The Specific Activity of Plasminogen Activator Inhibitor-1 in Disseminated Intravascular Coagulation With Acute Promyelocytic Leukemia

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In disseminated intravascular coagulation (DIC) with acute promyelocytic leukemia (APL) in the absence of severe infection, marked fibrinolysis was noted in comparison with normal levels of antithrombin III, which is a major inhibitor of the coagulation system. Increased plasminogen activator inhibitor-1 (PAI-1) antigen levels in plasma from patients with septicemia decreased the ratio of the plasma clot lysis rate induced by an anti-α2-plasmin inhibitor monoclonal antibody to the tissue-type plasminogen activator (t-PA) concentration. This decrease was not as prominent in plasma from patients with DIC, especially those with APL. To explore the character of PAI-1 in these plasmas, we measured the specific activity of PAI-1 by determining the ratio of active PAI-1 antigen to t-PA-unbound PAI-1 antigen. To calculate the amount of active PAI-1 antigen, the amount of t-PA/PAI-1 complex before and after the addition of a fixed amount of t-PA to the sample was measured by a sandwich solid-phase enzyme-linked immunosorbent assay using anti-PAI-1 and anti-t-PA monoclonal antibodies. The assay to measure total PAI-1 antigen used three monoclonal anti-PAI-1 antibodies and had similar sensitivities to free active, latent, vitronectin-bound and t-PA-bound PAI-1. The specific activity of PAI-1 decreased in patients with DIC (43.7% ± 30.6%) and in DIC cases with APL (10.3% ± 6.0%) in comparison to patients with septicemia (83.7% ± 20.2%) or normal controls (85.8% ± 27.3%). In DIC associated with APL, degraded forms of PAI-1 were detected in plasma by immunoblotting. These results suggest that a decrease in the specific activity of PAI-1 and an increase in secondary fibrinolysis result in a hyperfibrinolytic state in DIC patients with APL.

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**MATERIALS AND METHODS**

Reagents. The following materials were purchased: Freund's adjuvant, polyethylene glycol 1540, dimethyl sulfoxide, 2,6,10,14-tetramethylpentadecane (Pristane), and Tween 20 (Wako Chemical, Osaka, Japan); penicillin and streptomycin (GIBCO, Grand Island, NY); calf serum, Dulbecco's modified Eagle's medium (DMEM), and 96-well polystyrene microtiter plates (NUNC) (Flow Laboratories, North Ryde, Australia); fetal calf serum (FCS; Filtron, Altona, Australia); tissue culture flasks (Corning Glass Works, Corning, NY); gentamicin, bovine serum albumin (BSA; essentially fatty acid-free, globulin-free), theophylline, and prostaglandin E1 (Sigma, St Louis, MO); solid-phase lactoperoxidase-glucose oxidase (Enzyme conjugates), horseradish peroxidase-conjugated goat antimouse IgG (BioRad, Richmond, CA); CNBr-

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activated Sepharose 4B, protein A-Sepharose CL-4B, and Sepharose PD10 (Pharmacia-LKB Biotechnology, Uppsala, Sweden); collagen (Niko Bioscience, Tokyo, Japan); and 2,2'-asino-di-3-ethyl-benthanzoline sulfonate] (ABTS) (Kirekegaard & Perry Laboratories, Gaithersburg, MD). The mouse myeloma cell line P3-X63-Ag8-U1 (P3U1) was provided by Dr T. Watanabe (Kyushu University, Fukuoka, Japan). Aprotinin and single-chain urokinase-type plasminogen activator (scu-PA) were gifts from Mochida Pharmaceutical (Tokyo, Japan).

t-PA was purified from medium conditioned by human melanoma cells with aprotinin as described previously and had a specific activity of 550,000 U/mg protein according to the t-PA International Standard (63/517). It migrated as a single band in SDS-polyacrylamide gel electrophoresis (SDS-PAGE) under reducing and nonreducing conditions. The t-PA International Standard was supplied by the National Institute for Biological Standards and Controls (London, England). Glu-plasminogen was purified from fresh frozen plasma in the presence of 100 ng unlabeled activating units (KIU/mL aprotinin as described. Fibrinogen was prepared according to the method of Blomback and Blomback. Vitronectin was initially prepared from fresh frozen plasma according to the method of Dahlback and Fodack. Immunopurified VN was also prepared from fresh frozen plasma by ammonium sulfate precipitation (50% saturation) and affinity chromatography on a column containing anti-VN polyclonal antibody immobilized on Sepharose. Human PAI-1 was purified from the conditioned media of cultured human umbilical vein endothelial cells basically as described by Hekman and Loskutoff. Except for immunopurified VN, the purified proteins appeared to be greater than 98% homogeneous on SDS-PAGE. Immunopurified VN had a significant amount of material that smeared from the top of the gel to the 65,000-dalton polypeptide on SDS-PAGE. Protein concentrations were determined from their respective extinction coefficients at 280 nm. Protein concentrations of t-PA and PAI-1 were determined by amino acid analysis by the phenylthiocarbamyl method as described using the PICO.TAG system (Waters Associates, Milford, MA) after hydrolysis for 24 hours in HCl vapor at 110°C. The quantities of t-PA and PAI-1 were calculated based on their amino acid composition as deduced from their cDNA sequences and assumed molecular weights of 70,000 and 50,000, respectively.

Monospecific antisera against t-PA, scu-PA, PAI-1, and VN were prepared in rabbits as described previously. The t-PA and PAI-1 complex was purified as follows: Confluent cultures of human endothelial cells (1.5 x 10⁶/cm²) were washed three times with serum-free medium 199 and then incubated in serum-free medium containing 1 µg/mL t-PA for 6 hours. This conditioned medium was applied to an affinity column containing anti-t-PA antibody-immobilized Sepharose. Bound antigen was eluted with 3 mol/L KSCN and was further purified using anti-PAI-1 antibody-immobilized Sepharose as described. The quantity of PAI-1 present as a complex was calculated from the amount of t-PA present, assuming a 1:1 stoichiometric complex. The amount of t-PA was measured by ELISA using anti-t-PA monoclonal antibodies (MoAbs) JTA-1 and JTA-2 as described. In this ELISA, the amount of t-PA complexed with PAI-1 can be measured by using the same standard curve constructed for free t-PA as described. The level of t-PA was also confirmed by the commercial ELISA kit Imulyste t-PA (Biopool AB, Umea, Sweden).

MoAbs against PAI-1, using either t-PA/PAI-1 complex or latent PAI-1 as an antigen, were raised by standard methods using BALB/c mice and the mouse myeloma cell line P3U1 as described. The dissociation constants (kd) of 125I-labeled MoAbs for the latent PAI-1 and t-PA/PAI-1 complex were assessed by a solid-phase assay as described, according to the Frankel and Gerhard method. The competition between MoAbs (JTAI-1, JTAI-2, and JTAI-3) against PAI-1 for the same epitope on PAI-1 was tested as previously described. Peroxidase-coupled MoAbs were prepared by coupling the antibodies to horseradish peroxidase using m-piodylate by the method of Nakane and Kawaoi.

Clinical groups. DIC patients (n = 58) in this study were separated into two groups: groups with (n = 15) or without APL (n = 43). The presence of DIC was diagnosed for each patient by marked reductions in platelet counts, fibrinogen, antithrombin III (except in APL without sepsis), plasminogen, and a2-plasmin inhibitor levels and elevated levels of fibrin degradation products. In this study, DIC patients with APL and without sepsis (n = 13) were carefully chosen and separated from the two cases with severe sepsis (see Fig 5, lanes g and h). The DIC patient group without APL included 23 patients with leukemia, 12 patients with lung cancer, 6 patients with gastric cancer, and 2 patients with biliary tract infection. Thirty-two patients had either high- or low-grade fever. A sepsis group without DIC included five patients with malignant tumor, two with respiratory tract infection, and five with peritonitis.

Plasma. Blood was collected from the antecubital veins of healthy volunteers and patients into 0.1 vol 3.8% sodium citrate with or without theophylline (final 0.001 mol/L) and prostaglandin E₂ (final 0.5 mg/L) with a 30-second handcuff at about 9 AM. The collected blood was immediately cooled with ice water and centrifuged at 2,000g for 20 minutes at 4°C to prepare platelet-poor plasma (PPP). All plasma was stored at −80°C until use. Gel-filtered platelets and platelet-rich plasma were prepared as described. PAI-1–depleted plasma was prepared from pooled healthy volunteers’ plasma by passage through rabbit anti-PAI-1 IgG-immobilized Sepharose (5 mg IgG/mL Sepharose), which had been equilibrated with a mixture of 0.9 vol 0.15 mol/L NaCl and 0.1 vol 3.8% sodium citrate. Single time point plasma samples were taken from patients with DIC.

Plasma clot lysis induced by MoAb against α2-plasmin inhibitor (α2-PI). Plasma was supplemented with 2 μmol/L Glu-plasminogen and 1.2 μmol/L of anti-α2-PI MoAb JTP-1, which induces spontaneous plasma clot lysis. α-Thrombin (final 0.1 U/mL) and CaCl₂ (final 0.015 mol/L) were added, and the solution was allowed to clot for 15 minutes at 37°C in a glass tube. The clots were wound onto a bamboo stick and squeezed to remove as much fluid as possible. Clots were subsequently resuspended in the same sera. Fibrinolysis was monitored by measuring the amount of fibrin degradation products in the supernatant after 3 hours of incubation at 37°C using counter-immunoelectrophoresis as described. The amount of FDP before incubation was subtracted from this value.

Solid-phase ELISA for total PAI-1. Polystyrene chloride microtiter plates were coated with a mixture of two MoAbs against PAI-1 (10 µg/mL JTAI-1 and JTAI-2) for 16 hours at 4°C as described. The plates were emptied and the wells were treated for 2 hours at 25°C with 0.05 mol/L Tris-HCl, 0.1 mol/L NaCl, pH 7.4 (TBS buffer) containing 1% BSA. After washing twice with TBS containing 0.05% Tween 20 (TBS-Tween) and 0.1% BSA, samples were diluted in the same buffer and aliquots were added to the coated wells. When samples were plasma, the samples were diluted with the same buffer four times and the standard curves were constructed with various concentrations of purified PAI-1 diluted with a final 25% PAI-1 immunodepleted plasma. Variability due to dilution of plasma protein was adjusted by using PAI-1 immunodepleted plasma. The plates were then incubated for 10 hours at 4°C. After the plates were washed with cold PBS-Tween, peroxidase-anti-PAI MoAb JTAI-3 was added to each well and incubated for another 10 hours at 4°C. After washing twice with cold PBS-Tween, hydrolysis of the ABTS substrate by adsorbed JTAI-3 was
monitored at 25°C on an ELISA analyzer ETY-96 (Toyo Sokki, Tokyo, Japan) for 30 minutes.

**Solid-phase ELISA for t-PA/PAI-1 complex.** The ELISA for t-PA/PAI-1 complex in samples was performed basically as described above using JTAI-1 as the coating antibody and peroxidase-anti-t-PA MoAb JTA-1. To assess the level of active PAI-1 antigen in a sample, single-chain t-PA (final 75 ng/mL) and aprotinin (final 500 U/mL) were added to the sample and the mixture was incubated for 30 minutes at 37°C. After incubation, samples were diluted four times with TBS-Tween containing 0.1% albumin, and differences due to interference of plasma proteins were adjusted with PAI-1-depleted plasma added just before ELISA. The quantity of active PAI-1 was calculated from the amount of complex in the standard in the original sample and assuming a 1:1 stoichiometry.

**SDS-PAGE.** SDS-PAGE in slab gels was performed using resolving gels of 10% acrylamide and stacking gels of 4% according to Laemmli.30 SDS-PAGE in tube gels was performed using gels of 7% acrylamide according to Weber and Osborn.31 Molecular weight standards were obtained from Bio-Rad. To observe the profile of PAI-1 from plasma, plasma was immunoadsorbed with anti-PAI-1 polyclonal antibody immobilized on Sepharose. The column was eluted with sample buffer for SDS-PAGE containing 2% SDS, and samples were fractionated by SDS-PAGE. The gel was processed for reverse fluorography or Western blotting. Western blotting of the samples was performed after electrical transfer to nitrocellulose, using anti-PAI-1 IgG, as previously described.32

**Miscellaneous.** Human umbilical vein endothelial cells were isolated from human umbilical cord veins by the method of Jaffe et al33 and were cultured as described.34 Plasma concentrations of scu-PA were measured basically as described previously by ELISA using anti-urokinase MoAb JS-1 (final 10 ng/mL) as a coating antibody, peroxidase MoAb JS-2 as a second antibody, and purified scu-PA as the standard urokinase-type plasminogen activator (u-PA). This ELISA measures the amount of single-chain u-PA, two-chain u-PA that was activated by plasmin, and u-PA complexed with PAI-1 by using the same standard curve for free scu-PA (data not shown). Purified PAI-1 was reactivated with either 8 mol/L urea or 4 mol/L guanidine hydrochloride at pH 5.5 for 30 minutes at 37°C, and dialyzed for 12 hours at 4°C against 0.05 mol/L sodium acetate pH 5.5 containing 0.1 mol/L NaCl and 0.05% Tween 20 and for 2 hours at 4°C against TBS-Tween pH 7.4. PAI activity in a sample was assessed by the method as described13,14 (conventional functional assay of PAI) using single-chain t-PA, plasminogen (1 μmol/L), CNBr-digested fibrinogen (1 μmol/L), and S-2251 (Kabi Diagnostica, Stockholm, Sweden) (0.5 mmol/L). One unit of PAI is defined as the amount of PAI that neutralizes 2 ng of single-chain t-PA in 10 minutes at room temperature. MoAbs against PAI-1 were radiolabeled using lactoperoxidase-glucose oxidase and Na125I (20 Ci/mg) (New England Nuclear, Boston, MA). The labeled MoAbs JTAI-1, -2, -3, -4, and -5 had specific activities of 6.8 × 105, 7.2 × 105, 7.7 × 105, 8.3 × 105, 6.3 × 105, and 8.8 × 105 cpm/μg, respectively. The concentration of α2-PI/plasmin complex in plasma was measured using the ELISA kit TD-80C (Teijin Co, Tokyo, Japan).35

**RESULTS**

Coagulation and fibrinolysis in DIC patients with APL. DIC associates frequently with APL even in the absence of severe infection and results in a high mortality from hemorrhage, especially intracranial bleeding.1 Although thrombin generation was reported in DIC with APL,34 neither activity and antigen levels of protein C nor antithrombin III levels decreased in our DIC patients with APL in the absence of sepsis. Because sepsis is one of the major triggers of DIC, we carefully selected DIC patients for those with APL and without sepsis.

As shown in Table 1, the α2-PI antigen and fibrinogen levels in cases of DIC with APL were significantly lower and the level of FDP was significantly higher than observed in the group of DIC patients with other diseases. These results suggested that a marked increase in secondary fibrinolysis occurs in DIC patients with APL. However, in these patients the consumption of antithrombin III and protein C, and the increase in PAs and plasmin-α2-PI complex levels were not as great as those found in the control DIC group.

**Plasma clot lysis induced by an MoAb against α2-PI.** In DIC, "secondary fibrinolysis" that occurs on the fibrin clots plays a role not only in the removal of formed clots but also in the prevention of fibrin deposition. To test the capacity for initiation of fibrinolysis in these patients, we observed secondary fibrinolysis by adding a small amount of thrombin and 1.2 μmol/L MoAb against α2-PI (MoAb JTPI-1) as described in Materials and Methods. This MoAb induces complete spontaneous clot lysis when incubated at 37°C for 10 hours after clotting, as described.36 Because the concentration of plasminogen in plasma affected this spontaneous clot lysis and this effect leveled off in the presence of more than 2 μmol/L plasminogen (data not shown), we added 2 μmol/L Glu-plasminogen to each plasma sample before clotting. As shown in Fig 1, the ratio of the amount of fibrin degradation products produced after 3 hours of incubation at 37°C relative to the antigen level of t-PA were significantly different in patients with different disease states. The ratios in patients with septicemia were remarkably low and

| Table 1. Various Parameters Relating to Coagulation and Fibrinolysis in DIC Patients |
|----------------------------------------|----------|----------------|-------------------|--------------------|
|                                      | DIC (APL-) (N = 43) | DIC (APL-) (N = 13) | Sepsis (N = 12) | Normal (N = 40) |
| AT III (%)                            | 68.8 ± 26.4 | 109.2 ± 12.3 | 98.6 ± 10.3 | 100 ± 10         |
| Fibrinogen (mg/100 mL)                | 277.6 ± 158.4 | 80 ± 40.7 | 414.1 ± 152.0 | 250 ± 50        |
| FDP (μg/mL)                           | 78.2 ± 68.1 | 217.1 ± 206.0 | <10              | <10             |
| α2-PI (%)                             | 76.3 ± 21.2 | 42.9 ± 11.2 | 95.5 ± 12 | 100 ± 10         |
| PIC (μg/mL)                           | 5.2 ± 6.3 | 2.2 ± 3.1 | <0.5 | <0.5             |
| u-PA (ng/mL)                          | 2.9 ± 1.6 | 3.3 ± 2.1 | 2.6 ± 1.7 | 2.2 ± 1.8        |
| t-PA (ng/mL)                          | 78.2 ± 68.1 | 110.0 ± 6.4 | 19.3 ± 15.0 | 8.92 ± 7.89      |

Plasma was analyzed as described in Materials and Methods. Values are mean ± SD.

Abbreviations: APL, without APL; ATIII, antithrombin III; FDP, fibrinogen and/or fibrin degradation products; PIC, plasma α2-PI complex; u-PA, urokinase-type plasminogen activator.
those in DIC patients without APL were relatively low in comparison with the level of t-PA. Interestingly, the ratios in DIC plasma with APL were not lower than those in normal plasma. Because this assay was intended to observe the regulation of the fibrinolysis in the initiation step by neutralizing antiplasmin activity, we examined the antigen and activity level of PAI-1.

Development of ELISA for total PAI-1 and t-PA/PAI-1 complex. We selected three MoAbs against PAI-1 (JTAI-1, JTAI-2, and JTAI-3) whose kds were similar for free PAI-1 or the t-PA/PAI-1 complex: JTAI-1, \( k_d = 9.7 \times 10^{-9} \text{ mol/L} \) for latent PAI-1 and \( 4.8 \times 10^{-9} \text{ mol/L} \) for complex; JTAI-2, \( k_d = 7.8 \times 10^{-9} \text{ mol/L} \) and \( 11.2 \times 10^{-9} \text{ mol/L} \); and JTAI-3, \( k_d = 4.7 \times 10^{-9} \text{ mol/L} \) and \( 12.3 \times 10^{-9} \text{ mol/L} \), respectively. Furthermore, these three MoAbs did not compete with one another for binding to free PAI-1 or t-PA/PAI-1 complex in a competition assay (data not shown).

To determine whether an ELISA using any pair of the MoAbs could detect both t-PA-unbound PAI-1 and the t-PA/PAI-1 complex with a similar efficiency, standard curves were constructed using PAI-1–depleted pooled normal plasma supplemented with either purified PAI-1 or t-PA/PAI-1 complex. In any combination of two MoAbs, the reactivities toward free PAI-1 and the t-PA/PAI-1 complex were at least one and a half times different (data not shown). Therefore, we used a mixture of JTAI-1 and JTAI-2 as the coating antibodies and peroxidase-labeled JTAI-3 to develop an assay for total PAI-1 antigen. Using this system, free PAI-1 and t-PA/PAI-1 complex were recognized equivalently when diluted in normal PAI-1–depleted plasma (Fig 2A). However, perhaps due to the double antibody-coating, this assay was not so sensitive (about 0.6 ng/mL). Because plasma had to be diluted at least 32 times to minimize the interference of plasma proteins and the interdilution coefficient of variation of four serial dilutions (4, 8, 16, and 32 times) of 20 different samples was 58%, the samples were diluted with PAI-1–depleted plasma. Under these modified conditions, the interdilution coefficient of variation of 20 different samples was 5.1%. When adding purified latent PAI-1 or t-PA/PAI-1 complex to PAI-1–depleted plasma of final concentrations of 10, 50, or 100 ng/mL, recoveries from this assay were 110% ± 5%, 95% ± 4%, and 101% ± 6% for latent PAI-1 and 95% ± 7%, 102% ± 7%, and 90% ± 4% for t-PA/PAI-1 complex, respectively (mean ± SD, n = 3). Inter-assay and intra-assay coefficients of variation were 7.9% and 5.2% (n = 10), respectively.

To develop an ELISA for the t-PA/PAI-1 complex, JTAI-1 was used as the coating antibody and peroxidase-labeled anti-t-PA MoAb (JTA-1) was used to detect t-PA antigen as it recognizes both free t-PA and the t-PA/PAI-1 complex with a similar efficiency. Figure 2B illustrates a dose-response curve of purified t-PA/PAI-1 complex diluted to varying concentrations in normal PAI-1–depleted plasma. In this sandwich ELISA, addition of a greater than 50-fold mol/L excess of free latent PAI-1 to the samples just before assay did not interfere with the measurement of t-PA/PAI-1 complex in the samples (data not shown), and the assay sensitivity was 0.2 ng/mL. The samples were diluted with PAI-1–depleted plasma to measure both t-PA/PAI-1 complex and total PAI-1 antigen. When adding purified t-PA/PAI-1 complex to final concentrations of 10, 50, or 100 ng/mL, recoveries were 105% ± 6%, 101% ± 5%, and 98% ± 6%, respectively (mean ± SD, n = 3). Inter-assay and intra-assay coefficients of variation were 6.8% and 4.9%, respectively (n = 10).

Development of modified ELISA to measure active PAI-1 antigen. t-PA forms a 1:1 stoichiometric complex with active PAI-1 but does not complex with latent PAI-1, and the amount of \( ^{125} \text{I}-\text{labeled t-PA/PAI-1 complex} \) generated by the addition of \( ^{125} \text{I}-\text{labeled t-PA} \) to the sample, correlates well with the conventional functional assay of PAI. Therefore, we determined the amount of active PAI-1 antigen by adding a fixed amount of t-PA to biologic fluids and measuring the newly generated t-PA/PAI-1 complex by the ELISA method described above. To determine the optimum t-PA concentration and incubation time, various concentrations of t-PA and aprotinin were added to samples of normal plasma supplemented with either purified PAI-1 or t-PA/PAI-1 complex.
SPECIFIC ACTIVITY OF PAI-1 IN DIC WITH APL

Fig 2. Dose-response curves of total PAI-1 and t-PA/PAI-1 complex and correlation between PAI activity and active PAI-1 antigen. (A) Various concentrations of purified latent PAI-1 (○) or t-PA/PAI-1 (●) complex were added to normal PAI-1-depleted plasma and their differential reactivities were determined by the ELISA for total PAI-1 as described in Materials and Methods. The abscissa shows the amount of latent PAI-1 and calculated PAI-1 in the complex assuming a 1:1 complex between t-PA and PAI-1. (B) Various concentrations of purified t-PA/PAI-1 complex were added to normal PAI-1-depleted plasma and concentrations of complex were determined as described in Materials and Methods. The data represent the mean of three experiments. (C) Urea- or guanidine-activated PAI-1 samples (two each) were diluted with either TBS-Tween containing 0.1% BSA (○) or PAI-1-depleted plasma (●) as described in Results, and active PAI-1 levels were determined by a conventional method (ordinate) and by our modified ELISA (abscissa). One unit of PAI is defined as the amount of PAI that neutralizes 2 ng of single-chain t-PA. (D) Ten sepsis patient plasma without DIC, 16 DIC patient plasmas, and 14 normal plasmas were collected as described in Materials and Methods, and active PAI-1 levels were determined by ELISA as described above.

ple samples that contained activated PAI-1 or to three different plasma samples and the amount of t-PA/PAI-1 complex was measured after various incubation times. Added t-PA rapidly complexed with PAI-1 in samples and the complex was stable at least 60 minutes at 37°C. A final concentration of 75 ng/mL single-chain t-PA and 500 U/mL aprotinin was sufficient to minimize proteolytic degradation of PAI-1 and dilution of samples (data not shown). Therefore, 75 ng/mL t-PA (final concentration) was added to samples and the mixtures were incubated for 30 minutes at 37°C.

To test the correlation between our assay and the conventional method using t-PA, plasminogen, fibrinogen fragments, and S2251 as described,12 purified PAI-1 was activated with urea (n = 2) or guanidine chloride (n = 2) for assay. Samples were adjusted to a similar level of activity as determined by the conventional method and diluted several times with either TBS Tween containing 0.01% BSA or PAI-1-depleted plasma. As shown in Fig 2C, there was a good correlation between PAI-1 antigen calculated from the generated complex and PAI activity determined by the conventional method. Furthermore, in the plasma of 10 nonpregnant patients with sepsis without DIC, 16 DIC plasmas, and 14 normals, there was also a good correlation between the two assay methods (Fig 2D).

Effect of complexation of PAI-1 to vitronectin and the decay of active PAI-1 to the latent form on the measurement of PAI-1 antigen. Because plasma vitronectin binds to active PAI-1 and apparently stabilizes PAI-1 activity twofold to threefold in blood,19 it was critical to know to what degree the ELISA for total PAI-1 measures vitronectin-complexed PAI-1. Furthermore, because there may be free active and latent PAI-1 in biologic fluids, it was also important to test the reactivities of the ELISA toward active and latent PAI-1. To these ends, total PAI-1 antigen and the time-dependent decay of active antigen levels were determined in the presence or absence of exogenous vitronectin (Fig 3A). The decay rate of activated PAI-1 (40% of the antigen was activated by measuring activity and antigen using the conventional activity method and our total PAI-1 method), purified from conditioned media of endothelial cells, was compared with that of inherent active PAI-1 present in conditioned media. The results were essentially the same.
active PAI-1 antigen in purified PAI-1, platelet releasates, PPP, or blood. (A) Purified guanidine-activated PAL1 (final concentration, 25 kg/mL) in TBS-Tween containing 0.1% BSA. At the indicated times alone (○) or mixed with purified VN (△) (final concentration 0). Platelet releasates were obtained from gel-filtrated platelets by treatment with thrombin (0.5 U/mL) for various times. At the indicated times, aliquots were removed and immediately cooled. The levels of total PAI-1 antigen (closed symbols) and active PAI-1 antigen (open symbols) were measured as described in Materials and Methods. The releasate from the different methods for aggregating platelets did not show any distinct differences in their rates of loss of PAI-1 activity. At 37°C, PAI-1 activity also declined at a similar rate (half-life: 3 hours) for normal plasma, DIC patient plasmas, or for normal citrated whole blood (Fig 3C).

Evaluation of plasma PAI-1 in patients. Total PAI-1 levels in plasma from DIC patients and from patients with sepsis were higher than those from normal controls (Fig 4A). However, the specific activity of PAI-1 (ratio of active PAI-1 to t-PA unbound PAI-1) in plasma from DIC patients, especially those with APL, was greatly reduced in comparison with that observed from sepsis patients or normal controls (Fig 4B). The level of t-PA-unbound PAI-1 was calculated by subtracting the value of 5/12 × (t-PA/PAI-1 complex) from the total PAI-1 concentration. During DIC, platelets are continuously activated by thrombin or other agonists in vivo and released PAI-1 from platelets may contribute to the increase in PAI-1 antigen. The specific activity of platelet PAI-1 determined by our ELISA method was about 6%, a result almost identical to that described by Declerck et al (about 5%). These results suggest that an increasing level of platelet PAI-1 released into the plasma may be one explanation for the observed low PAI-1 specific activities in DIC patients. However, because the APL patients had low platelet counts and an extremely low PAI-1 specific activity, the profiles of PAI-1 antigen in patient plasmas were examined by Western blotting (Fig 5). As shown in Fig 5, lane f, PAI-1 appeared to be cleaved, presumably by some protease(s) present in the patient plasma. Additionally, in DIC associated with APL and sepsis (Fig 5, lane h), the high molecular weight (MW, ~110,000) form of PAI-1, which also reacted with anti-t-PA antibody by Western blotting, and degraded forms of PAI-1 were present as well as an apparent overall increased level of PAI-1 antigen.

DISCUSSION

APL has been characterized by proliferation of morphologically abnormal promyelocytes and a severe bleeding diathesis. The bleeding diathesis is due mainly to DIC initiated by release of procoagulant activity from abnormal granules in the promyelocytes. Thrombin generation and using either form of PAI-1, although active PAI-1 in the conditioned media was 8.3% ± 2.1% (mean ± SD, n = 3) of total PAI-1 antigen. The level of active PAI-1 antigen decreased gradually in the absence of vitronectin (half-life: 1.5 hours), while in the presence of vitronectin the half-life was prolonged (3 hours) at 37°C. By using a sandwich immunoradiometric assay we could detect complexes of PAI-1 and vitronectin in the presence of exogenous vitronectin. In contrast, total PAI-1 antigen did not change even if activated PAI-1 decayed into the latent form, or if activated PAI-1 complexed with vitronectin (Fig 3A).

Because PAI-1 in plasma or platelet releasates were supposed to bind to VN, the effect of decay of active PAI-1 in these samples on the measurement of total PAI-1 antigen by this assay was determined at 37°C. The specific activity of PAI-1 from platelet releasates decayed with a half-life of 3 hours, whereas total PAI-1 antigen did not change (Fig 3B). The releasate from the different methods for aggregating platelets did not show any distinct differences in their rates of loss of PAI-1 activity. At 37°C, PAI-1 activity also declined at a similar rate (half-life: 3 hours) for normal plasma, DIC patient plasmas, or for normal citrated whole blood (Fig 3C).
the beneficial effect of heparin in the management of patients with APL were reported. However, if we carefully selected the DIC group with APL to eliminate patients with sepsis, which often accompanies the reduction of leukocyte count by chemotherapy against abnormal promyelocytes, the level of antithrombin III (Table 1) and protein C did not decrease as much as in DIC with other diseases. In this group, marked hyperfibrinolysis was noted (Table 1), and heparin had no beneficial effect on the bleeding diathesis in these patients (data not shown). Although we did not examine the possibility that abnormal promyelocytes produce antithrombin III and protein C, these results may imply that this hyperfibrinolysis is not simply explained by an increase in secondary fibrinolysis. To simulate "secondary fibrinolysis," thrombin and CaCl₂ were added to each patient plasma and the fibrinolytic potential in plasma was analyzed by simple clot lysis assay using MoAb JTPI-1, which neutralizes α2-PI activity and induces spontaneous clot lysis. α2-PI activity was neutralized in this assay so that the ratio of fibrinolysis to t-PA (FP ratio) reflects the PAI activity in plasma. Because PAI-1 is the primary physiologic inhibitor of t-PA in clot lysis, we measured the amount of PAI-1 in patient plasma to explain the different FP ratios in the DIC groups with APL or without APL, in the patient group with sepsisemia, and in normal controls (Fig 1). Because PAI-1 in biologic fluids exists in a free active, latent, VN-bound and t-PA-bound forms, total PAI-1 antigen levels have been difficult to measure using murine MoAbs, although a number of assay systems have been developed. For example, the Imulyse PAI-1 antigen assay from Biopool detects both active and latent forms of PAI-1, whereas the t-PA/PAI-1 complex is recovered with about 10-fold lower efficiency according to the manufacturer's specifications. By using two different MoAbs as the first antibodies in our assay system, we can measure free active, latent, t-PA-bound and VN-bound PAI-1 with similar efficiencies (Figs 2 and 3).

However, because it is almost impossible to purify only
inherently active PAI-1 from the conditioned media of HUVEC, we cannot exclude the possibility that the relatively low specific activity of the PAI-1 in the conditioned media may lead to the invisibility of an existing differential reactivity of this assay between inherent active PAI-1 and the other forms of PAI-1. A number of assay systems have been developed that estimate PAI activity in biologic fluid. For example, measurement of residual PA activity, either directly or by measuring plasminogen activating activity in the presence of fibrin fragments after adding a fixed amount of PA to the samples, is one of the typical PAI activity assays. However, these assays cannot discriminate PAI-1 activity from other PA inhibitors, in particular PAI-2 activity during pregnancy and PAI-3 in the presence of heparin. We can measure the concentration of active PAI-1 antigen specifically by measuring t-PA/PAI-1 complex after adding a fixed amount of t-PA to the samples. A similar approach for the measurement of active PAI-1 has also been described by several groups.

Any measurement of PAI-1 levels in biologic fluids must rely on an appropriate and stable standard material for accurate determinations. For functional PAI-1 assays, the choice of a standard is particularly complicated because it is a trace plasma protein whose activity decays into latency. In addition, PAI-1 is present primarily in the latent form in most conditioned media from cultured cells and in platelets. The current assay measures active PAI-1 by virtue of its ability to bind a known amount of added t-PA, and the standard was purified t-PA/PAI-1 complex. This design obviated the need to activate latent PAI-1 for a standard, and the complex is inherently more stable than t-PA–unbound active PAI-1 because the complex is acyl-enzyme complex. I-labeled t-PA/PAI-1 complex dissociated somewhat (from about 5% to 35%) under nonreduced conditions according to the size of the gel and the time of the electrophoresis by Lacmml's SDS-PAGE method, and this may be due to nucleophilic attack on the acyl-enzyme complex by the positively charged Tris ion at high pH as described. However, less than 5% of the complex dissociated by the Weber and Osborn SDS-PAGE method or by gel-filtration chromatography using a TSK-GEL G3000 SW column and a CCP-8010 HPLC system (Toso Soda, Tokyo, Japan) after 18 hours at room temperature or 2 months of storage at -20°C (data not shown). We cannot rule out the possibility that a small amount of degraded t-PA/PAI-1 complex may be generated by adding t-PA to the samples. In fact, the slope of the calibration curve between active PAI-1 levels measured with the current assay and those with the conventional method in a purified system (Fig 2C: slope = 0.65) or in patient plasmas (Fig 2D: slope = 0.57) was smaller than the estimated one (slope = 0.7) (Fig 2C). However, the correlation was satisfactory enough, so that we used the present modified ELISA method to analyze the total PAI-1 antigen, t-PA/PAI-1 complex, and active PAI-1 antigen in the patient plasmas.

The PAI-1 antigen levels increased in DIC patients and in patients with septicemia (Fig 4A), and the FP ratios in these patients were lower than those in normal controls (Fig 1). These results may suggest the retardation of clot lysis in vivo by the elevated PAI-1 in these diseases. The ratio in DIC patients was higher than that in patients with septicemia. This difference may be explained by the fact that the specific activity of PAI-1 in patients with DIC decreased significantly in comparison to patients with septicemia or in normal controls (Fig 4B). In DIC patients the PAI-1 antigen levels may increase because of release from platelets by thrombin and other agonists, or increased production by endothelial cells after endotoxin, interleukin, or ischemic stimulation. In general, the plasma from a DIC patient without APL contained a small amount of degraded PAI-1 as observed on Western blots. However, the majority of the PAI-1 migrated as an intact Mr 50,000 band. Because PAI-1 in the DIC plasma decayed into the latent form at a rate similar to PAI-1 in normal plasma (Fig 3C), it is unlikely that the low specific activity results from an instability of the active PAI-1. However, it is possible that this inactive PAI-1 originated from activated platelets, assuming increased platelet activation during DIC. Although the level of total PAI-1 antigen in DIC patient plasma with APL was higher than that in normal controls, the FP ratio was similar to that in normal control. This may be due to the low specific activity of PAI-1 in DIC patient plasma with APL (Fig 4B).

In DIC patient plasma with APL, marked hyperfibrinolysis in vivo was suggested by the increase of FDP and the decrease of fibrinogen and a2-PI activity, and the level of plasmin/a2-PI complex did not increase (Table 1). Because the antigen levels of PAs were not greatly elevated, hyperfibrinolysis in DIC with APL may be due to at least two factors: (1) proteolytic cleavage of fibrinogen, a2-PI, and PAI-1 (Fig 5) by leukocyte elastase or other proteases from abnormal promyelocytes; and (2) increased secondary fibrinolysis. There are several reports of decreased levels of either a2-PI or PAI-1, and the patients had bleeding tendencies. In these cases, the antigen levels of t-PA were normal and treatment of these individuals with epsilon-aminocaproic acid was effective. This suggestion is further supported by the degraded forms of PAI-1 observed in plasmas from DIC patients associated with APL and sepsis (Fig 5, lanes g and h). In one case (Fig 5, lane h), the combined effects of endotoxin and APL protease on the marked increase of t-PA and PAI-1 and proteolysis of PAI-1 might result in the appearance of t-PA/PAI-1 complex and the degraded forms of PAI-1.

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The specific activity of plasminogen activator inhibitor-1 in disseminated intravascular coagulation with acute promyelocytic leukemia

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