Bleeding Diathesis Due to Decreased Functional Activity of Type 1 Plasminogen Activator Inhibitor

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Abstract

We evaluated an elderly patient with a lifelong history of severe bleeding after surgery or trauma and with evidence of persistent hyperfibrinolysis. Routine coagulation studies were normal. Serum plasminogen (40%, normal 72–128%) and α2-antiplasmin (55%, normal 70–145%) activities were decreased. Euphlobulin clot lysis was abnormally shortened (50 min) and normalized in vitro with L-aminocaproic acid (EACA). The patient was treated with EACA with prompt cessation of bleeding. Patient tissue-plasminogen activator (t-PA) levels in serum were normal (4.7 ng/ml, control 3.5–7.2) as detected by a two-site immunoradiometric assay (IRMA). Patient fibrinolytic inhibitor activities were assessed by incubating 125I-labeled t-PA with either whole blood or serum followed by SDS-PAGE and autoradiography to identify the resultant protease/protease inhibitor complexes. In comparison to blood samples obtained from normal donors, patient plasma and serum demonstrated reduced binding of a fast-acting plasminogen activator inhibitor to 125I-labeled t-PA. Immunoprecipitation experiments indicated diminished complex formation between type I plasminogen activator inhibitor (PAI-1) in patient serum and 125I-labeled t-PA. Low patient PAI-1 activity was confirmed in serum (0.36 U/ml, control 0.87–1.81; n = 3) and in platelet lysates using a functional IRMA to quantitate PAI-1 binding to immobilized t-PA. However, patient serum PAI-1 antigen was within the normal range when analyzed by IRMA (31.8 ng/ml, control 19.6–42.2); this result was confirmed in both serum and platelets by Western blot (n = 3). Mixing experiments using purified PAI-1 as well as patient and control sera did not show evidence for an inhibitor against PAI-1. We conclude that this patient’s bleeding diathesis was due to hyperfibrinolysis and defective PAI-1. This patient provides the first demonstration of a link between decreased in vivo PAI-1 activity and disordered hemostasis, and supports a role for PAI-1 in control of in vivo fibrinolysis.

Introduction

Regulation of fibrinolysis is critical during hemostasis, wound repair, neoplasia, inflammation, and other biologic processes (1–3). Proteolytic degradation of fibrin clots is mediated by the enzyme plasmin. Plasmin is formed in the circulation from the zymogen plasminogen through the proteolytic action of plasminogen activators (PAs).1 PAs are highly specific serine proteases found in normal or neoplastic tissues and can be divided into two distinct groups, tissue-type (t-PA) and urokinase-type (u-PA) plasminogen activator (4). The principal physiologic inhibitor of plasmin is α2-antiplasmin. Inhibitors of plasminogen activation also appear to play an important role in the physiologic control of fibrinolysis (5). At least four immunologically distinct molecules with plasminogen activator inhibitor (PAI) activity have been identified (6). Type 1 PAI (PAI-1), formerly termed the endothelial cell PA inhibitor, inhibits both t-PA and u-PA by forming 1:1 stoichiometric complexes with them. It has been detected in a large variety of cells cultured in vitro, and is also found in plasma and in the α-granules of blood platelets (5–7). A number of experimental observations support the hypothesis that PAI-1 is the principal physiologic inhibitor of t-PA and a primary regulator of in vivo fibrinolysis (6–8). In this study we report our investigations of an elderly man with a lifelong history of postoperative bleeding who was found to have sustained hyperfibrinolysis. Our results suggest that his bleeding diathesis is due to decreased functional activity of PAI-1.

Methods

Case report

The patient is a 76-yr-old man seen in consultation for persistent bleeding after transurethral resection of the prostate. He had a long history of bleeding after surgery. In 1969 severe postoperative bleeding occurred after total hip replacement. In 1978 revision of his hip prosthesis was associated with postoperative bleeding requiring 10 U of packed red blood cells. From 1978 to 1981, five episodes of upper gastrointestinal bleeding occurred requiring multiple blood transfusions. No specific site of bleeding was identified despite upper gastrointestinal series and endoscopy. In 1984, the patient fell on his side and developed a massive hemoptoma; the prothrombin time (PT), partial thromboplastin time (PTT), and platelet count were all normal. In August 1986, he developed obstructive urinary symptoms and underwent a transurethreal resection of the prostate. Pathology revealed benign hyperplasia. The postoperative course was complicated by bleeding requiring 4 U of packed red blood cells. The PT, PTT, platelet count, and bleeding time were all normal. Bleeding resolved but the patient was readmitted 3 d after discharge with persistent gross hematuria. Cystoscopy was negative and clot was evacuated. The patient continued to bleed and was treated with 8 U of red blood cells and 8 U of fresh-frozen plasma without cessation of bleeding. Consultation was requested. Routine coagulation studies were normal including the PT and PTT. There was no evidence of ongoing disseminated intravascular...

1. Abbreviations used in this paper: EACA, e-aminocaproic acid; IRMA, immunoradiometric assay; PA, plasminogen activator; PAI-1, type 1 plasminogen activator inhibitor; p-APMSF, p-amidinophenylmethylsulfonyl fluoride; PT, prothrombin time; PTT, partial thromboplastin time; t-PA, tissue-type plasminogen activator; u-PA, urokinase-type plasminogen activator.
lar coagulation with normal fibrinogen (256 mg/dl, normal 160–350), thrombin and reptilase times and D dimer, and only modestly (20–40, normal < 10) elevated fibrin degradation products. The bleeding time and von Willibrand factor antigen, ristocetin cofactor, and factor VIII coagulant activity were normal, suggesting that a diagnosis of von Willibrand’s disease was unlikely. Although routine coagulation tests were normal, there was evidence of ongoing fibrinolysis with a shortened euglobulin clot lysis time (50 min, normal no lysis at 2 h) which partially corrected in vitro (1 h, 45 min) after the addition of antifibrinolytic agents, e-aminoacapric acid (EACA). Additionally, serum plasminogen (40%, normal 72–128%) and α2-antiplasmin (55%, normal 70–145%) were persistently decreased. The patient was presumed to have a hyperfibrinolytic state of undetermined etiology and was treated empirically with EACA with prompt resolution of his bleeding. When seen in the clinic over the next 2 wk to 10 months, he continued to have laboratory evidence of hyperfibrinolysis with decreased levels of α2-antiplasmin and plasminogen, despite no clinical evidence of bleeding. Plasma urokinase by radioimmunoassay was normal. An intensive search for a primary or metastatic tumor, including numerous radiographic and laboratory tests, was unrevealing.

Review and resection of the patient’s excised prostate showed no evidence of malignancy. The patient’s antithrombin III level in stored serum from his hospital admission was normal (91%, normal range 86–120%), thus providing further evidence against disseminated intravascular coagulation; interestingly, his α2-antiplasmin (55%) and plasminogen (68%) levels continue to be abnormally low 2 yr after his diagnosis of disordered fibrinolysis while his antithrombin III levels continue to remain within the normal range.

Both of the patient’s parents were deceased at the time of evaluation and the patient had no siblings. The patient’s mother was known to have had unexplained severe hemorrhage after a hysterectomy. The patient has two children; both sons are alive and well and have no history suggestive of a bleeding dyscrasia. However, neither son has undergone any surgical procedure nor experienced severe trauma. Both sons have normal serum levels of plasminogen and α2-antiplasmin.

Procedures

Preparation of plasma, serum, and platelets. Whole-blood serum was prepared by collecting blood into glass tubes, allowing the blood to clot at 37°C for 1 h, then removing the clot by centrifugation at 1,800 g for 20 min at 4°C. The resulting serum was collected and stored at −70°C until used. Platelet-poor plasma was prepared by collecting blood in 3.8% sodium citrate, followed by centrifugation at 1,800 g for 30 min at 4°C. Platelet-rich plasma was obtained by centrifugation of citrated blood at 160 g for 15 min at 23°C. The platelet-rich plasma was then centrifuged at 660 g for 15 min at 23°C and the platelet pellet resuspended in calcium-free Heps-Tyroside buffer, pH 6.5. Platelets were washed by gel-filtration through Heps-Tyroside buffer, pH 7.3, as described (9). Alternatively, the platelet pellet was diluted to 105 platelets/ml and lyed by addition of 1.0% Triton X-100.

Reagents and fibrinolytic proteins. Natural human t-PA was purified from media conditioned by human melanoma cells, according to published protocols (10). Recombinant t-PA was purified from media conditioned by transfected Chinese hamster ovary cells (11). The t-PA International Standard (82/517) was kindly provided by the National Institute for Biological Standards and Controls (London). Goat anti-serum to t-PA was obtained from Bio-Pool (Horserows, Sweden) and the IgG fraction was prepared as described (10). Bovine and human PAI-I were purified as described from media conditioned by either bovine aortic endothelial cells or by a transformed human lung fibroblast cell line (SV40 W138 VA13 2RA) (12). PAI-I was isolated primarily in latent form and activated by treatment for 1 h at 37°C with 4 M guanidinium-hydrochloride followed by dialysis at 4°C (12). PAI-I activity was quantitated by measuring its ability to inhibit t-PA-mediated lysis of 125I-labeled fibrin (13, 14). One unit of PAI-I is defined as the amount required to inhibit the activity of 2 IU of t-PA by 50%. Antiserum to purified PAI-I was raised in New Zealand rabbits and was shown to be monospecific by immunoprecipitation and by two-dimensional gel electrophoresis (15). Monoclonal antibody (MAB 2D2) directed against human PAI-I was produced according to described protocols (10). Thrombin was a generous gift from Dr. J. Fenton (New York State Department of Health, Albany, NY). Pancreatic elastase was purchased from Elastin Products Co. (Pacific, MO). Molecular mass markers for SDS-PAGE were purchased from Sigma Chemical Co. (St. Louis, MO) and included myosin (205,000 D), α-galactosidase (116,000 D), BSA (66,000 D), egg albumin (45,000 D), and carbonic anhydrase (29,000 D).

Blood proteinase inhibitor complex assay. This assay was performed to assess the ability of inhibitors in whole blood to form complexes with 125I-labeled t-PA (13-16) (PAI-1-t-PA) (11). Purified recombiant t-PA was radiolabeled with Enzymobeads (Bio-Rad Laboratories, Richmond, CA) as described (11). Freshly drawn, citrated whole blood (500 μl) was incubated with 2 × 105 cpm of 125I-labeled enzyme at 37°C for 0–24 h. Plasma samples were then removed and centrifuged to remove cells. Less than 5% of the label was associated with the cellular fraction. Samples were diluted with 10 vol of 2% SDS buffer and analyzed by SDS-PAGE (500 cpm per lane). Complexes were detected by autoradiography.

Formation and immunoprecipitation of 125I-t-PA/PAI-I complexes in serum. Whole-blood serum (30 μl) was incubated with 125I-t-PA (106 cpm) for 0–60 min at 37°C. The reaction was rapidly terminated by the addition of either SDS sample buffer or the synthetic serine protease inhibitor; phosphoseryl-methylsulfonyl fluoride (pAPMSF; 10 mM final concentration). Samples treated with SDS sample buffer were analyzed without further treatment by SDS-PAGE and autoradiography, while samples treated with pAPMSF were immunoprecipitated according to the procedures described by Erickson et al. (16). Briefly, either antisemur against PAI-I (40 μl) or nonimmune serum (40 μl) was added to protein A–coated Sepharose CL-4B beads (80 μg Pharmacia, Uppsal, Sweden) that had been rehydrated and washed according to the manufacturer’s instructions. 125I-t-PA–treated serum samples were added to the beads, diluted into buffer, and then incubated at 37°C for 1 h in a total volume of 250 μl. The beads were removed by centrifugation, the supernatant fluid was collected, and the beads were washed three times by centrifugation with 1 ml of phosphate-buffered saline (PBS). Proteins bound to the beads were extracted by incubation with an equal volume of SDS-sample buffer. The resulting immunosupernatant fluids and/or the extracts from the immunoprecipitates were analyzed by SDS-PAGE followed by autoradiography.

Immunoradiometric assays (IRMA). To assess t-PA antigen, samples were diluted in IRMA buffer (PBS, 3% BSA, 5 mM EDTA, 0.1% Tween 80, and 0.02% NaN3) and incubated for 1.5 h at 37°C in microtiter wells precoated with goat anti-t-PA (50 μl per well, 10 μg/ml). Plates were blocked with 3% BSA and washed three times after each step with PBS containing 0.1% BSA, 0.05% NaN3, and 0.05% Tween 80. Bound t-PA was quantitated radiometrically after incubation with rabbit anti-t-PA, followed by 125I-labeled goat anti-rabbit IgG (105 cpm per well) (13). PAI-I activity was quantitated by a t-PA binding assay (17). Briefly, samples and standard curves of purified PAI were prepared in IRMA buffer and incubated for 1 h at 37°C in microtiter wells precoated with purified t-PA (50 μl per well, 1 μg/ml, overnight at 4°C). Plates were blocked and washed three times following each step as described above. PAI bound to t-PA–coated wells was detected radiometrically after incubation (1.5 h, 37°C) with rabbit antiserum to PAI-I (1:75 dilution, 50 μl per well) followed by 125I-labeled goat anti-rabbit IgG (2.5–5 × 105 cpm per well; 1.5 h at 37°C). PAI-I antigen was quantitated by a two-site IRMA: purified monoclonal antibody (2D2) against human PAI-I (5 μg/ml) in PBS was coated onto microtiter wells (4°C for 16 h). After incubation with 3% BSA and washing, samples diluted in IRMA buffer were incubated in the antibody-coated wells at 37°C for 1.5 h. Bound PAI-I was quantitated by incubation with rabbit anti–PAI-I antibody followed by 125I-labeled goat anti-rabbit IgG. In some experiments, serum samples were first mixed with purified PAI-I (100 μl of 3.25 U/ml) and incubated at 37°C for 15 min before assay for PAI-I activity. Circulating t-PA/PAI-I complexes were assayed in a two-site IRMA using goat anti-t-
PA as the immunoabsorbent and rabbit anti–PAI-1 as the detecting antibody (3).

Immunoblotting. Immunoblotting for PAI-1 antigen was performed as described (14). Samples were first subjected to SDS-PAGE. Proteins were then electrophoretically transferred to nitrocellulose using a buffer containing 50 mM Tris base, 95 mM glycine, 20% methanol, and 0.01% SDS. The nitrocellulose sheets were soaked in PBS containing 1% casein for 1 h at 23°C to block additional protein binding sites and were then incubated overnight at 4°C in 1% casein containing rabbit anti–PAI-1 (t:500 dilution). The nitrocellulose was washed three times and then incubated for 2 h at 23°C with 125I-labeled goat anti–rabbit IgG (2 × 107 cpm/ml). After washing, the nitrocellulose sheets were dried and exposed to XAR x-ray film (Eastman Kodak Co., Rochester, NY) at −70°C.

Results

Enhanced in vivo fibrinolysis has been associated with excess release of endothelial plasminogen activator (t-PA) (18, 19). Patient serum was assayed on three separate occasions by IRMA for t-PA antigen. Patient t-PA antigen levels were within normal range (Table I). To determine if excessive fibrinolysis was mediated by deficient PAI activity, we examined the ability of inhibitors in the patient’s blood to form SDS-stable complexes with 125I-labeled t-PA. In this analysis, patient or control whole blood was incubated with radiolabeled t-PA for 0–24 h at 37°C followed by centrifugation, separation of plasma t-PA-inhibitor complexes on SDS polyacrylamide gels and autoradiography (Fig. 1). The three bands at the top of the gel are complexes of t-PA with α2-antiplasmin. The band in the middle of the gel represents α2-antiplasmin complexes. The upper band of the doublet is an unidentified protein which also binds t-PA. The lower bands in the gel are free 125I-labeled t-PA. The formation of 125I-t-PA inhibitor complexes in both the patient and control blood are basically identical except for the lower band of the doublet at 92,000 D which is missing in the patient’s samples (arrow). The absence of this 125I-t-PA–inhibitor complex band at early time points (i.e., 0.25 h) suggested that the patient’s bleeding diathesis may be caused by abnormally low levels of a fast-acting PAI.

The presence of low amounts of the 125I-t-PA–fast-acting inhibitor complex in control plasma is consistent with the observation that normal plasma contains quite low levels of fast-acting PAI activity, whereas normal serum and platelets usually exhibit high levels of this inhibitor (6, 19). To analyze in further detail the activity of the patient’s fast-acting PAI, we incubated 125I-t-PA for short periods of time in serum prepared from the patient and from normal donors. Fig. 2A indicates that the addition of 125I-t-PA to normal human serum results in the rapid formation of a 92,000-D band. In comparison, less 125I-t-PA–inhibitor complex (92,000 D) was formed when 125I-t-PA was incubated in serum prepared from the patient’s blood (Fig. 2B). Thus, the majority of 125I-t-PA added to the patient’s serum was found to migrate in parallel with free 125I-t-PA (Fig. 2A, lane 1) even after 0.5–1-h incubations (e.g., Fig. 2B, lanes 6 and 7), whereas the amount of free 125I-t-PA was markedly reduced after a 15-min incubation with control serum (i.e., Fig. 2A, lane 5).

Immunoprecipitation experiments were performed to demonstrate that this 125I-t-PA–inhibitor band was composed of 125I-t-PA bound to PAI-1 (Fig. 3). For this purpose, 125I-t-PA was again incubated (30 min, 37°C) with both control and patient serum, the reaction was terminated by the addition of PAPMSF and the samples were immunoprecipitated with either rabbit anti–PAI-1 or nonimmune serum. Supernatants of 125I-t-PA-treated patient serum (lane 1) or control serum (lane 3) immunoprecipitated with nonimmune serum contained both free 125I-t-PA and a 92,000-D 125I-t-PA–inhibitor complex, whereas supernatants of samples immunoprecipitated with anti–PAI-1 serum contained only free t-PA (lanes 2 and 4, respectively). As described above, the patient samples contained fewer 125I-t-PA–inhibitor complexes than the controls. Elution of the protein A-Sepharose bead immunoprecipitates with SDS-sample buffer revealed a small amount of free 125I-t-PA using both nonimmune and anti–PAI-1 serum. However, a 92,000-D band also was eluted from the patient and control samples immunoprecipitated with rabbit anti–PAI-1 (lanes 7 and 9, respectively). Less 125I-t-PA–inhibitor complex was recovered from the patient immunoprecipitate than from control. These data suggest deficient complex formation between patient PAI-1 and 125I-t-PA.

To quantitate deficient patient PAI-1 activity, a number of additional immunologic assays were performed. When patient PAI-1 functional activity was assessed in an independent immunoradiometric t-PA binding assay, it was found to be mark-
Figure 2. Analysis of normal serum and patient serum for fast-acting PAI activity. Whole blood serum was prepared from blood collected from either a normal human donor (A) or the patient (B). $^{125}$I-t-PA in PBS (lane 1: $10^4$ cpm) was incubated at 37°C in the respective serum samples (30 μl) for the indicated times (lane 2, 0 min; lane 3, 5 min; lane 4, 10 min; lane 5, 15 min; lane 6, 30 min; lane 7, 60 min). Samples were subjected to SDS-PAGE on a 4–10% polyacrylamide gradient following by autoradiography. Positions of molecular mass standards in kilodaltons are indicated.

Figure 3. Immunoprecipitation of $^{125}$I-treated control serum and patient serum with antibodies to PAI. $^{125}$I-t-PA (10$^4$ cpm) was incubated in either patient serum (lanes 1, 2, 6, 7) or control serum (lanes 3, 4, 8, 9) for 30 min at 37°C. The reaction was terminated by the addition of pAPMSF (10 mM, final concentration) and immunoprecipitated with either nonimmune rabbit serum (NRS; lanes 1, 3, 6, 8) or antiserum against PAI-1 (lanes 2, 4, 7, 9) as described in Methods. The resulting immunosupernatant fluids (lanes 1–4) or extracts from the immunoprecipitates (lanes 6–9) were analyzed by SDS-PAGE (4–10% gradient) followed by autoradiography. Lane 5 represents free t-PA. Positions of molecular mass markers in kilodaltons are indicated.

dedly reduced (0.36±0.12 U/ml) compared to control donors (0.87±1.81 U/ml; n = 5) (Table I). Thus by both IRMA and by immunoprecipitation experiments, patient PAI-1 demonstrates diminished functional activity as evidenced by an inability to form normal complexes with t-PA. This decreased activity was found in both serum and plasma, as well as in platelet-rich plasma and platelet lysates (0.01 U/ml PAI-1 activity in patient platelet lysates vs. 0.09–0.11 U/ml in control lysates, n = 2). However, PAI-I antigen levels were found to be within the normal range in serum (Table I) and in platelet lysates. Patient circulating t-PA–PAI-1 complexes were modestly reduced (1.9 ng/ml vs. 2.5–5.3 ng/ml in control plasma; n = 2).

A number of proteases (e.g., thrombin, activated protein C, elastase) (6) could potentially have reduced patient PAI-1 activity either via complex formation with PAI-1 or by direct cleavage of the inhibitor. We assessed these possibilities initially by Western blot analysis of patient serum and plasma. As mentioned above, little PAI-1 antigen is present in control platelet-poor plasma under normal conditions (Fig. 4 A, lanes 2, 4, 8, and 10). In contrast, PAI-1 in serum from four normal donors (Fig. 4 A, lanes 3, 5, 9, and 11), as well as PAI-1 in patient serum (lane 7), was detected primarily at the molecular mass of 46,000 D. Unfortunately, the large amount of albumin present in plasma and serum samples distorts the electrophoretic mobility of PAI-1 (50,000 D) to a slightly lower molecular mass (6), thus potentially interfering with one's ability to distinguish between the native form and a proteolytically cleaved form of PAI-1 in this system. To accurately examine the molecular characteristics of the patient’s PAI-1, we also analyzed patient and control platelet lysates for PAI-1 by immunoblotting. Fig. 4 B indicates that the patient's platelet-derived PAI-1 migrated as a single band (50,000 D; lane 6) in parallel with the PAI-1 present in control platelets (lane 1). A high-M$_r$, immunoreactive band could be detected by this technique if t-PA was incubated with the platelet lysates prior to electrophoresis (lanes 2 and 7). Furthermore, cleavage of PAI-1 was also observed following a 10-min incubation of control or patient platelet lysates with either thrombin (lanes 3 and 8) or elastase (lanes 4 and 9), respectively. Thus, no evidence is provided by Western blot to indicate that patient PAI-1 had either been cleaved or had complexed with a protease.

Circulating antibodies against plasma PAI-1 could explain the patient's decreased serum PAI-1 functional activity. This possibility was addressed in three separate sets of experiments. Purified PAI-1 was incubated for 15 min at 37°C with an equivalent volume of either buffer, patient, or control serum and then assayed for PAI-1 activity in the immunoradiometric t-PA binding assay (Table II). Control serum containing normal endogenous PAI-1 produced the expected augmentation in PAI-1 activity. Patient serum had little demonstrable effect on the activity of purified PAI-1, thus demonstrating again minimal patient PAI-1 serum activity and providing no evidence for an inhibitor to PAI-1 in patient serum. Similarly, patient serum had no inhibitory effect on PAI-1 activity when incubated with control serum before activity determinations. In separate experiments, a direct assay for the presence of antibodies to PAI-1 was performed. Purified PAI-1 was bound to microtiter plates, then incubated with patient plasma followed by radiolabeled goat anti-human IgG. Utilizing this assay, no antibodies to PAI-1 were detected in patient plasma (data not shown).
circulating cess associated with (20–22). Lifelong to patients with 65%.

a2-antiplasmin antigen (20, 23) almost causes a diathesis. Disordered fibrinolysis are described in Methods. Patient samples are in lanes 6 and 7, while control samples from four different normal donors are in lanes 2 and 3, 4 and 5, 8 and 9, and 10 and 11, respectively. Under normal conditions, little PAI-1 antigen is detected in control platelet-poor plasma. (B) Control platelets (lanes 1–4; 40 µl per lane) or patient platelets (lanes 6–9; 40 µl per lane) were lysed with Trisonic X-100 and incubated (10 min, 37°C) in the absence or presence of various proteases: lanes 1 and 6, PBS alone; lanes 2 and 7, 10 ng t-PA; lanes 3 and 8, 10 U thrombin; lanes 4 and 9, 1 µg elastase. Lane 5 contained a mixture of t-PA, thrombin, and elastase in the absence of platelet lysates. Samples were subjected to SDS-PAGE and immunobotted for PAI-1 antigen. Positions of molecular mass markers in kilodaltons are indicated.

Discussion

Disordered fibrinolysis has infrequently been associated with a bleeding diathesis. Homozygous α2-antiplasmin deficiency causes a severe hereditary diathesis and is associated with almost no plasma antiplasmin functional activity and little detectable α2-antiplasmin antigen (20, 21). Our patient’s α2-antiplasmin antigen and activity levels ranged from 52% to 65%. Patients with heterozygous α2-antiplasmin deficiency infrequently have a history of excessive bleeding and, in contrast to our patient, have normal serum levels of plasminogen (20–22). Lifelong bleeding disorders have also rarely been associated with persistent evidence of hyperfibrinolysis and excess circulating levels of t-PA (17, 18). Our patient had normal circulating levels of t-PA antigen, measured on three separate occasions, normal levels of urokinase antigen, and did not have an increase in circulating levels of t-PA–PAI-1 complexes. Additionally, excess release of plasminogen activators would not be expected to deplete functional PAI-1 compartmentalized in platelets, nor would a primary excess release of t-PA be likely to explain low platelet PAI-1 activity coupled with normal platelet PAI-1 antigen by immunoblotting analysis.

Low PAI-1 activity, confirmed by two independent methods of assay, in combination with normal levels of t-PA and PAI-1 antigen, suggests a qualitative defect in PAI-1. The explanation for the decreased functional activity of this patient’s PAI-1 is unclear. PAI-1 interacts not only with plasminogen activators but also with a wide variety of other proteolytic enzymes (1, 2, 4). It is thus possible that excessive protease activity in the patient’s blood might have inactivated PAI-1, before assay, by either proteolytic cleavage or by the formation of inactive protease-PAI-1 complexes. Immuno-blotting analysis (Fig. 4) of plasma and serum, as well as analysis of platelet-derived PAI-1, was particularly helpful in eliminating this possibility since it indicated that both patient and control PAI-1 migrated similarly on SDS-PAGE.

Another potential explanation for decreased blood PAI-1 activity is the presence of circulating inhibitors to PAI-1. Franci et al. (23), in a preliminary report, have described a patient with amyloidosis, accelerated fibrinolysis, and a bleeding diathesis. This patient had normal t-PA antigen levels but PAI-1 activity was undetectable owing to the presence of neutralizing antibodies to PAI-1. The patient also had depressed levels of both plasminogen and α2-antiplasmin and, like our patient, responded to the therapeutic administration of EACA.

Functional and immunologic assays coupled with the use of purified PAI-1 failed to demonstrate an anti-PAI-1 inhibitor in our patient’s serum or plasma.

Our data suggest that diminished PAI-1 activity in this patient is most probably associated with an abnormality in the PAI-1 molecule itself. Further experiments will be required to determine the exact defect in PAI-1. An abnormality in the patient PAI-1 molecule could be due to alternative processing of PAI-1 mRNA or to a defect in the 12.2-kb PAI-1 gene located on the long arm of chromosome 7 (6, 24).

Elevated levels of serum PAI-1 have been associated with a

Table II. Effect of Patient Serum on Activity of Purified and Blood-derived PAI-1

<table>
<thead>
<tr>
<th>Sample</th>
<th>PAI-1 activity (U/ml)</th>
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<tbody>
<tr>
<td>PAI-1 (5 ng) + buffer*</td>
<td>1.07±0.13*</td>
</tr>
<tr>
<td>PAI-1 (5 ng) + control serum</td>
<td>2.12±0.23</td>
</tr>
<tr>
<td>PAI-1 (5 ng) + patient serum</td>
<td>1.15±0.10</td>
</tr>
<tr>
<td>Buffer + control serum</td>
<td>1.01±0.17</td>
</tr>
<tr>
<td>Patient serum + control serum</td>
<td>0.95±0.20</td>
</tr>
<tr>
<td>Patient serum + buffer</td>
<td>0.14±0.05</td>
</tr>
</tbody>
</table>

* Purified PAI-1 was incubated with 100 µl of either buffer or serum for 15 min at 37°C before IRMA for PAI-1 activity.

1 Data represent mean±SD from triplicate determinations.

2 Serum and/or buffer samples (100 µl) were mixed and incubated at 37°C for 15 min before IRMA for PAI-1 activity.
postulated hypofibrinolytic state and with an increased risk for thrombosis (25, 26), suggesting that PAI-1 may play an important role in mediating vascular fibrinolysis. Most t-PA in blood is complexed to PAI-1. Since t-PA antigen levels in our patient were normal, these data suggest that the decrease in PAI-1 activity disrupts the normal fibrinolytic balance in plasma, resulting in a relative increase in free t-PA and accelerated fibrinolysis. The successful treatment of our patient’s hemorrhagic episode with EACA, an antifibrinolytic agent which blocks t-PA, plasminogen, and plasmin binding to fibrin substrate, supports this conclusion (27). The evidence for persistent accelerated fibrinolysis in our patient (moderate consumption of both α2-antiplasmin and plasminogen, as well as a shortened euglobulin lysis time) suggests that ongoing production of functionally active PAI-1 is necessary to complex with small amounts of continuously released free plasminogen activator. In a similar manner, recent data suggests that activated procoagulants such as factor Xa and Va are continuously formed within the vascular system and are opposed by natural anticoagulant mechanisms (28).

We conclude that our patient’s bleeding diathesis was most likely due to hyperfibrinolysis and deficient PAI-1 activity. The data suggest that decreased PAI-1 activity results in an increase in free t-PA, which in turn causes spontaneous plasmin generation and predisposes to excessive bleeding. To our knowledge, the evidence presented provides the first demonstration of a link between decreased PAI-1 activity and disordered hemostasis, and supports a role for PAI-1 in the control of in vivo fibrinolysis.

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References