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Distal hinge of plasminogen activator inhibitor-I involves its latency transition and specificities toward serine proteases

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Abstract

Background: The plasminogen activator inhibitor-I (PAI-I) spontaneously converts from an inhibitory into a latent form. Specificity of PAI-I is mainly determined by its reactive site (Arg346-Met347), which interacts with serine residue of tissue-type plasminogen activator (tPA) with concomitant formation of SDS-stable complex. Other sites may also play roles in determining the specificity of PAI-I toward serine proteases.

Results: To understand more about the role of distal hinge for PAI-I specificities towards serine proteases and for its conformational transition, wild type PAI-I and its mutants were expressed in baculovirus system. WtPAI-I was found to be about 12 fold more active than the fibrosarcoma PAI-I. Single site mutants within the Asp355-Arg356-Pro357 segment of PAI-I yield guanidine activatable inhibitors (a) that can still form SDS stable complexes with tPA and urokinase plasminogen activator (uPA), and (b) that have inhibition rate constants towards plasminogen activators which resemble those of the fibrosarcoma inhibitor. More importantly, latency conversion rate of these mutants was found to be ~3–4 fold faster than that of wtPAI-I. We also tested if Glu351 is important for serine protease specificity. The functional stability of wtPAI-I, Glu351Ala, Glu351Arg was about 18 ± 5 , 90 ± 8 and 14 ± 3 minutes, respectively, which correlated well with both their corresponding specific activities (84 ± 15 U/ug, 112 ± 18 U/ug and 68 ± 9 U/ug, respectively) and amount of SDS-stable complex formed with tPA after denatured by Guanidine-HCl and dialyzed against 50 mM sodium acetate at 4°C. The second-order rate constants of inhibition for uPA, plasmin and thrombin by Glu351Ala and Glu351Arg were increased about 2–10 folds compared to wtPAI-I, but there was no change for tPA.

Conclusion: The Asp355-Pro357 segment and Glu351 in distal hinge are involved in maintaining the inhibitory conformation of PAI-I. Glu351 is a specificity determinant of PAI-I toward uPA, plasmin and thrombin, but not for tPA.

Background

Plasminogen activator inhibitor-1 (PAI-1) is a serine proteinase inhibitor in the serpin superfamily [1–3]. This 50 kDa glycoprotein is apparently the most important physiological inhibitor of tissue-type plasminogen activator (tPA) and of urokinase plasminogen activator (uPA). [4] It was shown to play a crucial role in the regulation of vascular thrombosis, tumor invasion, neovascularization, inflammation and wound healing. [5,6] Increased plasma levels of PAI-1 were found to be correlated with an increased risk for cardiovascular diseases. [6–8] The essential function of PAI-1 in the fibrinolytic system has been confirmed by studies with transgenic animals. [7,9]

PAI-1 shares a 35% homology with 40 other serpins. [10] The X-ray structure of active PAI-1 [11,12] consists of three sheets (A-C), nine helices (a-i) and a reactive center loop (RCL). The RCL contains the residues Ser331 to Arg356 (P16-P10'), and within it there is a peptide loop (Glu 351 to Pro 357, P5'-P11'), which is defined as the distal hinge (Table 1). It should be noted that the placement and conformation of the RCL in active PAI-1 is quite different from that of latent PAI-1. [13] The movement of the distal hinge and the insertion of the RCL into sheet A as strand 4A (s4A) should take place during the transition from the active into the latent form of PAI-1.

The catalytic center of tPA interacts with the reactive site of PAI-1 with concomitant formation of 1:1 molar, SDS-stable complex. The P1 (Arg346) residue in the RCL of PAI-1 is an essential determinant for its target specificity and inhibitory activity. [14,15] Since many serpins with identical amino acid residue at P1 position display different specificities, it is unlikely that the P1 residue is the sole determinant for target protease specificity, other regions of PAI-1 may also affect its specificity toward target proteases. The variable region in serine proteases domain determines their specificity for PAI-1. Alignment of variable region 1 of the serine proteases shows a remarkable correlation between the composition of this area and their susceptibility to inhibition by PAI-1 as shown in Table 2. [16] Both tPA and uPA, the target proteases of PAI-1, contain a relatively extensive variable region 1, consisting of four to five positively charged amino acid residues present at identical position. Plasmin and thrombin with moderate reactivity toward PAI-1 have relatively short variable regions which contain only two to three positively charged residues. For example, positively charged amino acids in the variable region of tPA, uPA and thrombin play dominant roles in the specific interaction with PAI-1. [17–19] Furthermore, Glu350 (P4') of PAI-1 has been shown to mediate the interactions with tPA. [18]

In this study, By producing wtPAI-1 and several single site mutations in the distal hinge of PAI-1 in a baculovirus

expression system, we were able to find out the role of the distal hinge in conformational transition of PAI-1 and to elucidate the role of Glu351 in PAI-1 for serine protease specificities.

Results

Characterization of wtPAI-1

To confirm the validity of recombinant PAI-1 as a model to study the native inhibitor secreted from human fibrosarcoma cells, we compared the recombinant PAI-1 and the fibrosarcoma PAI-1 first by SDS-PAGE and immunoblot with anti-PAI-1 monoclonal antibodies. The molecular weight of wtPAI-1 was found to be lower than native PAI-1 (~50 kDa, Fig. 1B). We assumed that this is due to a lower (probably different) extent of glycosylation in the baculovirus expression system. Indeed, when both proteins were deglycosylated with N-glycanase they were found to have an essentially identical M.W. (~48 kDa) and to cross-react immunochemically (Fig. 1B). Furthermore, when this wtPAI-1 was chromatographed on heparin-Sepharose it was found to be > 98% pure by silver staining of the gel (Fig. 1A lane 2).

Interestingly, the specific inhibitory activity of wtPAI was 63 ± 15 U/ μ g just after purification, and was increased to 84 ± 21 U/ μ g after denatured by 6 M Gdn-HCl and refolded in 50 mM sodium acetate (pH 5.6) at 4°C. In contrast, the specific activity of the fibrosarcoma PAI-1 was only 5 ± 3 U/ μ g before activation, and was only increased to 45 ± 16 U/ μ g after denaturing and refolding. The high specific inhibitory activity of wt-PAI-1 could be due to the fact that it was secreted from insect High-5 cells cultured at 27°C, while the fibrosarcoma preparation of PAI-1 was obtained from the medium of fibrosarcoma cells cultured at 37°C [20]. Exposure to the higher temperature could accelerate the conversion of the inhibitor into its latent form.

The functional stability of wtPAI-1 and the fibrosarcoma PAI-1 in the absence or in presence of vitronectin are illustrated in Figure 1C. wtPAI-1 lost its inhibitory activity at the same pace as the fibrosarcoma PAI-1, indicating that its functional half life as an inhibitor and its ability to be stabilized by vitronectin are similar to those of the fibrosarcoma PAI-1.

The second-order rate constants (in $M^{-1}s^{-1}$ units) of the fibrosarcoma PAI-1 and the wtPAI-1 towards t-PA were $1.4 \pm 0.5 \times 10^7$ and $1.7 \pm 0.6 \times 10^7$, and towards uPA were $4.1 \pm 1.4 \times 10^7$ and $4.4 \pm 1.6 \times 10^7$, respectively. These results show that the wtPAI-1 could inhibit tPA or uPA at the same rate as the fibrosarcoma PAI-1. In addition, it is known that PAI-1 inhibits either tPA or uPA by forming a 1:1 SDS stable complex [4], wtPAI-1 indeed could form

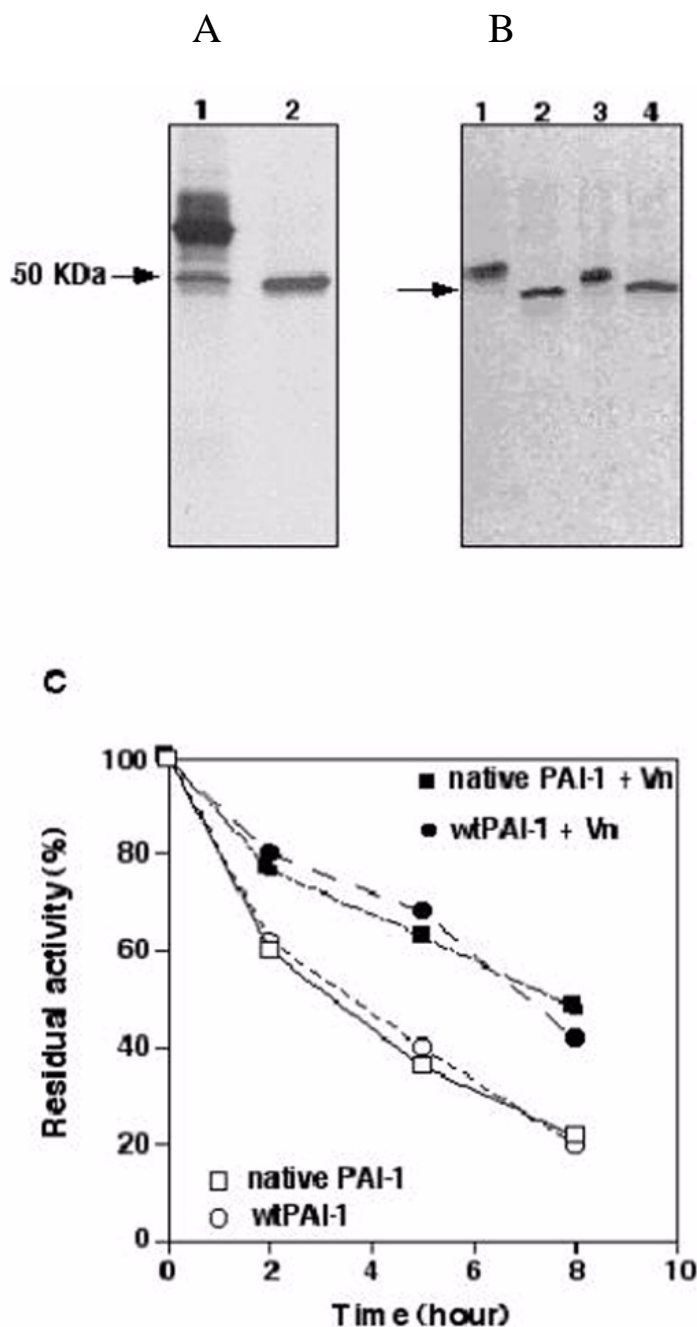


Figure 1

Purification and characterization of wtPAI-1. Panel A, Purification of wtPAI-1. The medium from High five cells was harvested 48 hours post infection and PAI-Is were purified on a heparin-Sepharose column and stained as described under Methods. Lane 1, the medium from infected High five cells, lane 2, wtPAI-1 eluted from the heparin-Sepharose column. Panel B, Treatment of fibrosarcoma- and wtPAI-1 with N-glycanase as described under Methods. Lane 1 and 2 were fibrosarcoma PAI-1, lane 3, and 4 were wtPAI-1. Fibrosarcoma PAI-1 (lane 2) and wtPAI-1 (lane 4) were treated with N-glycanase. Panel C, Stability of fibrosarcoma PAI-1 or of wtPAI-1. Fibrosarcoma PAI-1 or wtPAI-1 was incubated (37°C) in the absence (empty circle and square) or in the presence of a two molar fold of vitronectin (filled circle and square) in PBS. At the indicated time periods, aliquots were removed and the inhibitory activity of the PAI-Is towards uPA was measured as described under Methods. The inhibitory activity prior to the incubation at 37°C was taken as 100%.

Table 1: Amino acid sequence alignment at the carboxyl-terminal edges of various serpins

	s1C	s4B	s5B
PAI-1	EII <u>MDR</u> PFLFVVRHNPTGTVLFMGQVMEP		
Antitrypsin	EVKF <u>NK</u> PFVFLMIEQNTKSPLFMGKVVNPTQK		
Antithrombin	TFK <u>ANR</u> PFLVFIREVPLNTIIFMGRVANPCVK		
Antichymotrypsin	IVRF <u>NRP</u> FLMIIVPTDTQNIFFMSKVTNPKQA		
Ovalbumin	EFRA <u>DH</u> PFLFCIKHIATNAVLFFGRCSVSP		
Antiplasmin	SFSV <u>NR</u> PFLFFIFEDTTGLPLFVGSVRNPNSAPREL		

Table 2: Comparison of variable region of protease domain in serine proteases

Protease	Constant region 1	Variable region	Constant region 2
tPA	PWQAAI	FAK <u>HRR</u> SPGERF	LCGG
uPA	PWF AAI	Y <u>RR</u> H <u>R</u> GGSVTY	VCGG
Plasmin	PWQVSL	<u>R</u> TRFGMH	FCGG
Thrombin	PWQVM	<u>FR</u> KSPQEL	LCGA

Alignments of the amino acids sequences of the first variable region and parts of the surrounding constant regions of the protease domain of tPA, uPA, plasmin and thrombin. In the variable region positively charged amino acid residues are underlined.

SDS stable complex with tPA or uPA (Fig. 3A) just as fibrosarcoma PAI-1.

Single site mutations within the distal hinge of PAI-1

Since a proline occurs not only in position 357 of PAI-1, but also in the corresponding position of several other serpins (Table 1) we attempted to find out how does a mutation of P357 affect the inhibitory activity of PAI-1. Mutation of Pro357Gly blocked the inhibitory activity of PAI-1 (Fig. 2A). Similarly, a mutation into proline of either Asp355 or Arg356 also inactivated the inhibitor. It should be noted that all these mutants could be activated to a similar extent by 6 M Gnd-HCl (~20% inhibitory activity of the wtPAI-1, activated under the same conditions). These results suggest that a single site mutation at either one of the positions 355, 356 or 357 significantly inactivates PAI-1.

To elucidate the mechanism through which such mutations affect the activity of PAI-1, we determined their effect on the functional stability of PAI-1 and on its ability to form an SDS stable complex with either tPA or uPA. As seen in Fig. 2B, the rate of the spontaneous inactivation of the inhibitor was significantly increased upon mutation: the functional half life of the mutants Asp355Pro, Arg356Pro and Pro357Gly was 23.8 ± 4.5 min, 26.2 ± 3.7 min and 24.7 ± 4.8 min, respectively, while that of wtPAI-1 was 89.8 ± 8.7 min. Interestingly, there was no significant difference between the mutants tested and the wtPAI-1 in their ability to inactivate either tPA or uPA (Table 3).

They could also form SDS-stable complexes with tPA or uPA just as well as wtPAI-1 (Fig. 3A).

An interesting structural similarity among several serpins is known to occur at the position corresponding to 355 in PAI-1. While in PAI-1 this position accommodates a negatively charged aspartic acid residue, in other serpins it sometimes accommodates its non charged analog asparagine (Table 1). Asp355His, Asp355Lys, Asp355Glu, Asp355Gln and Asp355Asn were prepared and used to show that the negatively charged aspartic acid in PAI-1 is essential (though not sufficient) for its inhibitory activity. Mutation of Asp355 by asparagine (the non charged analog of aspartic acid), or by histidine or lysine (that have a positively charged side chain) or by alanine (neutral), inactivated PAI-1. Replacement of this aspartic acid residue even by the negatively charged glutamic acid (a very close analog of aspartic acid) did not significantly suffice to restore a full inhibitory activity of PAI-1 (Fig. 3B). After activation with 6 M Gnd-HCl, each of the six mutants exhibited a relatively high inhibitory activity, but their specific activities were still much lower than that of wtPAI-1 (Fig. 3B). Among them, Asp355Lys (5.1 ± 4.1 U/ug) had the lowest specific activity, while the Asp355Asn (20 ± 4.2 U/ug) had the highest specific activity (Fig. 3B). Although they had lower specific activities, all of the mutants mentioned above could still form SDS-stable complexes with tPA or uPA just as the wtPAI-1 (data not shown).

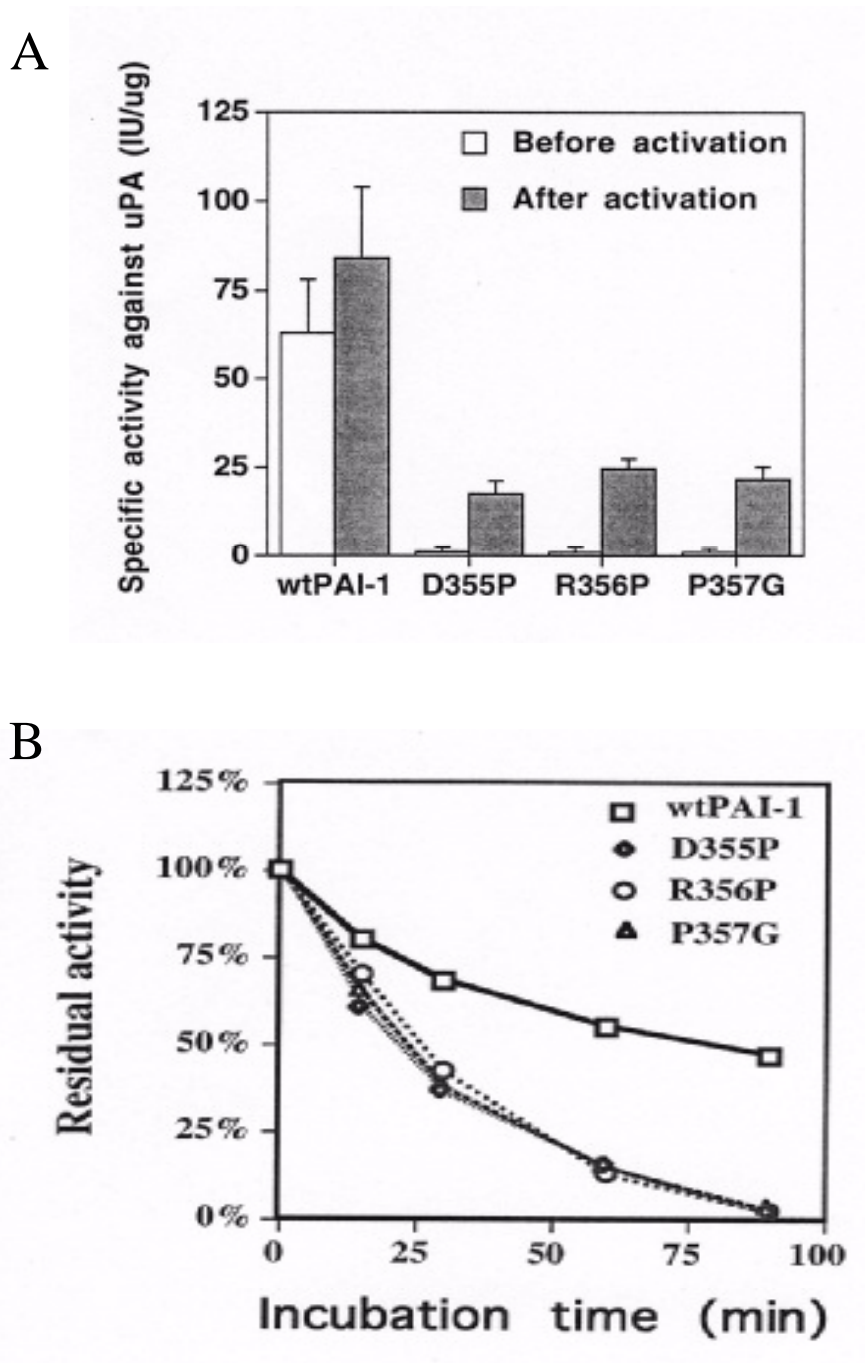


Figure 2

The specific inhibitory activity and functional stability of wtPAI-1, D355P, R356P and P357G Panel A: The specific inhibitory activity of wtPAI-1, D355P, R356P and P357G towards uPA was measured immediately after their purification and after their activation with 6 M Gdn-HCl as described under Methods. The data represent the mean \pm SEM of three separate determinations. Panel B: wtPAI-1, D355P, R356P and P357G were incubated in PBS at 37°C. At the indicated time periods, aliquots were removed and the inhibitory activity of the PAI-Is towards uPA was measured as described under Methods. The inhibitory activity prior to the incubation at 37°C was taken as 100%.

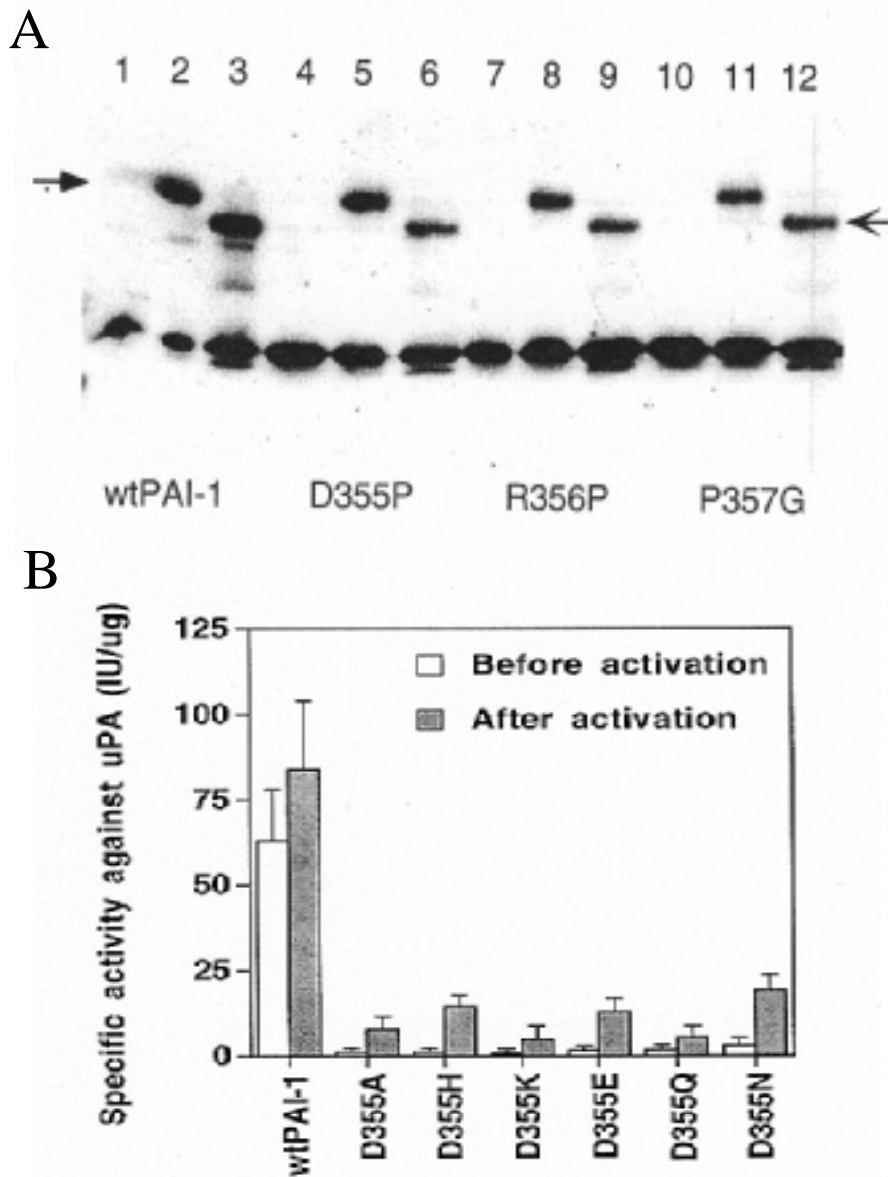


Figure 3

Complex formation and specific inhibitory activity of PAI-1s Panel A: The PAI-1s were incubated for 30 min at 37°C in the absence or in the presence of a two molar excess of either single-chain tPA or uPA. They were then subjected to SDS-PAGE and immunoblotting using anti-PAI-1 monoclonal antibodies. Lanes 1, 4, 7, 10 are PAI-1s only, lanes 2, 5, 8, 11 are PAI-1s with tPA, and lanes 3, 6, 9, 12 are PAI-1s with uPA. The filled arrow indicates the complexes between PAI-1s and tPA, and the regular arrow indicates the complex between PAI-1s and uPA. Panel B: The specific inhibitory activity of D355A, D355H, D355K, D355E, D355Q, and D355N towards uPA was measured immediately after their purification and after their activation with 6 M Gdn-HCl as described under Methods. The data represent the mean ± SE of three separate determinations.

Table 3: Second-order rate constants of wtPAI-1, D355P, R356P, P357G toward tPA or uPA.

	tPA(M ⁻¹ s ⁻¹)	uPA(M ⁻¹ s ⁻¹)
wtPAI-1	1.7 ± 0.6 × 10 ⁷	4.4 ± 1.6 × 10 ⁷
D355P	2.0 ± 0.7 × 10 ⁷	5.1 ± 1.7 × 10 ⁷
R356P	1.3 ± 0.5 × 10 ⁷	4.7 ± 1.7 × 10 ⁷
P357G	1.6 ± 0.7 × 10 ⁷	4.9 ± 2.5 × 10 ⁷

The second-order rate constants of inhibition, expressed in M⁻¹ s⁻¹ were determined as described in the Materials and Methods section. Values are averages of at least four independent experiments.

Two additional mutations, Arg356Ala and Arg356Glu, were prepared to further elucidate the role of Arg356. The specific inhibitory activity of Arg356Ala was 7.2 ± 2.5 U/μg, while Arg356Glu exhibited essentially no inhibitory activity. This specific activity was increased to 35.4 ± 8.7 U/μg for Arg356Ala and 6.4 ± 2.3 U/μg for Arg356Glu after denature and refolding. Just like ARG356Pro, Arg356Ala and Arg356Glu were much more labile than wtPAI-1. These results suggest that the role of the Asp355-Pro357 segment is to stabilize the inhibitory conformation of PAI-1.

Specific activity, functional stability and complex formation of Glu351Ala and Glu351Arg

The specific inhibitory activities of wtPAI-1, Glu351Ala and Glu351Arg were assessed with uPA and S-2444. As shown in Fig. 4A, the specificity of wtPAI-1, Glu351Ala and Glu351Arg after denatured by 6 M Gdn-HCl and dialyzed against 50 mM sodium acetate, pH 5.6, was 84 ± 15, 112 ± 18 and 68 ± 9 U/μg, respectively. Glu351Ala had a higher, and Glu351Arg lower specific activity than wtPAI-1. The specific activities of PAI-1s determined immediately after purification displayed similar pattern as that of inhibitors reactivated. There was about 15–20% increase after reactivation compared to each untreated protein itself. As shown in Fig. 4B, the two mutants also spontaneously lost their inhibitory activity (convert to latency) as wtPAI-1, but at different rates. The functional stability of wtPAI-1, Glu351Ala, Glu351Arg was about 18 ± 5, 90 ± 8 and 14 ± 3 minutes, respectively. It suggested that the specific activities measured above, to some extent, proportionally represent the functional stability. More importantly, amount complexes formed by tPA and PAI-1s remained after incubation at 37°C (Fig. 4C) were proportionally concordant with the results obtained from enzymatic stability assay (Fig. 4B). This proves that results from enzymatic assay are valid. The result of functional stability suggests that Glu351 is indeed involved in the conformational transition of PAI-1.

Inhibition of thrombin by PAI-1s and Neutralization of PAI-1s by Thrombin

Both mutants inhibited thrombin more efficiently than wtPAI-1, more importantly, Glu351Ala was better than Glu351Arg in inhibiting thrombin (Fig. 5A); Inhibition of thrombin by Glu351Arg was almost as same as by Glu351Ala in presence of vitronectin (Fig. 5B), which facilitated the thrombin inhibition by both mutants. When excess equal amount of thrombin was incubated with equal amount of PAI-1s in term of inhibitory activity against uPA, mutant inhibitors were neutralized faster by thrombin than wtPAI-1, and vitronectin also facilitated the neutralization of Glu351Ala or Glu351Arg by thrombin, just similar to the wtPAI-1 (Fig. 5C).

Inhibition of plasmin by PAI-1s

When plasmin was incubated with increased amount of equal active PAI-1s in absence or in presence of vitronectin, it was inhibited faster by mutants than by wtPAI-1 (Fig. 6 upper panel); and both of mutant also displayed improved inhibition toward plasmin in presence of vitronectin (Fig. 6 lower panel).

Kinetic analysis

The rate constants of tPA for wtPAI-1, Glu351Ala and Glu351Arg were similar (Table 4), however, The rate constants of inhibition toward uPA by Glu351Ala and Glu351Arg were 4 and 7-fold higher than by wtPAI-1, respectively. Moreover, The rate constants of Glu351Ala and Glu351Arg for thrombin inhibition were 2 to 4 folds higher than that of wtPAI-1 and rate constants of Glu351Ala and Glu351Arg to plasmin were 2 fold higher than that of wtPAI-1. This indicates that Glu351 is not essential for the interaction with tPA, but it is an very important determinant for the optimal interaction with uPA, thrombin and plasmin.

Discussion

The validity of human recombinant wtPAI-1 in representing the physiological form of the inhibitor is shown here by the following: (a) The inhibitory activity of this wtPAI-1 is actually higher than the inhibitory activity of the PAI-1 isolated from fibrosarcoma cells (12 fold more

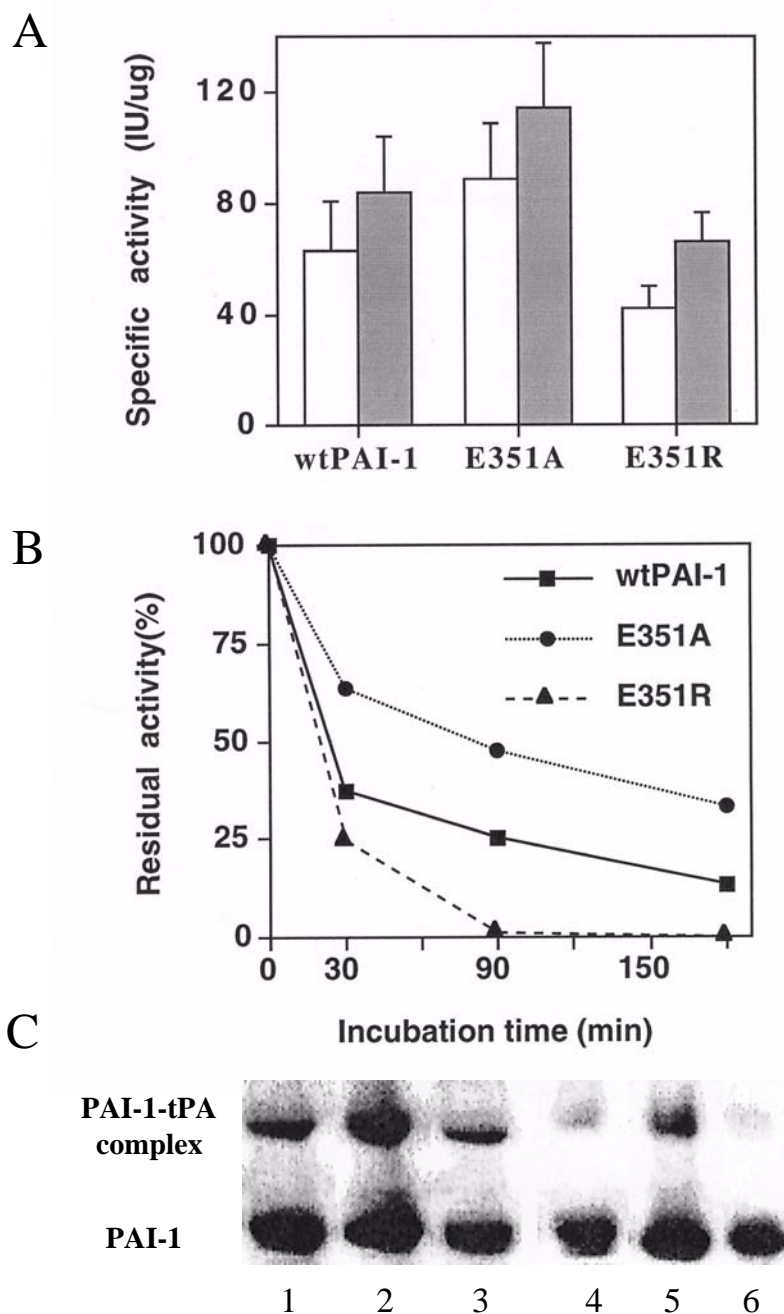


Figure 4

Specific activity, functional stability and complex formation of E351A and E351R. Panel A, Specific activities of wtPAI-1 and mutants were measured by a direct chromogenic assay using uPA and substrate S-2444 in microtiter plate as described under Methods. One unit of PAI-1 activity is defined as the amount that completely neutralizes 1 U of u-PA. Panel B, wtPAI-1, or mutants in assay buffer were incubated at 37°C. At times indicated, aliquots were taken out and determined their activity immediately as described above. The specific activity obtained at time zero was assigned a value of 100%. Panel C, complex formation of PAI-1s with tPA: certain amount of wtPAI-1, E351A and E351R were incubated with equal amount of tPA at 37°C for 30 minutes before put on ice, at same time these PAI-1s were incubated in PBS at 37°C for 3 hrs, respectively, before they were incubated with same amount of tPA at 37°C for 30 minutes, then the protein samples prepared at two time points were subjected to SDS-PAGE, and probed the membrane blot with anti-PAI-1 monoclonal antibody. Lane 1, 2 and 3 for wtPAI-1, E351A and E351R at zero time point, lane 4, 5, 6 for wtPAI-1, E351A and E351R after 3 hours' incubation at 37°C.

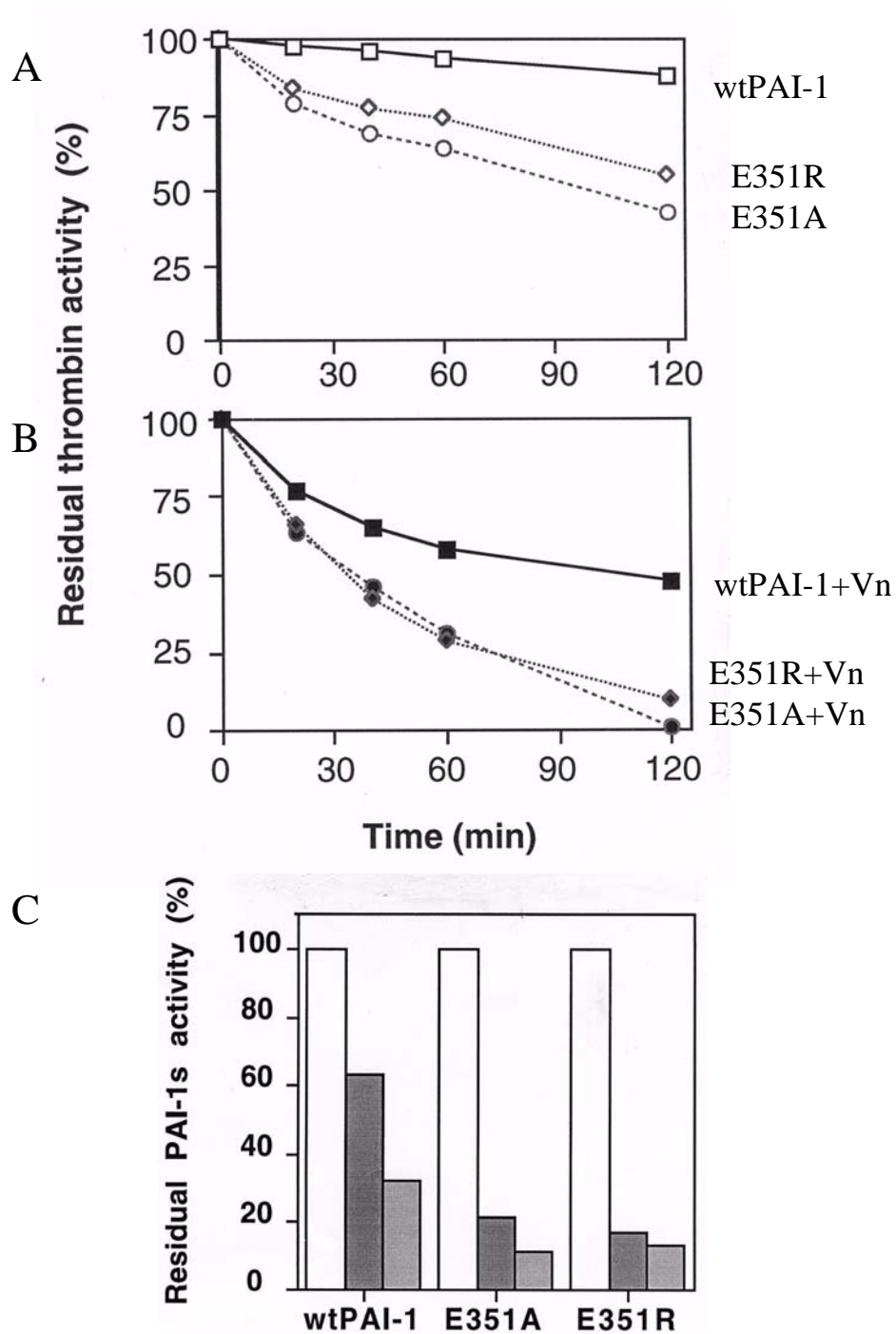


Figure 5

Inhibition of thrombin by wtPAI-I, E351A and E351R and Neutralization of PAI-I-s by thrombin Thrombin (50 ul, 0.5 U/ml) was incubated with equal amount of wtPAI-I, E351A, or E351R in term of inhibitory activity toward uPA (panel A), or in presence of vitronectin(panel B). After 20, 40, 60 and 120 min, remaining thrombin activities were determined with S-2238. The initial activity of thrombin was taken as 100%. Panel C, Forty nanomolar of thrombin were incubated with 10 nM of active PAI-I alone, or in presence of 40 nM vitronectin at 37°C. After 30 min, residual PAI-I activity was determined by titration on uPA. The residual PAI-I activity without incubation with thrombin was taken as 100%. Empty bars, PAI-I-s, filled black bars, PAI-I-s + thrombin, and filled light bars, PAI-I-s + thrombin + vitronectin.

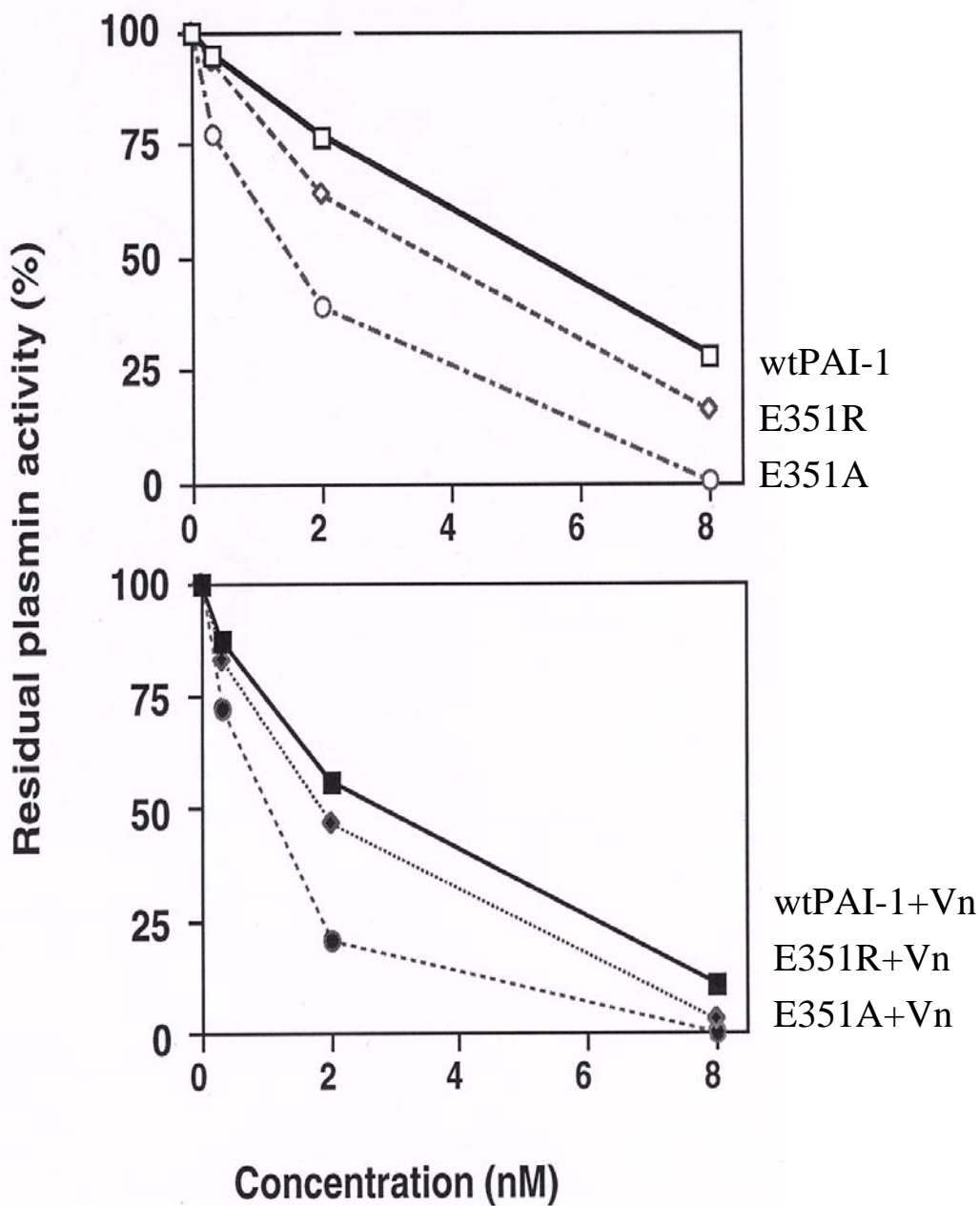


Figure 6

Inhibition of plasmin by wtPAI-1, E351A and E351R Plasmin (50 μ l, 100 nM) was incubated with equal amount of wtPAI-1, E351A, or E351R alone in term of inhibitory activity toward uPA (0, 0.32, 1.6 and 8 unit in 50 μ l, respectively, upper panel), or in presence of two molar excess of vitronectin(lower panel). Residual plasmin activity was calculated according to the absorbance at 405 nm after 60 minutes incubation with plasmin substrate (final concentration, 0.5 mM). The activity of plasmin without PAI-1s was taken as 100%.

Table 4: The second-order rate constants of wtPAI-1, E351A and E351R towards tPA, uPA, plasmin and thrombin.

	tPA (10^7 M-ls-l)	uPA (10^7 M-ls-l)	Thrombin (10^3 M-ls-l)	Plasmin (10^4 M-ls-l)
wtPAI-1	2.11 ± 0.83	0.48 ± 0.27	0.72 ± 0.31	3.10 ± 1.15
E351A	1.25 ± 0.49	2.64 ± 0.53	1.74 ± 0.35	6.29 ± 1.83
E351R	1.52 ± 0.35	3.62 ± 0.74	3.30 ± 0.42	7.12 ± 1.96

The second-order rate constants of inhibition, expressed in M-l s-l were determined as described in the Materials and Methods section. Values are averages of at least four independent experiments.

inhibitory prior to activation by 6 M Gnd HCl, and 2 fold more inhibitory after activation); (b) The inhibition rates of both tPA and uPA obtained with wtPAI-1 are essentially identical to those obtained with PAI-1 secreted by fibrosarcoma cells; (c) wtPAI-1 forms SDS-stable complexes with tPA and uPA that are similar to those obtained with the fibrosarcoma PAI-1; (d) The stabilization of the inhibitory conformation of wtPAI-1 by vitronectin is essentially identical to that of the fibrosarcoma PAI-1; (e) The molecular weight of the protein moiety of wtPAI-1 is similar to that of the fibrosarcoma PAI-1, if the two inhibitor preparations are deglycosylated with N-glycanase. In view of the above, we used mutated recombinant PAI-1s to assess the importance of specific amino acid residues in the inhibitory function of PAI-1.

This study focuses on the functional role of Glu351, Asp355, Arg356 and Pro357 within the distal hinge region of PAI-1. We suspected that the "bending" property of the prolyl residue at position 357 and the rigidity in the polypeptide backbone imposed by prolyl residues in general might have a role in maintaining the inhibitory conformation of this segment. Indeed, we found that a mutant of Pro357 in which the proline residue was replaced by glycine is non inhibitory. This mutant could, however, be activated by 6 M Gdn-HCl, supporting the notion that the lack of activity of Pro357Gly may indeed be due to a conformational effect on the structure of the inhibitor.

The structural importance of the local conformation of the Asp355-Pro357 segment is further illustrated by two single site mutants in which the original Pro357 was preserved, but an additional proline residue was introduced (by mutation) at positions 355 or 356. These two mutants were inactive but they were activated considerably by treatment with 6 M Gdn-HCl. The enhanced lability of these mutants was evident from a more rapid conversion into their latent structure. These mutants were found to form SDS-stable complexes with either tPA or uPA after treatment with 6 M Gdn-HCl. This treatment presumably allows them to restore (at least in part) their inhibitory structure, and their ability to inhibit plasminogen activa-

tors with second order rate inhibition constants similar to those obtained with the wtPAI-1. Since these mutants have faster latency transition time, we assume that these mutations do so by loosening the structural restrictions normally imposed on the mobility of the RCL. [21] However, we show here that in spite of the fact that these mutants are structurally more fragile, their second-order rate constants towards their target proteinases are very similar to those of wtPAI-1. This implies that a facilitated latency transition does not necessarily result in slow rate inhibition towards tPA or uPA. This is in line with the findings that stable PAI-1 mutants (72-fold or 245-fold longer half life than of wtPAI-1) have similar inhibition rate constants towards uPA. [22,23] We therefore conclude that the segment Asp355-Pro357 is not involved directly in the complex formation, but rather in the latency transition.

The results presented here with the mutants expressed in High 5 cells show that indeed the conversion of the negative into a positive charge (Asp355His and Asp355Lys) yields an inactive inhibitor protein, but the preservation of the negative charge does not suffice by itself. Mutation of this aspartic acid residue into a negatively charged glutamic acid (also negatively charged, and differs from Asp355 by having an extra methylene group only) still yields a mutant PAI-1 that has a very low inhibitory activity. The fact that the negative charge is an important factor in the inhibitory activity is illustrated here by the finding that Asp355Asn and Asp355Gln have both a very low activity, yet they are more active than Asp355Lys and Asp355His. Generally speaking, it is clear from our studies that any mutation of Asp355, whether it preserves the negative charge or not, yields an inactive inhibitor.

The markedly enhanced functional stability of Glu351Ala (Fig. 4B) may be due to the unfavorable interactions resulted from Ala instead of Glu, which delays the latency conversion, while the decreased functional stability of Glu351Arg may be caused by positively charged Arg, which may facilitate the conformation transition. Here we show that different amino acid substitution at the same position result in opposite effect on the PAI-1 stability,

suggesting that the network in which Glu351 is involving does provide PAI-1 stability. Moreover, the structural stabilization needs not necessarily affect its mechanism of reactive center loop insertion, but instead may occur through subtle optimization of interactions. Indeed, results of studies with other proteins have generally supported the notion that thermodynamic stabilization of a protein is global. This is also consistent with studies of mutant T4 lysozymes with multiple substitutions at one position demonstrated a direct relationship between the hydrophobicity of the substituted residue and the resulting increase in stability. [24]

Cognate protease specificity

Improvements in specificity of mutants for uPA indicates that Glu351 involve in the interaction with uPA. To the best of our knowledge, this is the first time that the specificity of PAI-1 toward uPA is partially mediated by the negatively charged residue located in the C-terminal of cleavage site in PAI-1. The interaction of Glu351 with uPA is much more important than that of Glu351 with tPA, which is demonstrated by improved reactivity of Glu351 mutants toward uPA, but not toward tPA, showing that tPA is not directly interact with Glu351 of PAI-1, whereas Glu351 is necessary for the optimal interaction with uPA. Our results are concordant with previous study, [25] which suggests that serpin body-protease body interactions play significant roles in determining serpin inhibitory activity against target proteases.

Non-cognate protease

Substitution of Glu351 also altered the specificity toward non-cognate proteases: plasmin and thrombin. Glu at 351 may provide steric hindrance that slows the rate of thrombin or plasmin inhibition. Mutations at Glu351 may decrease the original repulsive interactions with variable region of thrombin, which may slow the rate of thrombin inhibition, or results in favorable interactions that may not exist when wtPAI-1 react with thrombin. So, both Glu351Arg and Glu351Gln are better thrombin inhibitors than wtPAI-1. Similar results were observed for plasmin. With single mutation at position Glu351, we improved reactivity of PAI-1 to thrombin and plasmin without significantly affecting its specificity to tPA. This substantiates previous mutagenesis studies, which demonstrated that repulsive interactions and/or lack of productive electrostatic interactions between Glu39 and Glu192 of thrombin and PAI-1 are responsible for the slow reaction of thrombin with this serpin.[26] Furthermore, the rapid inhibition of plasmin requires a non-covalent interaction between an amino-terminal site of plasmin and a carboxyl-terminal on anti-plasmin, and a second site outside the RCL of PAI-1 contributes to its specific interaction with proteases.[15,27] The specificity of the two mutants toward thrombin and plasmin implicates

that Glu351 of PAI-1 is involved in its specificity toward thrombin and plasmin.

Conclusion

Single site mutations within the segment Asp355-Arg356-Pro357 of PAI-1 yield Gdn-HCl activatable inhibitors that can still form SDS stable complexes with plasminogen activators, with second order inhibition constants that are similar to the native inhibitor. Interestingly, the conversion of these mutants to latent forms was ~3–4 fold faster than wtPAI-1, suggesting that the Asp355-Pro357 segment is involved in maintaining the inhibitory conformation of PAI-1. Glu351 contributes to the optimal functional stability of PAI-1 and participates its conformational stability. More importantly, Glu351 of PAI-1 is a specificity determinant for uPA, thrombin and plasmin, but not for tPA.

Methods

The following materials were obtained from commercial sources: native PAI-1, secreted into the cell culture medium from human fibrosarcoma cell line HT 1080 (Biopool); single-chain tPA and two-chain tPA standard (Biopool); uPA, plasmin and thrombin were from Calbiochem; the chromogenic substrate for uPA, pGlu-Gly-Arg p-Nitroanilide (S-2444), the chromogenic substrate for tPA and H-D-Ile-Pro-Arg-pNA-2HCl (S-2288) from Chromogenix AB, chromogenic substrate for thrombin, H-D-Phe-pipecolyl-Arg-pNA(S-2238) and chromogenic substrate for plasmin, N-p-Tosyl-Gly-Pro-Lys-p-Nitroanilide from Sigma); heparin-Sepharose, monoclonal antibody against PAI-1 from American Diagnostica, Inc.; recombinant N-glycanase (Genzyme diagnostics); the pRSET(A) vector, baculovirus transfer vector pAcSG2 and Baculo-Gold transfection kit from Pharmgen; Sf9 cells (*Spodoptera frugiperda*), High Five cell (*Trichoplusia ni*) from Invitrogen; Restriction enzymes, T₄ ligase and shrimp phosphatase were purchased from New England Biolab. BCA protein assay reagent from Pierce; A full-length cDNA of human PAI-1 (pPAI-1-AI) [28] was generously provided by Prof. Keld Danö, The Finsen Laboratory.

General DNA techniques

Plasmid DNA was purified using either Promega or QIA-GEN miniprep kits. The DNA was sequenced using the dideoxy chain termination reaction method using an ABI 373 DNA sequencer. Oligonucleotides were synthesized using a 380 B DNA synthesizer (ABI model). DNA manipulation techniques were carried out according to standard procedures [29].

Construction of wtPAI-1 and its mutants

The pPAI-1-AI was digested with EcoRI and BglII to release a 1426 basepair (bp) PAI-1 coding sequence, which was isolated, purified, and ligated into EcoRI and BglII

digested pVL1393. The resultant pVL1393-PAI-1 was analyzed by restriction mapping and sequencing. The mutants at E351A, E351R, D355, R356, and P357 of PAI-1 were generated by PCR. The occurrence of the resulting mutations D355A, D355E, D355H, D355K, D355N, D355P, D355Q, R356P, P357G as well as E351A and E351R in pRSET was confirmed by sequencing. The PAI-1 sequence containing the desired mutations was digested with EcoRI and BglII, and then subcloned into pVL1393. The recombinant baculoviruses containing either wtPAI-1, or one of the above mutants was obtained as described by Kjoller et al. [30]

Expression and purification of wtPAI-1 and its mutants

High 5 cells, grown at 27°C in SF-900 serum-free medium, were infected with high-titer virus stocks ($1-3 \times 10^8$ /ml). The medium was collected 48 hrs after the infection (found to be the optimal harvesting time for wtPAI-1) and then centrifuged for 5 minutes ($125 \times g$). The supernatants (50 ml) were collected and sodium chloride (final concentration 0.25 M) was added. They were then applied on a heparin-Sepharose column (bed volume 5 ml) that was equilibrated and run (flow rate 20 ml/h) with a buffer consisted of 50 mM sodium acetate, pH 5.6, containing 0.25 M NaCl and 0.01% Tween-20. The columns were washed and the PAI-1s were eluted with a 40 ml gradient of 0.25 – 1 M NaCl in the same buffer.

SDS-PAGE, immunoblotting and deglycosylation of PAI-1

The medium of the infected cells and the eluent from heparin-Sepharose were separated on 10% SDS-PAGE, and monitored by silver staining. Fibrosarcoma PAI-1 and wtPAI-1 were transferred onto nitrocellulose paper and the proteins were revealed with anti-human PAI-1 monoclonal antibodies, followed by horseradish peroxidase (HRP) conjugated rabbit anti-mouse antibody, and enhanced chemiluminescence (ECL). 5 ug of Fibrosarcoma PAI-1, or wtPAI-1 was denatured by boiling for 5 min in a buffer consisted of 0.5% SDS, 50 mM β -mercaptoethanol and 50 mM Tris-HCl, pH 7.4. They were then digested (18 hr, at 37°C) with 0.5 units of recombinant N-glycanase in the presence of 2.5% NP-40, and subjected to SDS-PAGE and silver staining. Protein concentrations were determined by the Bio-Rad protein assay reagent.

Functional stability of fibrosarcoma PAI-1, wtPAI-1 and its mutants

The inhibitory activities of fibrosarcoma PAI-1, wtPAI-1, and its mutants were measured by a direct chromogenic assay using uPA (100 U/ μ g) and its substrate S-2444 in a microtiter plate as described earlier [20]. Briefly, fibrosarcoma PAI-1, wtPAI-1, or its mutants were diluted 1:10 in the activity measuring assay buffer (50 μ M Tris-HCl, pH 7.5, containing 0.15 M NaCl and 100 μ g/ml BSA). The PAI-1 samples were incubated in the absence or in the

presence of a two molar excess of vitronectin (the fibrosarcoma PAI-1 and the wtPAI-1 were incubated with vitronectin) at 37°C. At the times indicated, aliquots were removed and analyzed immediately for their inhibitory activity. One unit of PAI-1 activity is defined as the amount that completely neutralizes 1 U of uPA. The specific activity obtained at time zero was assigned a value of 100%.

Activation of PAI-1 and kinetic analysis

Fibrosarcoma PAI-1, wtPAI-1 or its mutants were denatured with 6 M guanidine-hydrochloride (Gdn-HCl) (45 min at 37°C), followed by dialysis (4–16 hours at 4°C) against 50 mM sodium acetate, pH 5.6, containing 0.5 M NaCl and 0.01% Tween-20. This process which PAI-1 regained its activity is called activation by Gdn-HCl. The inhibitory activity of each of the PAI-1s was assayed immediately. The second-order rate constants for the interaction of the fibrosarcoma PAI-1, wtPAI-1 or the indicated mutants with single-chain tPA or uPA were determined in a single-step assay as previously described. [20]

Specific activity, functional stability and complex formation of PAI-1s

The inhibitory activities of wtPAI-1 and mutants were measured by a direct chromogenic assay using uPA (100 U/ μ g) and substrate S-2444. [17] Briefly, serially diluted wtPAI-1, or mutants were incubated with equal volume of uPA (50 μ l, 50 U/ml) for 30 min at 37°C, followed by addition of 100 μ l, 0.5 mM S-2444. Residual uPA activity was quantitated by measuring the change in absorbance at 405 nm with ELISA reader. The specific activity of PAI-1 was calculated based on the amount of uPA inhibited by PAI-1. The assay for PAI-1 inhibitory activity toward two-chain tPA was the same for uPA except that the samples were incubated with tPA and S-2288. PAI-1 inhibitory activity is expressed as the number of international units of PAs inhibited by PAI-1. Specific activity of uPA and two-chain tPA are 100 U/ μ g and 764 U/ μ g, respectively. For the functional stability, wtPAI-1, or mutants were incubated at 37°C. Aliquots were taken out at times indicated and analyzed immediately as described above. The specific activity obtained at time zero was assigned a value of 100%. To provide the evidence to support the conclusion obtained from enzyme assay, certain amount of wtPAI-1, E351R and E351A were incubated with equal amount of tPA at 37°C for 30 minutes before put on ice, at same time these PAI-1s were incubated in PBS at 37°C for 3 hours, respectively, before the PAI-1s were incubated with same amount of tPA at 37°C for 30 minutes, then the protein samples prepared at two time points were subjected to SDS-PAGE and western blot as described before.

Inhibition of thrombin and plasmin by PAI-1s

Thrombin (0.025 U), or plasmin (0.1 µg/ml) was incubated at 37°C for 30 min with same amount of wtPAI-1, E351A and E351R in term of inhibitory activity toward uPA (20 U in the concentration range of 4–10 µg/ml) in absence or presence of vitronectin (60 µg/ml). Linear increase of absorbance at 405 nm were recorded after adding S-2238 (final concentration 0.25 mM) for thrombin, or N-p-Tosyl-Gly-Pro-Lys-p-Nitroanilide (final concentration 0.5 mM) for plasmin. Thrombin, or plasmin activity without PAI-1 was taken as 100%.

Neutralization of PAI-1s by Thrombin

wtPAI-1 and mutants were incubated with four molar excess of thrombin (40 nM) in the absence or presence of vitronectin (30 nM). Control experiments were carried out in the absence of thrombin. After 20 min at 37°C, the activity of thrombin was quenched by the addition of hirudin (final concentration, 60 units /ml), resulting in inhibition of more than 95% of the amidolytic activity of thrombin. Subsequently, the residual PAI-1 inhibitory activity was determined by titration of aliquots on uPA, using S2444 as substrate. The increase of the optical density at 405 nm in the samples was corrected for the values measured with (hirudin-inactivated) thrombin alone, since the amidolytic activity of thrombin on S2444 was not completely blocked by hirudin,

Complex formation of PAI-1s with tPA or uPA

Complex formation of the fibrosarcoma PAI-1, wtPAI-1 or its mutants with single-chain tPA or uPA were performed as described earlier [20]. Samples of PAI-1s (5 U) were incubated in the absence or in the presence of a 2-fold excess of uPA or tPA in the activity assay buffer. Following a 30 min incubation at room temperature, the samples were subjected to a 10% SDS-PAGE and immunoblotting as described above.

Abbreviations

tPA, tissue-type plasminogen activator; uPA, urokinase-type plasminogen activator; PAI-1, plasminogen activator inhibitor type 1; serpin, serine protease inhibitor; RCL, reactive center loop; Vn, vitronectin, SDS, sodium dodecylsulfate; PAGE, polyacrylamide gel electrophoresis. Gdn-HCl, guanidine-hydrochloride; HRP, horseradish peroxidase; ECL, enhanced chemiluminescence; PBS, phosphate-buffered saline; BSA, bovine serum albumin.

Authors' contribution

Qingcai Wang carried out the experiments and drafted the manuscript;

Professor Shmuel Shaltiel sponsored and supervised the project and revised the manuscript.

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