Plasminogen deficiency causes severe thrombosis but is compatible with development and reproduction

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Plasminogen (Plg)-deficient mice were generated to define the physiological roles of this key fibrinolytic protein and its proteolytic derivatives, plasmin and angiostatin, in development, hemostasis, and reproduction. Plg−/− mice complete embryonic development, survive to adulthood, and are fertile. There is no evidence of fetal loss of Plg−/− mice based on the Mendelian pattern of transmission of the mutant Plg allele. Furthermore, embryonic development continues to term in the absence of endogenous, sibling-derived, or maternal Plg. However, Plg−/− mice are predisposed to severe thrombosis, and young animals developed multiple spontaneous thrombotic lesions in liver, stomach, colon, rectum, lung, pancreas, and other tissues. Fibrin deposition in the liver was a uniform finding in 5- to 21-week-old mice, and ulcerated lesions in the gastrointestinal tract and rectal tissue were common. A remarkable finding, considering the well-established linkage between plasmin and the proteolytic activation of plasminogen activators, was that the level of active urokinase-type plasminogen activator in urine was unaffected in Plg−/− mice. Therefore, Plg plays a pivotal role in fibrinolysis and hemostasis but is not essential for urokinase proenzyme activation, development, or growth to sexual maturity.

[Key Words: Plasminogen, gene targeting, thrombosis, fertility, proenzyme activation]

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Mignatti et al. 1986; Ossowski et al. 1991b; Crowley et al. 1993; Kook et al. 1994). The recent finding that uPA- and tPA-deficient mice develop and exhibit relatively modest phenotypic abnormalities, other than an age-related risk of thrombosis, suggests that one important role of these PAs is directing fibrinolysis (Carmeliet et al. 1994). However, considerable uncertainty still remains as to the roles of Plg outside fibrinolysis because other PAs of unknown biological significance have been identified, including plasma kallikrein, (Kluft et al. 1987), factor XIIa (Goldsmith et al. 1978), and factor XIa (Mandle and Kaplan 1979). In addition, although Plg disorders are encountered frequently in some populations [Iichino et al. 1991; Kikuchi et al. 1992], no homozygotes with total Plg deficiency have ever been documented. Finally, potent biological activities have been associated with the noncatalytic portions of Plg, including the angiogenesis- and metastasis-suppressing activity of the Plg derivative, angiostatin, which is made up solely of kringle domains (O’Reilly et al. 1994).

To directly establish the role of Plg and its derivatives in development, hemostasis, reproduction, and tumor cell invasion, we have disrupted the Plg gene in mice. We report that Plg-deficient mice are viable and fertile but experience severe thrombotic problems with an early onset.

Results

Disruption of the Plg gene

To compromise expression and secretion of all portions of the Plg polypeptide a replacement-type targeting vector was constructed using a combination of 5’-flanking and intron 2 sequences from the cloned murine Plg gene (see Fig. 1A). Incorporation of this targeting vector into the murine genome by homologous recombination results in a 9-kb deletion within the Plg gene that includes proximal promoter sequences, exon 1 [encoding the signal peptide], and exon 2 [encoding 43 residues of the “preactivation peptide”]. The targeting vector was introduced by electroporation into the HPRT-deficient embryonic stem (ES) cell line E14TG2a (Hooper et al. 1987) and 2 of 46 stable transfectants picked for analysis were found to have incorporated the transgene by homologous recombination based on both PCR and Southern blot hybridization analysis of genomic DNA extracts (data not shown). Each of these targeted ES cell clones was microinjected into 3.5-day-old C57Bl/6 blastocysts to raise chimeric founder transgenic mice. Four of six transgenic males that sired litters passed the mutant Plg gene to their offspring. Crosses of heterozygous mice resulted in the birth of mice that were homozygous for the mutant Plg allele (Plg−/−), 83 (49%) were heterozygous (Plg+/−), and 45 (26%) were homozygous for the wild-type Plg allele (Plg+/+).

Homozygous mutant mice do not express detectable Plg mRNA or plasma Plg

Plg expression is highly liver-specific and represents a major fraction of total protein secreted into the plasma (Robbins 1992). Therefore, hepatic Plg mRNA was measured by Northern blot hybridization analysis of total RNA extracts from 5-week-old Plg−/−, Plg+/−, and Plg+/+ mice (Fig. 2A). Plg mRNA could be detected easily in RNA preparations from both Plg+/+ and Plg+/− mice. The mRNA levels in Plg+/− livers were approximately half that of Plg+/+ livers based on quantitative PhosphorImager analysis (data not shown). However, no hybridizing transcript of the size of Plg mRNA could be detected in Plg−/−-derived liver RNA samples when analyzed in parallel [Fig. 2A]. Even very long exposures of blots to PhosphorImaging screens indicated little hybridizing material throughout the Plg−/− RNA lane. Based on the sensitivity of the PhosphorImaging system in detecting Plg mRNA in control liver samples, the level of any hepatic Plg transcripts derived from the mutant Plg gene was estimated to be at least 3000-fold less than the Plg mRNA level found in control livers. In addition, no Plg transcripts could be detected in either RNA extracts from adult brain, heart, spleen, lung, and kidney or RNA extracts from entire Plg−/− newborns, suggesting that there are no secondary sites of expression of the mutant Plg gene in Plg−/− mice (Fig. 2B).

Plg enzymatic activity was analyzed in whole plasma by using a gel zymography assay. In this assay system, the casinolytic activity of plasmin is detected after electrophoretic separation of plasma proteins by SDS-gel electrophoresis. The Plg zymography assay is very sensitive and offers the advantage that the many plasmin inhibitors found in plasma [i.e., a1-antitrypsin (54 kD), α2-antiplasmin (70 kD), and α2-macroglobulin (725 kD)] are electrophoretically separated from Plg (92 kD) prior to activity detection. A plasminogen activator-stimulated casinolytic activity with an apparent molecular weight identical to Plg was readily detectable in whole plasma collected from Plg+/+ and Plg+/− mice [Fig. 3A]. This casinolytic activity appears to be plasminogen-mediated based on the diminished activity in parallel zymography gels that were cast without PA. In contrast, no PA-stimulated casinolytic activity was observed in whole plasma samples from Plg−/− mice [Fig. 3A]. Given that Plg activity could be detected in Plg+/−-derived samples containing as little as 8 nl of plasma, and no PA-activatable casinolytic activity could be detected in as much as 5 μl of plasma from Plg−/− mice, these data indicate that any plasma Plg activity in the Plg−/− mice must be at least 600-fold less than Plg+/+ mice. The well-known affinity of Plg for lysine–Sepharose (Deutsch and Mertz 1970) was also exploited to establish whether any residual Plg antigen could be detected in plasma collected from Plg−/− mice. Although plasma fractions eluted from lysine–Sepharose with the lysine
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Figure 1. Targeting of the Plg gene by homologous recombination. (A) Structure of the Plg targeting vector, wild-type Plg allele, and targeted Plg allele. Exons are indicated as black boxes, the 5' flanking and intron sequences are hatched. Sequenced areas are indicated with arrows. Exon 1 and 2 sequences determined from the cloned Plg gene were identical to the published sequence of the mouse Plg cDNA (Degen et al. 1990). The phosphoglycerate kinase promoter (PGK)-HPRT minigene cassette (van der Lugt et al. 1991) was introduced in the opposite orientation to the Pig gene and replaces a 9-kb portion of the Pig gene, including the 5'-flanking sequences and the first two exons. The locations of primers used for screening ES cell clones by PCR analysis (see Materials and methods) are indicated with arrows. The downstream Plg primer was complementary to sequences outside those used in the targeting vector. The probe used for Southern blot analyses of tail biopsy DNA also was complementary to Plg sequences outside the targeting vector (bottom). The hybridizing EcoRI fragments expected from the endogenous and correctly targeted Plg alleles are indicated. (B) Southern analysis of EcoRI-digested tail DNA from offspring of Plg+/+ x Plg+- mice. The examples shown were from mice genotyped as wild-type (Plg+/+), heterozygous (Plg+-), and homozygous mutant (Plg-/-) based on PCR analyses (see Materials and methods). The size and relative position of molecular mass marker fragments are shown (left).

The endogenous Plg gene appears to eliminate Plg gene transcription and gives rise to a Plg-null phenotype.

Hematological parameters in Plg-deficient mice

The loss of plasma Plg does not have any significant impact on general hematological parameters when measured in 4- to 5-week-old mice. In particular, no major

analog, e-amino caproic acid, were markedly enriched for Plg antigen in samples from Plg+/+ mice, no Plg antigen was detected in Western blot analyses of lysine-Sepharose eluates or whole plasma from Plg-/- mice (Fig. 3B). Based on the sensitivity of the Western blot assay for Plg antigen, Plg levels in plasma samples from Plg-/- mice were, again, more than two orders of magnitude below those in Plg+/+ mice. Thus, the deletion introduced in

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Figure 2. Absence of detectable Plg mRNA in mice homozygous for the targeted Plg allele. (A) Northern blot of total liver RNA from two wild-type mice (lanes 1, 2), two heterozygous mice (lanes 3, 4), and two mice homozygous for the targeted allele (lanes 5, 6). Positions of 28S and 18S rRNAs are indicated. Ethidium bromide staining of the agarose gel prior to blotting (right) indicated similar RNA loading in all cases. (B) Northern blot of total RNA extracted from various tissues collected from 4-week-old Plg−/− (lanes 3–7) and Plg+/+ (lanes 9–13) mice, and from whole newborn Plg−/− (lane 2), and Plg+/+ (lane 8) mice. Brain (lanes 3, 9), heart (lanes 4, 10), spleen (lanes 5, 11), lung (lanes 6, 12), and kidney (lanes 7, 13). Total liver RNA isolate from an adult Plg+/+ mouse was analyzed in parallel as a positive control. Similar amounts of RNA were loaded as indicated by the ethidium bromide-stained gel (right).

Differences were found between control and Plg−/− mice with regard to platelet counts, red and white cell counts, and hematocrit (Table 1). In addition, no differences were found in plasma levels of uPA or tPA based on gel zymography of whole plasma (data not shown). Furthermore, thrombin time measurements (Table 1) and pre-
Figure 3. Absence of Plg activity and antigen in plasma of Plg-targeted mice. (A) Gel zymography of fivefold serial dilutions of plasma from Plg<sup>+/+</sup> mice (lanes 1–5), Plg<sup>+/−</sup> mice (lanes 6–10), and Plg<sup>−/−</sup> mice (lanes 11–15). Gels were cast with (top) and without (bottom) 20 IU/ml of human uPA. (B) Immunoblot analysis of Plg<sup>−/−</sup>-derived whole plasma (1 μl, lane 1), Plg<sup>+/+</sup>-derived whole plasma, (1 μl, lane 12), and fivefold serial dilutions of plasma fractions eluted from lysine-Sepharose (Plg<sup>+/+</sup> eluate, lanes 1–5; Plg<sup>−/−</sup> eluate, lanes 6–10). Plg was detected using a polyclonal antibody raised against rat plasminogen as described in Materials and methods. The relative sizes and the positions of protein molecular mass standards are indicated.

liminary measurements using a fibrinogen-specific enzyme-linked immunosorbent assay (ELISA) (K. Holm-bäck and T. Bugge, unpubl.) indicated that plasma fibrinogen levels in Plg<sup>−/−</sup> and Plg<sup>+/+</sup> mice were comparable. Thus, Plg deficiency per se does not result in appreciable anemia, thrombocytopenia, or change in steady-state fibrinogen levels. However, it remains to be established whether older mice that develop fibrin deposition in tissues (see below) also develop secondary alterations in these hematological parameters.

Proteolytic activation of uPA in urine of Plg-deficient mice

Plasmin is generally thought to have a physiologically important positive feedback on plasminogen activation by converting inactive, single-chain pro-uPA to active, two-chain uPA. However, the factors that contribute to initial pro-uPA activation and plasmin formation are unclear (Ichinose et al. 1986; Kobayashi 1991). The urine is a rich source of uPA, and a large fraction is two-chain uPA in normal animals (Kielberg et al. 1985; Sappino et al. 1991). To determine whether Plg contributes signifi-

Table 1. Hematological analysis of Plg-deficient transgenic mice

|                       | Plg<sup>+/+</sup> mice | Plg<sup>−/−</sup> mice |
|-----------------------|------------------------|------------------------|
| Blood analysis        |                        |                        |
| platelets (× 10<sup>4</sup>/liter) | 979 ± 130 (3)         | 953 ± 48 (4)          |
| RBC (× 10<sup>12</sup>/liter)  | 7.5 ± 0.7 (3)         | 7.3 ± 0.6 (4)         |
| WBC (× 10<sup>9</sup>/liter)   | 2.6 ± 1.3 (3)         | 2.9 ± 0.9 (4)         |
| hematocrit (%)        | 40.1 ± 1.3 (3)        | 44.3 ± 3.2 (4)        |
| hemoglobin (grams%/%) | 12.3 ± 0.5 (3)        | 13.1 ± 1.3 (4)        |
| Plasma thrombin time* | 13.1 ± 1 sec (3)      | 13 ± 2 sec (4)        |

Data presented is the mean ± s.d. with the number of mice analyzed shown in parentheses. All mice analyzed were 4–5 weeks of age. Control wild-type mice were adult Black Swiss.

*Thrombin times were measured in reaction mixtures containing equal volumes of plasma and 20 U/ml of bovine thrombin in PBS.
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cantly to the formation of this two-chain uPA in the urogenital tract, urine from Plg<sup>-/-</sup> and Plg<sup>+/+</sup> mice was compared using esterase and zymography activity assays and Western blot immunoassay. Urine samples were also collected from uPA<sup>-/-</sup> mice (Carmeliet et al. 1994) to control for the specificity of each assay. Surprisingly, the enzymatic activity of uPA in urine collected from Plg<sup>-/-</sup> and Plg<sup>+/+</sup> mice was similar (Fig. 4A, data not shown). Furthermore, the proportion of high molecular weight (45,000) and low molecular weight (30,000) uPA measured by gel zymography and immunoblot of nonreduced samples was similar between normal and Plg-deficient animals (Fig. 4B,C). While these assays indicate that the generation of low molecular weight uPA in the urogenital tract is not dependent on plasmin, they do not indicate the fraction of high molecular weight uPA initially present in a two-chain form. This question was addressed by immunoblot assay of reduced samples. In samples from both control mice and Plg<sup>-/-</sup> mice, reduced 45,000 high molecular weight uPA was largely separated into 30,000 heavy chain (Fig. 4C, right; data not shown) and 15,000 light chain (immunologically not detected; Kielberg et al. 1985) fragments. Therefore, at least in the urogenital system, plasmin does not appear to be essential for the conversion of pro-uPA to enzymatically active forms of uPA.

Figure 4. Enzymatically active two-chain uPA in Plg-deficient mice. (A) uPA-specific esterase activity in representative bladder urine samples collected from Plg<sup>+/+</sup>, Plg<sup>+-</sup>, and Plg<sup>-/-</sup> mice. Bladder urine from a uPA<sup>-/-</sup> mouse was assayed in parallel to establish assay specificity. (B) uPA activity detected by gel zymography of representative bladder urine samples (1 μl analyzed) collected from Plg<sup>+/+</sup> (lane 2), Plg<sup>+-</sup> (lane 3), and Plg<sup>-/-</sup> (lane 4) mice. Bladder urine from a uPA<sup>-/-</sup> mouse was analyzed in parallel (lane 1) to control for assay specificity. (C) Immunoblot detection of uPA in nonreduced (lanes 1–4) and reduced [lanes 5–8] bladder urine samples (5 μl analyzed) collected from Plg<sup>+/+</sup> (lanes 1,5), Plg<sup>+-</sup> (lanes 2,6), and Plg<sup>-/-</sup> mice (lanes 3,7). Bladder urine from a uPA<sup>-/-</sup> mouse was analyzed in parallel to control for specificity (lanes 4,8). Similar results were obtained in three independent experiments with bladder and void urine sample sets collected from Plg<sup>+/+</sup>, Plg<sup>+-</sup>, Plg<sup>-/-</sup>, and uPA<sup>-/-</sup> mice using all three assays (data not shown). The relative sizes and positions of protein molecular weight standards are indicated at right. [HMW uPA] High molecular weight uPA; [LMW] low molecular weight uPA.
Survival and fertility in Plg-deficient mice

Forty-three Plg−/− mice have now been maintained for up to 21 weeks with no obvious phenotypic abnormalities observed in the majority of animals. However, inflammatory rectal lesions were found in 7 Plg−/− mice (one at 5 weeks, two at 9 weeks, three at 13–14 weeks, and one at 21 weeks of age), possibly secondary to fibrin deposition and vasocclusion (see below). A rectal prolapse was observed in the oldest of these mice. Two 8- to 9-week-old mice were found to have penile lesions. With the exception of these nine mice, no differences have been observed between Plg−/− mice and littermate controls with regard to either outward appearance, behavior, weight gain, or survival characteristics (data not shown). Neither running nor premature death was evident in the 5-month window examined that extends well into adulthood. Furthermore, Plg−/− mice are fertile, with males siring multiple litters and females carrying their litters to term. Seven litters have been raised in test crosses between Plg−/− males and Plg+/+ females, and four litters have been raised between Plg+/+ males and Plg−/− females. The average litter sizes in these test crosses were 7.6 and 7.8 pups, respectively. These values are comparable to the average litter size of 8.1 recorded with control breeding pairs in our colony. Two test crosses between mates both genotyped as Plg-null were also productive, with four and five Plg−/− mice born. The generation of offspring from Plg−/− parents is particularly significant in that it demonstrates clearly that Plg is not essential for development to term, the development of offspring from crosses of this type have no access to either endogenous or exogenous (maternal- or sibling-derived) Plg.

Pathological consequences of Plg deficiency

Five Plg−/− mice with overt rectal lesions (see above) and 12 apparently healthy Plg−/− mice, including five juvenile animals (5–7 weeks old) and seven young adults (10–17 weeks old), were sacrificed for a detailed histological examination of tissues. Age-matched Plg+/+ and Plg−/− mice were analyzed in parallel as controls. A uniform feature in all 17 Plg−/− mice examined microscopically was the presence of multiple hepatic fibrin deposits that were recognized as fibrillar material in hematoxylin/eosin-stained sections (Fig. 5A) that immunolabeled intensely using a polyclonal antibody raised against mouse fibrinogen (Fig. 5E,F). These deposits showed varying degrees of organization with infiltration of spindle cells, possibly macrophages or fibroblasts (Fig. 5D). Necrosis was apparent within adjacent tissue in only a fraction of the hepatic lesions, suggesting that these deposits were generally not extensive enough to impair perfusion in these young mice. Although these deposits were found throughout the liver, they tended to be more numerous in the subcapsular region, possibly because of surface trauma (for examples, see Fig. 5, D and F). In addition, the lesions were more extensive in terms of number, size, and degree of adjacent necrosis in the older mice examined. No such deposits were found in either hematoxylin/eosin-stained (Fig. 5A) or anti-fibrinogen-immunostained (Fig. 5G; data not shown) livers collected from age-matched control mice.

Other sites of fibrin deposition and organ damage were also common in Plg−/− mice. Microscopic analysis of rectal tissues from Plg−/− mice with visible rectal lesions uniformly revealed anal–rectal tissue ulceration with surface exudate [mixture of necrotic debris, neutrophils, and bacteria] and surrounding fibrin deposition (e.g., see Fig. 6A,B). However, similar lesions were also discovered in four animals without gross rectal lesions (data not shown), implying that fibrin deposition and necrosis may precede and predispose to ulceration. Gastric ulcers were seen in 10 of the 17 Plg−/− mice examined (ranging from 5 to 21 weeks of age), including 7 mice with ulcerated lesions in the glandular portion of the stomach (e.g., see Fig. 6C,D), 2 with an ulcer in the squamous portion of the stomach (e.g., see Fig. 6E), and 1 with an ulcer at the junction of the two portions (data not shown). Colonic ulcers, proximal to the rectum, were seen in three Plg−/− mice (7 and 10 weeks of age) (data not shown). In each of these lesions, the surface epithelium was necrotic and covered by exudate, the underlying tissue was necrotic with diffuse fibrin deposition in and around the lesions, and the adjacent mucosa was reactively altered with epithelial hyperplasia and intravascular fibrin deposition in the lamina propria. The appearance of these ulcers is consistent with ischemic necrosis secondary to fibrin occlusion of small vessels. Seven mice (10–21 weeks old) showed pulmonary lesions, with capillary fibrin thrombi in four mice and organizing patches of alveolar fibrin in four animals (e.g., see Fig. 6F). In one 11-week-old mouse, organizing fibrin deposits were observed in multiple pancreatic lobules (Fig. 6G), with one centrally located lesion associated with lobule necrosis (data not shown). Fibrin deposits were detected in the thymus of one 5-week-old animal and in the adrenal tissue of one 14-week-old animal (Fig. 6H). Finally, the ovary of a 14-week-old female had fibrin deposits in small vessels, and the uteri of three females housed with males contained large aggregates of fibrin, possibly remnants of earlier or involuting pregnancies (data not shown). Brain, heart, spleen, small intestine, kidney, testicle, bone, muscle, tongue, oropharynx, and miscellaneous connective tissue were also surveyed, and no lesions were detected.

Discussion

The studies presented here provide direct evidence that Plg and its proteolytic derivatives, plasmin and angiostatin, are not essential for either mouse development, growth, or fertility. Each of these processes was unimpeded under conditions where no trace of residual Plg could be detected within a sensitivity range extending nearly three orders of magnitude below normal. Notably, any Plg available at this level would be of little biological significance given that the expected Plg concentrations would fall many orders of magnitude below the 2 μM $K_m$.
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Figure 5. Fibrin deposition in livers of Plg-deficient mice. (A) Hematoxylin/eosin-stained liver section from a control, Plg+/+ mouse. Magnification bar, 100 μm. (B) Hepatic fibrin deposits (arrows) in the liver of an 8-week-old Plg−/− mouse. Large fibrin patches contain necrotic hepatocytes. Magnification bar, 100 μm. (C) Fibrin deposition in the liver of a 10-week-old Plg−/− mouse with entrapped hepatocytes and infiltration of spindle cells (arrows) indicating early organization. Magnification bar, 35 μm. (D) Hepatic fibrin deposition in subcapsular sinusoids (arrows) of an 11-week-old Plg−/− mouse. Magnification bar, 1.5 μm. (E) Immunoperoxidase-stained sinusoidal fibrin(ogen) in a liver section from a 10-week-old Plg−/− mouse. The deposits (arrows) were rounded and scattered throughout the parenchyma. Magnification bar, 35 μm. (F) Higher magnification of immunostained sinusoidal fibrin deposits (arrows) in a subcapsular region similar to that illustrated in D. Magnification bar, 1.5 μm. (G) Immunostain of fibrinogen in a liver section from a control, 10-week-old Plg+/+ mouse indicating the normal sinusoidal staining pattern and the lack of luminal accumulation. Magnification bar, 35 μm. (H) Liver section from a Plg−/− mouse treated with preimmune serum prior to immunoperoxidase labeling.

for PAs when bound to either fibrin or the cell surface (Ellis and Danø 1992; Collen and Liinen 1994). Thus, Plg-mediated proteolysis either plays no fundamental role in developmental cell migration and tissue remodeling events or other extracellular proteolytic systems can fully compensate for the loss of Plg. Although a surprising number of partial functional overlaps between related proteins have been identified by using gene targeting (for examples, see Lohnes et al. 1993; Rudnicki et al. 1993; Carmeliet et al. 1994), no other secretory protease of similar specificity and abundance in the circulation stands out as a possible compensatory enzyme. The results reported here strongly imply that all facets of plasmin-mediated proteolysis, including degradation of extracellular matrices, and activation of proenzymes and latent growth factors, are not strictly needed for ovula-
Figure 6. Fibrin deposition in other organs of Plg-deficient mice. (A) Longitudinal section of the rectum at the squamocolumnar junction in a 14-week-old Plg\(^{-/-}\) mouse showing fibrin deposition and ulceration (arrows) with surface exudate and reactive hyperplasia (asterisk). Magnification bar, 350 μm. (B) Higher magnification of A showing fibrin deposition and surface ulceration. Magnification bar, 100 μm. (C) Ulceration in the glandular portion of the stomach (arrows) of a 14-week-old mouse. Magnification bar, 350 μm. (D) Higher magnification of C showing capillary fibrin deposition (arrows) at the base of the ulcer. Magnification bar, 35 μm. (E) Ulceration in the upper portion of the stomach of a 11-week-old mouse with reactive hyperplasia in the adjacent squamous mucosa (asterisks). Note fibrin deposition at the base of the ulcer (arrow). Magnification bar, 350 μm. (F) Fibrin deposition (arrows) in an area of organization in the lung of a 10-week-old mouse. Magnification bar, 35 μm. (G) Fibrin deposition in the periacinar capillaries of the pancreas (arrows) of an 11-week-old mouse. The ingrowth of spindle cells indicates early organization. Magnification bar, 100 μm. (H) Interstitial fibrin deposition (arrows) in thymus outlining areas of thymocyte dropout. Magnification bar, 100 μm.

Fibrin deposition is a common feature of Plg\(^{-/-}\) mice. The general predisposition to thrombosis documented here in Plg\(^{-/-}\) mice is reminiscent of the phenotype found in PA-deficient [PA\(^{-/-}\)] mice [Carmeliet et al. 1994] but with some notable differences. First, the various histological abnormalities in PA\(^{-/-}\) mice were only seen in animals older than 2-3 months of age and most significantly in ill, "preterminal" mice. In contrast, the Plg\(^{-/-}\) mice uniformly develop hepatic fibrin deposits by 5 weeks of age, and several of these recently weaned mice...
had thrombotic ulcerations in the stomach and rectal tissue as well as other thrombotic lesions. Second, stomach ulcers were found in more than half of the Plg-/- mice examined, but neither this nor thrombotic lesions in the pancreas, thymus, or adrenal tissue were described in PA-/- mice. Finally, fibrin deposits were documented in kidney, skin, ear, and small intestines of PA-/- mice, whereas these tissues were apparently free of fibrin deposits in the 5- to 21-week-old Plg-/- mice examined. Some of these differences may be related to differences in genetic background, the variable penetrance of specific thrombotic events in both PA-/- and Plg-/- lines, and/or the youth and overall good health of the Plg-/- mice that have been collected so far for histological analysis. However, it cannot be excluded at present that any phenotypic differences between PA-/- and Plg-/- mice may be related to the availability of alternative Plg activators or to noncatalytic functions of Plg in uPA- and tPA-deficient mice. The long-term impact of Plg deficiency on thrombotic complications and life span remains to be determined.

The development of fibrin deposits in Plg-/- mice is consistent with the severe thrombotic problems reported in patients identified with dysplasminogenemia (Aoki et al. 1978; Ichinose et al. 1991; Kikuchi et al. 1992; Robbins 1992; Azuma et al. 1993) and hypoplasminogenemia (Lottenberg et al. 1985; Girolami et al. 1986; Mannucci et al. 1986; Dolan et al. 1988; Patrassi et al. 1993), which include recurring venous thrombosis, pulmonary hypertension, retinopathy, and stroke. However, the total deficit of Plg established in transgenic mice is distinct from that in PA-/- and Plg-/- mice may be related to the availability of alternative Plg activators or to noncatalytic functions of Plg in uPA- and tPA-deficient mice. The long-term impact of Plg deficiency on thrombotic complications and life span remains to be determined.

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Although the multiple sites of fibrin deposition observed in Plg-/- mice might result from a primary defect in tissue development and architecture, no obvious alterations have been found under either gross or microscopic examination. Therefore, one hypothesis that emerges from this study is that the fundamental, and possibly only essential, physiological role of the Plg activation system is fibrinolysis. The finding that uPA and tPA functionally complement each other in vivo (Carmeliet et al. 1994) is consistent with this general notion. uPA may primarily participate in cell-associated fibrinolysis via its interactions with the high-affinity uPA receptor (Vassalli et al. 1991), and tPA may mediate cell-independent and cell-associated clot lysis via its fibrin binding (Collen and Lijnen 1994) and receptor binding (Cesarman et al. 1994) properties. The diverse contexts in which PAs are expressed may simply reflect the diverse contexts in which cells might encounter and must cope with fibrin. The known expression of PA by macrophages (Vassalli et al. 1992), keratinocytes (Morioka et al. 1987), capillary endothelial cells (Pepper 1987), granulosa cells (Sappino et al. 1989), renal tubule cells (Sappino et al. 1991), and many other cell types fits this overall concept. A critical tool in rigorously establishing any putative role of Plg outside of fibrinolysis will be mice lacking both fibrinogen and Plg. Abnormalities associated with Plg deficiency that are secondary consequences of a reduced ability to clear fibrin can be defined by comparing the phenotypes of Plg-/- mice with fibrinogen-free Plg-/- mice. In this regard, fibrinogen-deficient mice are already available [T.T. Suh and J.L. Degen, unpubl.], and we have begun the appropriate crosses to generate mice with combined Plg and fibrinogen deficiency.

If the only essential physiological role of Plg is fibrinolysis, then this does not preclude other important roles of Plg in disease pathophysiology. Notably, a linkage between PA or PA receptor expression and tumor cell invasiveness in vitro and in vivo is well established (Nosowski and Reich 1983; Dano et al. 1985, 1994; Mignatti et al. 1986; Nosowski 1988; Nosowski et al. 1991a,b; Crowley et al. 1993; Kook et al. 1994). Conceivably, tumor cells may depend or capitalize on the ability of plasmin to degrade both fibrin and other extracellular matrix components in the penetration of tissue barriers. Similarly, development of atherosclerotic lesions may depend on plasmin-mediated degradation of vessel wall components by smooth muscle cells. Each of these concepts can now be tested directly in vivo by breeding Plg-/- mice to existing lines genetically predisposed to either malignant tumor formation (Guy et al. 1992) or atherosclerosis (Plump et al. 1992).

Plg-/- mice also provide an opportunity to define physiological activators of single-chain uPA and tPA without the confounding background activity of plasmin. The finding that the level of activated, two-chain and low molecular weight uPA in the urine is unaffected by Plg deficiency argues that one or more proteases, other than plasmin, is available in the urogenital tract which is capable of activating pro-uPA and cleaving the light chain to produce enzymatically-active low molecular weight uPA. Proteases that are known to activate pro-uPA in vitro include plasma kallikrein, factor XIIa, and cathepsin B (Ichinose at al. 1986; Kobayashi et al. 1991). Because cathepsin B is known to activate both free and receptor-bound pro-uPA, cleaves uPA at the same site as plasmin (Kobayashi et al. 1991), and is expressed...
by proximal tubule cells and secreted into the urine (Olbricht et al. 1991), this enzyme is a strong candidate for a urinary pro-uPA activating agent. However, direct studies are necessary to resolve which proteases activate uPA in the urogenital tract and to establish whether plasmin-independent PA activation also occurs outside the urogenital system.

Materials and methods

Construction of targeting vector and generation of transgenic mice

A bacteriophage ADASHII genomic DNA library, prepared using DNA from 129/SvJ strain mice, was screened by in situ hybridization (Suh et al. 1994) with a 580-bp EcoRI–NsiI fragment from the mouse Pig cDNA (Degen et al. 1990) encoding the amino-terminal portion of Pig through the second kringle domain. Three positive isolates were found to contain both exons 1 and 2 by Southern blot analysis (Smith et al. 1980) with the following 32P-labeled exon 1- and 2-specific oligonucleotide probes: OligoEx1-5’, 5’-CGGTGCTGTTGGCCAGTCCC-3’; and OligoEx2-5’, 5’-GGGAGACTCGCTGGATGGC-3’. Hybridizing DNA fragments from one recombinant bacteriophage, λPlg-1, were subcloned into Bluescript II (Stratagene) plasmid vectors, and overlaps between each subclone were established based on restriction endonuclease cleavage sites and Southern blot analyses. DNA sequence was determined using plasmid templates (Suh et al. 1994) and both the Ex1-5’ and Ex2-5’ primers, and confirmed that the cloned gene encoded Pig; the gene organization over the first three exons was identical to the human Pig gene (Petersen et al. 1990), and the exon sequences were identical to the corresponding published mouse Pig cDNA sequences (Degen et al. 1990). The targeting vector was constructed by placing a 3.2-kb SaII cassette encoding HPRT (van der Lugt et al. 1991) between a 4-kb SacI–SmaI fragment derived from 5’-flanking sequences and a 1-kb PstI fragment derived from intron 2 sequences. The Pig targeting vector was further modified with an herpes simplex virus thymidine kinase [HSV tk] cassette to provide a means of selection against cells that randomly inserted the transgene into the genome (Mansour et al. 1988). The targeting vector was introduced into E14Tg2a ES cells [Hooper et al. 1987] by electroporating (800 V/cm and 200 μF; IBI GeneZapper) 105 cells suspended in 750 μl of media containing 40 μg of linearized targeting vector in a 0.4-cm cuvette. The cells were maintained on mitomycin C–treated primary mouse embryonic feeder fibroblasts (Li et al. 1994) for 24 hr and then placed in selection media containing 0.1 mm hypoxanthine, 16 μM thymidine, 0.4 μM aminopterin (HAT supplement; Gibco), and 2 μM ganciclovir (Syntex). After 7–11 days, 46 ES cell clones were picked and expanded in 24-well dishes, and DNA extracts were tested by PCR using an oligonucleotide primer complementary to a portion of Pig intron 2 that was not included in the targeting vector (Oligon2, 5’-AATGGCCAGGCATCTTTCAAGGC-3’) and a primer complementary to the HPRT cassette (OligoHP, 5’-GGGAGCTCCCTCCCTAGCTCACC-3’). Two isolates produced a 1100-bp PCR product matching the size expected for cells incorporating the targeting vector by homologous recombination. Pig gene targeting was verified in each of these isolates by Southern blot hybridization analysis (Li et al. 1994) of EcoRI- and KpnI-digested genomic DNA using a 170-bp PstI–EcoRI intron 2 probe that was external to the targeting vector sequences. Pig-targeted ES cells were injected into the blastocoe1 cavity of C57Bl/6-derived blastocysts and implanted into pseudopregnant females (Li et al. 1994). Chimeric male offspring were bred to NIH Black Swiss females (Taconic) to generate hemizygous offspring. These mice were subsequently interbred to generate homozygous Pig+/− progeny.

Genotype analyses

Genotypes of mice were established using tail biopsy DNA preparations [Li et al. 1994] by either Southern blot analysis of EcoRI digests (see above) or PCR analysis. In the PCR assays, mice with one or two targeted Pig alleles were identified using reaction mixtures containing the primers, Oligon2 and OligoHP (see above). Mice with one or two wild-type Pig alleles were identified using reaction mixtures containing the exon 2-specific primers OligoEx2-5’ (see above) and OligoEx2-3’ [5’-TTCACATTGGGCAAAACGT-3’], which yield a PCR product of 108 bp.

Hematological analysis

Blood was collected into one-tenth volume of 0.129 M sodium citrate anticoagulant (Becton Dickinson) by insertion of a 27-gauge hypodermic needle into the inferior vena cava of mice anesthetized with 0.1 ml per 30 gram of body weight of ketamine/xylazine/acepromazine (4:1:1). Blood cell counts and hemocrit were established using a Technicon H-1 blood cell analyzer. Plasma was prepared from citrated blood by centrifugation at 2500g for 10 min at room temperature. The thrombin-stimulated clotting time was determined using plasma samples preincubated for 2 min at 37°C in glass test tubes. The reaction mixtures were prepared by combining equal volumes of plasma and 20 U/ml of bovine thrombin diluted in PBS [Pacific Hemostasis].

Northern blot analysis

Total RNA was extracted from tissues frozen in liquid nitrogen using the guanidinium thiocyanate method (Chomczynski and Sacchi 1987). RNA concentrations were determined spectrophotometrically using an extinction coefficient of 20 at 260 nm for a 1 mg/ml solution. Samples (40 μg) were electrophoretically fractionated on denaturing agarose gels, blotted onto BA85 nitrocellulose filters (Schleicher & Schuell), and hybridized with [Bell et al. 1990] to a 32P-labeled [1.6 × 106 to 2.4 × 106 cpm/μg] 900-bp EcoRI fragment from the mouse Pig cDNA subclone MPL2 [Degen et al. 1990]. To control for sample loading, rRNAs (28S and 18S) were visualized by UV transillumination of ethidium bromide-stained gels. Hybridized material was visualized and quantitated using phosphor screens and a Molecular Dynamics PhosphorImager equipped with ImageQuant software.

Zymographic analysis of PA and Pig

Plasma was prepared from citrated whole blood as described above. PA was detected by gel zymography as described previously [Heckel et al. 1990]. Pig was detected by using the same experimental procedure as for PA detection except that 20 IU/ml of human high molecular weight uPA [American Diagnostics] was cast in the polyacrylamide gels rather than human Pig.

Western blot analysis

Proteins were fractionated by SDS–polyacrylamide gel electrophoresis (8% acrylamide) and transferred to Immobilon P membranes [Millipore] as described previously [Heckel et al. 1990]. Pig was detected with a sheep anti-rat Pig antisera that cross-
reacts with mouse plasminogen [kindly provided by E. Reich]. Mouse uPA was detected using affinity-purified rabbit anti-mouse uPA antibodies [Larsson et al. 1984]. Bound primary antibody was detected using alkaline phosphatase-conjugated dog anti-sheep IgG or goat anti-rabbit IgG [Sigma] and the 5-bromo-4-chloro-3-indolyl phosphate [BCIP]/nitro blue tetrazolium [NBT] [Sigma] staining system.

Histology and fibrinogen immunostaining of tissue sections

Immunostaining was performed using the Vectastain Elite ABC into paraffin, sectioned, and stained by standard techniques. In sample buffer containing 2% SDS for gel fractionation and repeated eight times with the OD at 280 nm of the third wash at room temperature for 30 min with continuous mixing. The eluates were diluted in sample buffer containing 2% SDS for gel fractionation and subsequent Plg detection.

Lysine-Sepharose enrichment of plasma Plg

Freshly prepared citrated plasma samples [150 µl] were diluted in an equal volume of buffer A [0.15 M potassium phosphate, 0.8 mM sodium phosphate, 14 mM NaCl, and 0.3 mM KCl [pH 7.3]] and then combined with 100 µl [settled volume] of lysine-Sepharose 4B [Pharmacia]. The suspensions were incubated at room temperature for 30 min with continuous mixing. The Sepharose beads were collected by centrifugation at 3000 g for ~30 sec and resuspended in 1 ml of buffer A. This wash step was repeated eight times with the OD at 280 nm of the third wash being <0.02. Bound material was eluted from the lysine-Sepharose by a 30-min incubation with an equal volume of buffer A containing 0.4 M e-amino caproic acid. The eluates were diluted in sample buffer containing 2% SDS for gel fractionation and subsequent Plg detection.

Histology and fibrinogen immunostaining of tissue sections

Mice were sacrificed under ketamine anesthesia [see above], and tissues were placed immediately into zinc-formalin fixative [U.S. Biotex]. Tissues were fixed for 24 hr and then processed into paraffin, sectioned, and stained by standard techniques. Immunostaining was performed using the Vectastain Elite ABC kit [Vector Laboratories] and nickel-enhanced diaminobenzidine stain.

uPA esterase activity in urine

Urine was collected directly [void urine] or by bladder puncture in ketamine-anesthetized mice [see above]. The urine collected was either snap frozen in liquid nitrogen or placed directly into gel sample buffer containing 2% SDS. For immunoblotting, 5-µl urine samples were fractionated by SDS-polyacrylamide gel electrophoresis [12% acrylamide], and uPA detected as described above. uPA esterase activity assays were performed using the uPA-specific chromogenic substrate, Spectrozyme-UK [Promega], the 5-bromo-4-chloro-3-indolyl phosphate (BCIP)/nitro blue tetrazolium (NBT) [Sigma] staining system.

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Plasminogen deficiency causes severe thrombosis but is compatible with development and reproduction.

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