were raised to human ERp57 in mice by the GenScript Company. Hybridomas expressing anti-ERp57 mAbs were identified by Western blot and the mAbs were purified using a protein A column. The Abs were
screened for inhibition of platelet aggregation, and an IgG2a Ab (3G1; hereinafter called Mab1) was found that inhibited platelet aggregation. Another IgG2a Ab (1B12; hereinafter called Mab2) reacted with Erp57 by Western blot but did not inhibit platelet aggregation. This Ab was used as a control and compared with the isotype-specific IgG2a control Ab.

**Preparation of the PDI substrate, Di-E-GSSG**

Dioscin glutathione disulfide (Di-E-GSSG) prepared by the method of Raturi and Mutus was kindly provided by Dr Mutus (University of Windsor, Windsor, ON). To make this PDI substrate, 2 esoin moieties are attached to 2 free N termini of GSSG. PDI reductase activity cleaves the SS bond of Di-E-GSSG resulting in the release of 2 esoin-GSH (EGSH) molecules and an increase in fluorescence.

**Preparation of EGSH and an EGSH standard curve**

To convert the fluorescent intensity of a sample to the amount of esoin-GSH (EGSH) generated, we used a standard curve of the fluorescent intensities of different EGSH concentrations. EGSH was prepared as described by treating Di-E-GSSG with DTT and separating the mixture with a Sephadex G-25 column. The eluted sample did not show any increase in the fluorescence after the addition of DTT (10 mM), confirming that the Di-E-GSSG has been converted to EGSH. The aliquots were pooled together and quantified using molar absorption coefficient \( \varepsilon = 88 \times 10^3 \) M\(^{-1}\) cm\(^{-1}\). The increase in fluorescence was monitored as a function of [EGSH] (\( \varepsilon = 88 \times 10^3 \) M\(^{-1}\) cm\(^{-1}\)). The standard plot was generated with excitation at 525 nm and emission at 545 nm and used to quantify the reduction of [Di-E-GSSG] to [EGSH].

**Assay of enzyme-dependent disulfide reduction**

Erp57, PDI, and Erp5 disulfide reductase activities against Di-E-GSSG were monitored in PDI assay buffer (0.1 M potassium phosphate buffer, 2 mM EDTA, pH 7.0) by adding Erp57, PDI, or Erp5 (20 nM) to Di-E-GSSG (150 nM) in the presence of 5 mM DTT. The increase in fluorescence was monitored at 545 nm with excitation at 525 nm. The degree of inhibition was relative to the total amount of EGSH formed over the time period of the assay or by the initial velocity of the reaction (nM EGSH formed per minute).

**Western blotting**

Western blotting was performed using the Odyssey Western blot kit according to the manufacturer’s instructions and developed using the Odyssey Infrared Imaging System (Li-Cor Biosciences).

**Preparation of catalytically inactive Erp57**

A catalytically inactive mutant Erp57, with the 4-cysteine residues of both active sites mutated to serine residues (C57S, C60S, C406S, and C409S) was prepared using the QuikChange Site-Directed Mutagenesis Kit (Stratagene). The mutant Erp57 is also designated as Erp57 (oo-oo; representing the Cys/SH change to Ser/OH in the 2 CGHC active sites). DNA sequencing confirmed the correct base substitutions.

**Platelet isolation and platelet aggregation**

All studies on human platelets were performed after approval by the Temple University Institutional Review Board. Platelets were prepared as described previously. Briefly, 8.6 mL of blood was drawn into 1.4 mL of acid-citrate dextrose solution, and the platelet-rich plasma, obtained after centrifugation of whole blood, was gel-filtered on a Sepharose 2B column equilibrated in Tyrode-albumin solution at pH 7.35. Calcium (1 mM) was then added to the platelets. Platelet aggregation was as described using the Chronolog lumaggregometer. The percentage of aggregation was calculated from the amplitude of the tracings and normalized to the response of the IgG control within an individual experiment.

**Flow cytometric studies of platelets**

Flow cytometric analysis was performed as described. Briefly, citrated platelet-rich plasma was diluted into an equal volume of PBS and incubated for 10 minutes with the rabbit anti-Erp57 IgG (30 \( \mu \)g/mL) or normal rabbit IgG and then with the SFLRN peptide for 5 minutes without stirring at room temperature. In some samples, the anti-Erp57 Ab was added after the SFLRN peptide had been incubated with the platelets. In other studies, the inhibitory monoclonal anti-Erp57 Ab (30 \( \mu \)g/mL) was used. A total of 10 \( \mu \)L of FITC-conjugated PAC-1 (BD Biosciences) was then incubated with 10 \( \mu \)L of platelets for 30 minutes in the dark. In some experiments, convulxin (Enzo Life Sciences) was used as the agonist and PE-conjugated anti-P-selectin Ab (eBioscience) was incubated with the platelets. To study surface expression of Erp57 on platelets, 20 \( \mu \)g/mL rabbit anti-Erp57 Ab or normal rabbit IgG was incubated with nonstimulated platelets for 10 minutes. FITC-labeled goat anti–rabbit Ab was then added for 10 minutes. The cells were fixed with 2% paraformaldehyde. Ten thousand platelets per sample were analyzed by flow cytometry. The data were combined to obtain the mean with the fluorescent intensity of Ab binding to nonactivated platelets subtracted out.

**Tail bleeding times**

Bleeding times were determined as previously described with some modifications. Mice were obtained from Charles River Laboratories and were performed in accordance with institutional guidelines using IP pentobarbital for anesthesia. Male mice (C57BL/6, 6-8 weeks old) were anesthetized and recombinant mutant Erp57 (oo-oo; 100 \( \mu \)g/mouse) in a volume of 100 \( \mu \)L of PBS, PBS only, or mAbs (200 \( \mu \)g/mouse) were injected through the iliac vein. After 5 minutes, the mouse tail was transected 5 mm from the tip with a razor blade. Because of the relatively long tail bleeding times with this method, the procedure was revised to transect the mouse tail 3 mm from the tip. The bleeding tail was immersed in a 12-mL test tube containing PBS, which was warmed in a water bath at 37°C. Bleeding times were determined when the bleeding stopped for > 10 seconds. In initial experiments, bleeding times exceeding 30 minutes, and in a second set of experiments 15 minutes, were noted as 30 or 15 minutes, respectively.

**Studies using FeCl3-induced thrombosis of the carotid artery**

Male mice weighing 20 to 25 g (C57BL/6 age 6-8 weeks) were anesthetized using sodium pentobarbital 120 mg/kg by IP injection as described elsewhere with some modifications. A midline incision was made in the neck, and the left carotid artery was surgically exposed by blunt dissection. A 1 × 2 mm patch of no. 1 Whatman filter paper, soaked in 5% FeCl3, was applied under and above the exposed artery for 2 minutes. After removal of the filter paper, the artery was washed with PBS and an imaging ultrasound gel (MS400-0090; VisualSonics) was placed in the surgical wound to allow Doppler monitoring. The artery was identified using a small animal blood flow transducer (MS400, 18-38 MHz; VisualSonics) and the color Doppler mode of the VisualSonics Vevo model 2100 flowmeter. The transducer was fixed in place using the Vevo Imaging Station (Integrated Rail System), a bench mounted adjustable rail system custom designed for small animal positioning and imaging. Mouse temperature was maintained at 37°C. Baseline systolic and diastolic blood velocity (mm/s) was recorded using the two-dimensional (2D) imaging Doppler mode. The occlusion time was defined as the time from the initiation of arterial injury to complete occlusion of blood flow. An occlusion was determined to be stable when the flow completely stopped for at least 5 minutes. Four hundred fifty micrograms of the anti-Erp57 Ab or control IgG2a were injected before the FeCl3 injury through the right common iliac vein as described elsewhere. The operator was blinded to the Ab infused while performing all experiments.

**Flow cytometric studies on mouse platelets after infusion of Erp57**

Mice were anesthetized and recombinant mutant Erp57 (oo-oo; 100 \( \mu \)g/mouse) in 100 \( \mu \)L of PBS or PBS only was injected using an insulin syringe into the iliac vein. After 5 minutes, 180 \( \mu \)L of blood was drawn into 20 \( \mu \)L of acid-citrate dextrose solution through the inferior vena cava as described elsewhere using a 21-gauge needle. Platelet-rich plasma (PRP) was made.
by centrifugation of whole blood at 80g for 10 minutes as described elsewhere. The PRP obtained was diluted into 2 volumes of Tyrode buffer. Convulxin (500 ng/mL) was added for 5 minutes and 1/100000 L of the PE-labeled JON/A Ab that recognizes activated mouse IIb3 (Emfret) or PE-conjugated anti–P-selectin (eBioscience) was then incubated with 10000000 L of platelets for 30 minutes in the dark. The platelets were fixed and analyzed as with human platelets.

**Statistics**

For bleeding time experiments, data were statistically analyzed by ANOVA (nonparametric) using GraphPad Prism software. If ANOVA conveyed a P < .05, a Dunn test was performed to further assess significance. The Student t test was used to calculate P values for differences in PAC1 binding and aggregation results. Differences were considered significant at P ≤ .05.

**Results**

**Characterization of ERp57, PDI, and ERp5**

The disulfide isomerase enzymes PDI and ERp5 have a role in platelet aggregation,23-25 ERp57, the closest homologue to PDI,25 is recruited to the platelet surface and secreted with platelet activation. Using the cDNAs for human ERp57, PDI, and ERp5 subcloned into a vector, we expressed the 3 proteins and purified them using a nickel column. ERp57, PDI, and ERp5 of appropriate sizes were seen on a gel (Figure 1A). ERp57 had greater disulfide reductase activity in the GSSG assay than PDI or ERp5 (Figure 1B) with initial velocities of 40, 20, and 16 nM EGSH/min, respectively.
Characterization of rabbit anti-ERp57 Abs by Western blot

The anti-ERp57 Abs raised from the 2 rabbits (Ab 1 and Ab 2) gave single bands against platelets by Western blot (Figure 1C). There was no cross-reactivity of Ab 1 to PDI (or ERp5-not shown) by Western blot (Figure 1C lane 2). Ab 2 had cross-reactivity to PDI by Western blot. Ab 1 provided the greatest inhibition (∼84% of the initial velocity) of ERp57 activity in the Di-E-GSSG assay (Figure 1D).

Characterization of rabbit anti-ERp57 Abs in the Di-E-GSSG assay

Dose response studies of the Abs against ERp57 showed that 30 μg/mL of either Ab provided greater inhibition than 10 μg/mL (Figure 2A). Because immunoreactivity to an epitope may be lost under reduced denaturing gel conditions but the Ab may still be inhibitory in a functional assay, we also tested the Abs for inhibition of PDI. There was minimal inhibition at 10 μg/mL by Ab 1 and a relatively small amount of inhibition of PDI activity at 30 μg/mL (23% of the initial velocity). Ab 2 inhibited more of the PDI activity (50% of the initial velocity using 30 μg/mL; Figure 2B). Therefore, Abs 1 and 2 have differential inhibitory activities against ERp57 and PDI. Ab 1 has substantially more inhibitory activity against ERp57 and substantially less inhibitory activity against PDI than Ab 2.

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ERp5 is another member of the PDI family that also has a role in platelet aggregation, but has much less homology to PDI than ERp57. There was no inhibition of ERp5 in the functional assay with the anti-ERp57 Abs (Figure 2C).
Anti-ERp57 Abs inhibit platelet aggregation

The differential inhibition of the 2 Abs against ERp57 and PDI provides a useful tool to test the role of ERp57 in platelet function.

The Ab that strongly inhibits ERp57 (Ab 1) also strongly inhibits aggregation to both collagen and SFLLRN (Figure 3A-B). The Ab that more weakly inhibits ERp57 (Ab 2) also weakly inhibits platelet aggregation. This is the Ab that cross-reacts with PDI by...
Western blot (Figure 1C lane 2) and has the greatest inhibition of PDI (Figure 2B). Although PDI has a role in platelet aggregation, the anti-PDI effect of Ab 2 is evidently insufficient to substantially inhibit platelet aggregation (Figure 3A-B). Therefore, the inhibition of platelet aggregation by the Ab that strongly inhibits ERp57 is likely primarily by inhibition of ERp57.

**Anti-ERp57 Abs inhibit activation of αIIBβ3**

Flow cytometric analysis showed binding of the monoclonal Ab 1 to resting platelets (Figure 3C). The anti-ERp57 Ab 1 substantially inhibited activation of the αIIbβ3 fibrinogen receptor measured by binding of an activation-specific Ab, PAC-1 (Figure 3D). This was true only if the anti-ERp57 Ab was added before the SFLLRN-agonist (Figure 3Di) but not if it was added after the SFLLRN agonist (Figure 3Dii). This indicates that inhibition of activation of αIIbβ3 by anti-ERp57 is not by steric effects on PAC-1 binding but by inhibition of ERp57. The fluorescent intensity of PAC-1 binding was inhibited by over 50% by anti-ERp57 compared with the IgG control (Figure 3Dii). Ab 1 also had an inhibitory effect on P-selectin expression (Figure 3Ei-ii) a measure of α-granule secretion.

**Wild-type ERp57 potentiates platelet aggregation while the catalytically inactive ERp57 inhibits platelet aggregation and prolongs the tail bleeding time in mice**

As expected, the catalytically inactive mutant ERp57 had no activity in the Di-E-GSSG assay (Figure 4A). We tested the effect of adding wild-type ERp57 or the catalytically inactive ERp57 to human platelets on platelet aggregation. Using concentrations of collagen and the SFLLRN peptide that induced submaximal aggregation, wild-type ERp57 potentiates platelet aggregation while the inactive mutant ERp57 inhibited aggregation (Figure 4B-C). The mean tail bleeding times of mice infused with the mutant ERp57 were prolonged by > 2-fold compared with mice infused with the buffer or wild-type controls (Figure 4D). Mouse platelets prepared from blood collected 5 minutes after infusing the inactive ERp57 showed inhibition of αIIbβ3 activation and P-selectin expression (Figure 4Ei-ii).

**Characterization of 2 monoclonal anti-ERp57 Abs by Western blot and the GSSG assay**

We raised mAbs to ERp57 and characterized 2 IgG2a Abs. Both Abs reacted by Western blot with ERp57 in platelet lysate and with purified ERp57 but not with purified PDI (Figure 5A). However, only one of the 2 Abs inhibited ERp57-dependent conversion of Di-E-GSSG to EGSH in a dose-dependent manner (inhibiting ~64% of the initial velocity at 30 μg/mL; Figure 5B). This Ab did not inhibit PDI or ERp5 in the GSSG assay (Figure 5C-D).

**Differential effect of the inhibitory anti-ERp57 Ab (Mab1) compared with the noninhibitory Ab (Mab2) on platelet aggregation and tail bleeding times in mice**

When we studied the 2 anti-ERp57 Abs in platelet aggregation, the inhibitory anti-ERp57 Ab (Mab1) inhibited aggregation to collagen, SFLLRN, and calcium ionophore (Figure 6A-B). The P2Y12 ADP receptor contains 2 thiols that are potentially redox reactive making it a possible target of ERp57. In the calcium ionophore studies, we excluded this possibility by removing the P2Y12 receptor (Figure 6A). We used a sensitive doppler flow probe that detects systolic and diastolic flow to measure tail bleeding times. The inhibitory anti-ERp57 Ab Mab1 prolonged tail bleeding times compared with the tail bleeding times in mice infused with the control mouse IgG2a or the noninhibitory anti-ERp57 Ab (Mab2; Figure 6E).
diastolic blood flow through the mouse carotid artery (Figure 6Fi). Blood flow ceases when complete occlusion occurs (Figure 6Fii). The time to complete occlusion after the FeCl3 injury was prolonged by infusion of Mab1 relative to normal mouse IgG2a (Figure 6Fiii).

The monoclonal anti-PDI Ab RL90 cross-reacts with ERp57 in the Di-E-GSSG assay

We have developed a mAb that inhibits ERp57 and does not cross-react with PDI. A commercially available mAb called RL90 has been used to demonstrate the role of PDI in platelet aggregation24,27 and thrombosis and hemostasis.5,6 Because of the high homology of ERp57 with PDI, we tested this Ab to PDI for cross-reactivity with ERp57. RL90 did not cross-react with ERp57 by Western blot (Figure 7A). As expected, RL90 inhibited PDI activity (Figure 7B) giving 61% inhibition of initial velocity at 30 μg/mL. These findings are in line with previous studies where RL90 was reported to provide up to 50% to 60% inhibition of PDI.5,29 However, RL90 also inhibited ERp57 activity (Figure 7C) giving ~39% inhibition of initial velocity at
Figure 7. The monoclonal anti-PDI Ab RL90 cross-reacts with ERp57 in the Di-E-GSSG assay. (A) Lack of cross-reactivity by Western blot to ERp57. (B) The inhibition of PDI. (C) The inhibition of ERp57 by RL90, at 10 μg/mL, and 30 μg/mL relative to an isotype specific monoclonal mouse IgG2a Ab (30 μg/mL). 60 μg/mL did not provide more inhibition of PDI or ERp57 with the curves overlapping with the curves for 30 μg/mL. The amount of EGSH formed over 120 seconds from Di-E-GSSG is shown with the enzyme added at 20 seconds.

Discussion

Although discovered relatively recently, ERp57 has quickly become one of the better-studied members of the PDI family.8 ERp57 catalyzes isomerization of disulfide bonds in glycoproteins and we hypothesized that extracellular ERp57 had a role in platelet aggregation, a process that is dependent on the cysteine-rich αIIbβ3 glycoprotein. PDI has a role in platelet function4 and has homologous and structural similarity with ERp57.30 We therefore needed to document that inhibition of platelet function by the Abs against ERp57 was not because of inhibition of PDI. Using 2 rabbit Abs raised to ERp57 we found that the best inhibitory Ab to ERp57 (Ab 1) only weakly inhibited PDI. It is possible that a small amount of the inhibition of platelet aggregation by Ab 1 in Figure 3A and B is because of inhibition of PDI. However, it is apparent that only Ab 2 reacts with PDI by Western blot (Figure 1C) and more strongly inhibits PDI (Figure 2B) yet only weakly inhibits aggregation. Because Ab 1 inhibits PDI only weakly, we conclude that it primarily inhibits platelet aggregation by inhibiting ERp57.

Using a polyclonal rabbit Ab to ERp57, we found a low but statistically significant level of ERp57 on the surface of resting platelets. Resting platelets display a relatively low level of active PDI on the platelet surface.27,31 Similarly, only low levels of ERp5 are expressed on resting platelets.7 A limited surface expression of the PDI enzymes has been proposed as a way of “setting the gain” or threshold for platelet activation.7 In principle, a small amount of enzyme could activate a large number of receptors. The surface amount of active PDI increases dramatically with thrombin stimulation of the platelets31; collagen-induced platelet activation also increases the active form of PDI on the platelet surface.27 In preliminary studies (using polyclonal Abs and mAbs), we have found that the level of ERp57 on the surface increases substantially with activation as previously reported.9 In vivo, other sources of ERp57 (such as ERp57 secreted from endothelial cells) at the site of vascular injury could increase the local concentration of ERp57 at the platelet surface further enhancing receptor activation.

To further document a role for extracellular ERp57 in platelet function, we examined the effect of wild-type ERp57 and a mutant ERp57 rendered catalytically inactive on platelet aggregation. The wild-type enzyme potentiated aggregation while the inactive enzyme had the opposite effect, further suggesting that ERp57 on the platelet surface has a role in platelet aggregation. These findings imply that the added ERp57 interacts with a platelet surface substrate of ERp57 to potentiate platelet aggregation. On the other hand, inactive ERp57 appears to compete with normal platelet surface ERp57 resulting in inhibition of aggregation. The prolongation of tail bleeding times in mice by infusion of catalytically inactive ERp57 suggests a role for circulating or intravascular ERp57 in hemostasis. Platelets collected after infusion of inactive ERp57 show inhibition of activation of αIIbβ3 and P-selectin expression. This suggests that some infused ERp57 binds to the platelet surface in vivo and continues to inhibit the platelets ex vivo. This may suggest a role for platelet ERp57 in the hemostatic mechanisms of tail bleeding times.

Most reports on the role of PDI in platelet aggregation and hemostasis were performed using only one inhibitory Ab3,5,6,24,27 although the role of PDI in regulation of the α2β1 collagen receptor was studied using both a polyclonal and mAb.32 Cross-reactivity of the anti-PDI Abs with ERp57 was not tested. Bacitracin has also been used to inhibit PDI5,24; however, this is not a specific inhibitor of PDI.33 We wanted to confirm a role for ERp57 using a third inhibitor specific for ERp57 and therefore generated 2 IgG2a mAbs to ERp57. Although both Abs reacted with ERp57 by Western blot, only one Ab inhibited ERp57 activity. This allowed us to use the noninhibitory Ab as a control. The inhibitory Ab did not react with PDI or inhibit PDI or ERp5 activity implying that there is no functional cross-reactivity with these enzymes. This is especially important in the case of PDI because there is 50% identity in the amino acid sequence of the active site containing a and a’ domains of the 2 enzymes.14

Using these Abs, we found that the inhibitory anti-ERp57 Ab inhibited platelet aggregation while the noninhibitory control anti-ERp57 Ab did not. This confirms a role for platelet surface ERp57 in platelet aggregation. The inhibitory Ab also prolonged the tail bleeding times in mice confirming a role for intravascular ERp57 in hemostasis. Furthermore, the inhibitory Ab prolonged the time to occlusion in a FeCl3-induced thrombosis model. This implies that intravascular ERp57 has a role in thrombosis.

The high homology between ERp57 and PDI raises the possibility that some of the inhibition observed by Abs against PDI may actually represent inhibition of ERp57. The original rabbit anti-PDI Ab we used to document a role for PDI in platelet aggregation reacted strongly and apparently specifically with PDI3,34 suggesting the major part of the inhibition effect was on PDI. However, we did not at the time test this Ab for cross-reactivity to ERp57. In subsequent inhibition studies the monoclonal anti-PDI Ab RL90 was also found to inhibit platelet aggregation3,24,27 indicating a role for the traditional PDI in platelet

30 μg/mL. Thus, a commonly used mAb to PDI has some cross-reactivity with ERp57 in a functional assay.
aggregation. However, we found some cross-reactivity of monoclonal anti-PDI Ab RL90 with ERp57 in our functional assay. It is therefore possible that previously reported effects of RL90 are due in part to inhibition of ERp57.

The fact that a specific inhibitory monoclonal anti-ERp57 Ab inhibits platelet aggregation does not exclude a role for the traditional PDI (or other PDIs recruited to the platelet surface) in platelet aggregation and hemostasis. It may be that both PDI and ERp57 have important roles in platelet aggregation. For example, numerous proteins interact with the cytoplasmic tails of αIIbβ3 and inhibition of the loss of the interaction of some of the proteins within the αIIbβ3 tail on an individual basis results in complete inhibition of platelet aggregation or thrombus formation. The selective loss of the Kindlin3 interaction with the cytoplasmic tail of β3 results in the complete loss of the aggregation response in mouse or human platelets. Similarly, loss of the interaction of talin with the cytoplasmic tail of β3 also results in the complete loss of aggregation and thrombus formation. These findings imply that both Kindlin3 and talin are important. Our findings with respect to inhibition by Abs to PDI and ERp57 also suggest that both are important and they work by different or separate mechanisms.

The substrate or target for ERp57 is not yet known. Because anti-ERp5 inhibits aggregation induced by several different agonists including calcium ionophore, a surface agonist receptor is unlikely the target. We performed some platelet aggregation experiments in which the dithiol containing P2Y12 receptor was excluded as the mechanism. Other possible targets of ERp57 in platelet aggregation include αIIbβ3 or the fibrinogen added to support platelet aggregation. Because fibrinogen was not added to the flow cytometric assay, the inhibition of PAC-1 binding by the anti-ERp57 Abs implies that fibrinogen is not the target of ERp57. Functional and physical associations of PDI and ERp5 with αIIbβ3 have been noted, raising the possibility that αIIbβ3 is a target of ERp57. This would be consistent with the proposed role for disulfide exchange in αIIbβ3 and disulfide isomerases in the final steps of physiologic agonist-induced activation of integrins. Extracellular ERp57 may have a role in conformational changes that occur in αIIbβ3 as this receptor transitions from a low-affinity conformation to a high-affinity ligand-binding conformation.

Conceivable mechanisms by which both ERp57 and PDI could be required for aggregation include the sequential catalysis of different reactions. For example, one enzyme could catalyze a required disulfide reduction or cleavage reaction thus allowing the other enzyme to catalyze a thiol disulfide exchange reaction. Alternatively, PDI and ERp57 could catalyze simultaneous reactions at different agonists sites of αIIbβ3. Interestingly, a recent report found that a “PDI family member” was required for the unfolding of a polymavirus protein. Specifically, PDI and ERp57 reduced the viral protein and ERp57 isomerized the protein in a process that required free thiols. Thus, in platelet reactions catalyzed by both ERp57 and PDI could be essential with inhibition of either enzyme leading to loss of activation of αIIbβ3. With platelet reactions catalyzed by both ERp57 and PDI could be essential with inhibition of either enzyme leading to loss of activation of αIIbβ3. With platelet activation or aggregation an increase in thiols in PDI is found suggesting that the thiol form of the active site is generated. It is this form that catalyzes disulfide reduction and isomerization. Although ERp57 catalyzes similar reactions as PDI, substantially more work is needed to determine the status of the active sites of ERp57 and the reactions it catalyzes in platelets. These interactions need to be defined, as little is currently understood.

We found that inhibition of ERp57 by rabbit and monoclonal anti-ERp57 Abs decreased convulxin-induced P-selectin expression on platelets. This suggests that both ERp5 and ERp57 have a role in P-selectin exposure. Jordan et al also found a role for PDI in P-selectin expression although Lahav et al documented a lesser role for PDI. The interrelationships of PDI, ERp57, and ERp5 in P-selectin expression will need further characterization.

The role of platelet ERp57 in vivo will require further study. In regard to PDI, it is known that PDI is found on platelet microparticles from patients and that platelets release PDI in vivo at the site of vascular injury. A laser-induced injury model found that endothelial cells could provide the extracellular PDI needed for thrombosis, although platelet microparticles were also a potential source of PDI contributing to thrombosis. Although this injury model provides evidence of a role for endothelial cell PDI in thrombosis, the issue of a contribution from platelet PDI was not addressed. The relative contribution of ERp57 from endothelial cells and platelets or other sources of ERp57 needs to be studied. Eventually, targeted knockouts of PDI and ERp57 in megakaryocytes/platelets will help define the platelet contribution of these PDIs in hemostasis and thrombosis.

In summary, using 3 different reagents we have found that ERp57 on the platelet surface mediates platelet aggregation. ERp57 is also required for hemostasis and thrombosis. Although the degree to which the different members of the PDI family have unique versus overlapping roles in platelet function remains to be further elucidated, in these studies the inhibitory Abs to ERp57 do not significantly inhibit PDI. Importantly, the newly generated inhibitory mAb to ERp57 does not show antigenic or functional cross-reactivity with PDI. Therefore, ERp57 appears to have a role in activation of the αIIbβ3 integrin and platelet aggregation that is distinct from the role of PDI.

Note added in proof: It has come to our attention after our manuscript was accepted for publication that similar findings with respect to the role of platelet ERp57 have been obtained by Holbrook et al, whose work is being published in the Journal of Thrombosis and Haemostasis.

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Authorship

Contribution: Y.W. helped with the design and supervised experiments; Y.W. and S.S.A. collected, analyzed, and interpreted data, and helped revise parts of the manuscript; Y.W., S.S.A., J.Z., L.W., and M.P.C. performed research; and D.W.E. designed the project, analyzed the data, supervised the research, and wrote the manuscript.

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The disulfide isomerase ERp57 mediates platelet aggregation, hemostasis, and thrombosis

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