Thrombin-activatable fibrinolysis inhibitor (TAFI) deficiency is compatible with murine life

Mariko Nagashima,1 Zheng-Feng Yin,2 Lei Zhao,1 Kathy White,3 Yanhong Zhu,4 Nina Lasky,2 Meredith Halks-Miller,3 George J. Broze, Jr.,2 William P. Fay,4 and John Morser1

1Department of Cardiovascular Research, Berlex Biosciences, Richmond, California, USA
2Division of Hematology, Washington University School of Medicine, Barnes-Jewish Hospital, St. Louis, Missouri, USA
3Department of Pharmacology, Berlex Biosciences, Richmond, California, USA
4University of Michigan Medical Center, Ann Arbor, Michigan, USA

Address correspondence to: M. Nagashima, Department of Cardiovascular Research, Berlex Biosciences, 15049 San Pablo Avenue, Richmond, California 94804, USA. Phone: (510) 669-4136; Fax: (510) 669-4246; E-mail: Mariko_Nagashima@berlex.com.

Received for publication January 5, 2001, and accepted in revised form November 13, 2001.

To investigate the consequence of deficiency in thrombin-activatable fibrinolysis inhibitor (TAFI), we generated homozygous TAFI-deficient mice by targeted gene disruption. Intercrossing of heterozygous TAFI mice produced offspring in the expected Mendelian ratio, indicating that transmission of the mutant TAFI allele did not lead to embryonic lethality. TAFI-deficient mice developed normally, reached adulthood, and were fertile. No gross physical abnormalities were observed up to 24 months of age. Hematological analysis of TAFI-deficient mice did not show any major differences including plasma fibrinogen level, prothrombin time, and activated partial thromboplastin time. TAFI-deficient mice did not suffer from excess bleeding as determined by blood loss following tail transection, although their plasma failed to prolong clot lysis time in vitro. In vivo, TAFI deficiency did not influence occlusion time in either an arterial or a venous injury model. TAFI deficiency did not improve survival rate compared with the wild-type in thrombin-induced thrombosis, factor X coagulant protein–induced thrombosis, and endotoxin-induced disseminated intravascular coagulation. Furthermore, TAFI deficiency did not alter kaolin-induced writhing response, implying that TAFI does not play a major role in bradykinin catabolism. The current study demonstrates that TAFI deficiency does not change normal responses to acute challenges.


Introduction

The fibrinolytic system is essential in the maintenance of blood vessel patency and removal of extravascular fibrin deposition. It is also thought to play a critical role in the degradation of extracellular matrices in concert with other proteinases in a number of physiological and pathological conditions (1–5). Plasmin is the key protease in the fibrinolytic system that degrades fibrin. It is converted from a circulating zymogen, plasminogen, by either tissue-type plasminogen activator (t-PA) or urokinase-type plasminogen activator (u-PA). Inhibition of the fibrinolytic system may occur at the level of plasmin by α2-antiplasmin or at the level of its activators by plasminogen activator inhibitors, predominantly PAI-1. Recently, another protein, thrombin-activatable fibrinolysis inhibitor, has been shown to inhibit the fibrinolytic system.

Thrombin-activatable fibrinolysis inhibitor (TAFI; EC 3.4.17.20) is a 60-kDa plasma glycoprotein that possesses carboxypeptidase B activity upon activation. This glycoprotein was originally described as a novel basic carboxypeptidase generated in serum following coagulation, which was distinct from carboxypeptidase N (EC 3.4.17.3), the key plasma enzyme responsible for degradation of anaphylatoxins (6, 7). Because of its instability, Hendriks et al. named it carboxypeptidase U (U referring to unstable; ref. 6), while Campbell et al. named it arginine carboxypeptidase (carboxypeptidase R; ref. 7). Meanwhile, a novel protein with a high affinity for plasminogen was isolated, cloned, and shown to have high-sequence homology to that of pancreatic procarboxypeptidase B (8). Following incubation with a serine protease such as trypsin or thrombin, this protein had carboxypeptidase B activity and was termed plasma procarboxypeptidase B. Subsequent characterization revealed the identity between carboxypeptidase U and plasma procarboxypeptidase B (9). The same protein was purified and characterized by Bazjar et al., who demonstrated that this protein inhibited fibrinolysis following its activation by thrombin, hence the name “thrombin-activatable fibrinolysis inhibitor” (10, 11). The potential role of this enzyme in the regulation of fibrinolysis was also suggested based on other in vitro and in vivo studies (9, 12–16).

In vitro TAFI is activated via proteolytic cleavage of a 92 amino acid–long activation peptide by a high con-
The turnover of biologically active peptides possesses a physiological challenge, we generated homozygous TAFI-deficient mice. The newly exposed C-terminal lysine residues of fibrin following partial degradation by plasmin bind both plasminogen and t-PA with high affinity, thereby potentiating plasmin generation at the surface of fibrin clot (19–22). Activation of TAFI during clot lysis results in removal of these C-terminal lysines, and consequently, reduced plasmin production, leading to a several-fold prolongation of clot lysis time. Numerous in vivo studies in the rabbit have demonstrated that inhibition of TAFI activity with carboxypeptidase inhibitor from potato (CPI) enhanced t-PA–induced thrombolysis as well as endogenous fibrinolysis (23–26). Furthermore, fibrin deposition in either the lung or kidney of the rat in response to endotoxin was reduced by inhibition of TAFI activity (27, 28). Therefore, accumulating evidence indicates that TAFI plays a role in the regulation of fibrinolysis in vivo with a caveat that in these studies CPI was used to preferentially block TAFI activity. While it is inactive against carboxypeptidase N, it inhibits a number of other carboxypeptidases present in the body (for review, see ref. 29).

Mouse models deficient in each component of the fibrinolytic system have been generated to examine their role in development, thrombosis, thrombolysis, reproduction, wound repair, and cell migration (30–34). Unlike deficiency in coagulation factors such as tissue factor or factor V, which are embryonic lethal, deficiency in a component of the fibrinolytic system results in animals that develop normally with no obvious phenotypic abnormalities. However, plasminogen-deficient mice suffered spontaneous thrombosis, and wound healing was grossly impaired (32, 33). Furthermore, plasminogen activator–deficient mice had impaired lysis of microthrombi in the lung and increased fibrin deposition in response to endotoxin challenge (31), while mice deficient in either PAI-1 or α2-antiplasmin had increased thrombolysis and were more resistant to endotoxin-induced thrombosis (30, 34). To examine the consequence of TAFI deficiency on endogenous fibrinolysis, as well as other prothrombotic challenges, we generated homozygous TAFI-deficient mice by targeted gene disruption. Such an animal model can also be used to determine if TAFI is involved in the turnover of biologically active peptides possessing C-terminal basic residues. Here we report the generation and characterization of TAFI-deficient mice.

Methods

Animals. All mice were maintained in an animal facility accredited by the Association for Assessment and Accreditation of Laboratory Animal Care International. The protocols for in vivo studies were approved by Berlex Biosciences Animal Care and Use Committee. Mice were in a mixed background of C57BL/6 and 129/Sv and used at an age between 8 and 12 weeks old. For most of the studies, wild-type littermates were used as controls. However, for tail bleeding, thrombin-induced pulmonary thromboembolism, and LPS-induced disseminated intravascular coagulation studies in which large numbers of mice were required, offspring of TAFI-deficient mice used for breeding were compared with offspring of their wild-type littermates used for breeding.

Materials. Human thrombin (3,000 NIH U/mg, obtained by a direct comparison to an NIH thrombin reference standard; 1 NIH unit equals 1.15 IU) was purchased from American Diagnostica Inc. (Greenwich, Connecticut, USA). Recombinant tissue-type plasminogen activator (580,000 IU/mg) was purchased from Genentech Inc. (South San Francisco, California, USA). Human thrombomodulin was purified as described previously (35). Endotoxin (Escherichia coli LPS, serotype 0111:B4), HEPES, and 3-[3-cholamidopropyl]dimethylammonio]-1-propane-sulfonate (CHAPS) were obtained from Sigma Chemical Co. (St. Louis, Missouri, USA). Furylacycloyl-alanyl-arginine (FA-Ala-Arg) was from Bachem AG (Bubendorf, Switzerland). D-Phe-Pro-Arg chloromethylketone (PPACK) and D-Val-Phe-Lys chloromethylketone were obtained from Calbiochem-Novabiochem Corp. (San Diego, California, USA). Lovenox (enoxaparin sodium) was purchased from Aventis Pharmaceuticals Inc. (Bridgewater, New Jersey, USA).

Generation of TAFI gene-disrupted mice. The full-length murine TAFI cDNA was isolated from a mouse liver cDNA library (lambda ZAP; Stratagene, La Jolla, California, USA) using a partial human TAFI cDNA as probe. The murine TAFI cDNA was used to screen a P1 genomic library prepared from the genomic DNA isolated from the 129/Sv mouse (Genome Systems Inc., St. Louis, Missouri, USA). A 9.2-kb SacI fragment of TAFI DNA containing exons 6 to 9 and a 6.4-kb Spel DNA fragment containing exon 10 were subcloned into pBluescript II KS (Stratagene) producing plasmids A and B, respectively. For the construction of the targeting vector, a HSV-thymidine kinase (tk) cassette was placed at the KpnI site within the multiple cloning site of pBluescript II KS using blunt-end ligation. A 0.9-kb AclI-HindIII TAFI DNA fragment spanning a portion of intron 7 and half of exon 8 from plasmid A was subcloned into pET-21b (Stratagene) at its multiple cloning site. A Xhol-HindIII fragment containing the TAFI DNA was removed from the resulting pET-21b plasmid and was inserted between Xhol and HindIII sites directly after the tk cassette in the targeting vector. A Sall fragment containing a 1.5-kb PGK-neomycin phosphotransferase (neo) cassette was inserted into plasmid B directly upstream of the 6.4-kb TAFI DNA. Construction of the targeting vector was then completed by inserting a 7.9-kb Xhol-NotI DNA fragment from plas-
mid B, containing the neo cassette and exon 10 of TAFI gene, between SalI and NotI sites of the targeting vector (Figure 1a). The targeting vector was linearized with NotI and introduced by electroporation into 129/Sv-derived R1 embryonic stem cells (gift from A. Nagy, University of Toronto, Ontario, Canada), and stable transfectants were selected as described previously (36). Of 420 clones, two contained the expected allele as determined by Southern blot analysis. An embryonic stem cell clone was injected into C57BL/6 blastocysts as described (36), and the resulting male chimeras were bred to C57BL/6 females to generate heterozygous F1 offspring. Heterozygous TAFI mice were intercrossed, yielding TAFI+/+, TAFI+/−, and TAFI−/− F2 littersmates.

**Genotypic analysis.** Genomic DNA was isolated from either tail clips or ear biopsies and used for genotyping by either Southern blot analysis or PCR analysis. For Southern blotting, a 0.45-kb HindIII fragment of TAFI genomic DNA encompassing exon 6 was used as a hybridization probe. PCR analysis of the TAFI gene was performed using three primers derived from (a) the upstream portion of exon 8 (5′-AGAAAGGTGCGTCAAGTTCC3′-3′), (b) a portion of exon 8 deleted in the mutant allele (5′-GGTCTCTCGGTGCTTGTG-3′), and (c) the neo cassette (5′-TTCTGACTAGGGGAGGAGAG-3′). The PCR product derived from the wild-type allele is 190 bp, and that derived from the mutant allele is 390 bp.

**RT-PCR.** mRNA was extracted from mouse liver using a FastTrack 2.0 kit (Invitrogen Corp., Carlsbad, California, USA) according to the manufacturer’s instructions. First-strand cDNA was synthesized by oligo (dT) priming using the First-Strand cDNA Synthesis kit (Roche Molecular Biochemicals, Indianapolis, Indiana, USA). The first-strand cDNA product was used in the subsequent PCR amplification with primers, one annealing to the 5′ end of TAFI cDNA (5′-CAAGTCACGTGTGGGATGAGGC-3′) and the second one annealing to the 3′ end of TAFI cDNA (5′-ATTACTGTCTCGTGTGAGCAGGCC-3′). All the PCR products were analyzed by electrophoresis on 1% agarose gels.

**Determination of TAFI antigen in plasma.** The presence of TAFI in plasma is demonstrated using either Western blot analysis or an ELISA. Mouse platelet-poor plasma was obtained by centrifugation of citrated blood (9:1 vol/vol with 3.8% sodium citrate) at 1,500 × g for 15 minutes. For Western blot analysis, plasma was diluted with 25 µl of water and allowed to clot at room temperature with addition of thrombin (4.5 NIH U/ml, final), thrombomodulin (50 nM, final), and calcium chloride (17 mM, final). Ten minutes later, the clot was squeezed and discarded, and activation of TAFI was terminated by addition of PPACK (1 µM, final). Total carboxypeptidase B activity (TAFIa and carboxypeptidase N activity) was measured by addition of a 40-µl aliquot of the remaining fluid to 960 µl of 0.02 M Tris-HCl, pH 7.8, 0.075 M NaCl, and 0.1% (wt/vol) CHAPS containing FA-Ala-Arg (0.4 mM final) as a substrate. The rate of hydrolysis of FA-Ala-Arg was determined by measuring changes in absorbance at 336 nm every 20 seconds for 10 minutes. Carboxypeptidase N (CPN) activity was measured similarly in the diluted plasma without addition of thrombin and thrombomodulin. The values for CPN thus obtained were same as those obtained by measuring the total carboxypeptidase B activity as described above, except in the presence of CPI.

**Figure 1**

**TAFI gene-disruption in mice.** (a) TAFI replacement construct and partial restriction map of the endogenous locus. Exon 8 and 9 encodes some of the zinc and substrate binding sites of TAFI. (b) Southern blot analysis of TAFI gene disruption. The probe indicated in a was used to detect 9.2-kb DNA fragment in the wild-type allele and 7.0-kb DNA fragment in the mutant allele following Sac1 digestion of genomic DNA. (c) Western blot analysis of plasma TAFI from the wild-type, heterozygous, and homozygous TAFI mice. Ab raised against TAFI peptide (CGIHAREWISPFA) hybridized to a band with a molecular weight similar to that of murine albumin only in the plasma from the wild-type and heterozygous mice. (d) RT-PCR analysis of liver mRNA isolated from TAFI+/+, TAFI+/−, and TAFI−/− mice. RT-PCR products obtained with primers that span the entire coding region of TAFI cDNA in the presence of reverse transcriptase (RT) are shown on the left-hand side and those in the absence of reverse transcriptase are on the right-hand side. Neo, neomycin phosphotransferase gene; tk, thymidine kinase gene; probe, TAFI-specific probe used in Southern blot analysis (which encodes exon 6).
TAFI antigen (nM, equivalent) 0.62 ± 0.23 (n = 5) 0.34 ± 0.12 (n = 6) 0.07 ± 0.03 (n = 7)

TAFI activity (mOD/min) 0.88 ± 0.33 (n = 5) 0.52 ± 0.29 (n = 6) 0.1 ± 0.15 (n = 7)

CPN activity (mOD/min) 3.72 ± 1.18 (n = 5) 3.63 ± 0.67 (n = 6) 4.7 ± 1.06 (n = 7)

TAFI antigen level in plasma was measured using an ELISA assay kit with purified human TAFI as a standard. TAFI activity in plasma was determined using FA-Ala-Arg as substrate following its activation with thrombin (4.5 NIH U/ml), thrombomodulin (50 nM), and CaCl2 (17 mM) as described in Methods. CPN activity was determined similarly without addition of thrombin, thrombomodulin, and CaCl2. The number of mice used in each group is indicated in parentheses. mOD, milli-optical density.

**Table 1**

<table>
<thead>
<tr>
<th>TAFI+/+</th>
<th>TAFI+/–</th>
<th>TAFI–/–</th>
</tr>
</thead>
<tbody>
<tr>
<td>TAFI antigen (nM, equivalent)</td>
<td>0.62 ± 0.23</td>
<td>0.34 ± 0.12</td>
</tr>
<tr>
<td>TAFI activity (mOD/min)</td>
<td>0.88 ± 0.33</td>
<td>0.52 ± 0.29</td>
</tr>
<tr>
<td>CPN activity (mOD/min)</td>
<td>3.72 ± 1.18</td>
<td>3.63 ± 0.67</td>
</tr>
</tbody>
</table>

In *vitro* plasma clot lysis. The clot lysis profile of mouse plasma was determined using a 96-well microtiter plate clot lysis assay. Fifteen microliters of citrated plasma was mixed with 3 µl of 2 µM thrombomodulin and 60 µl of 0.04 M HEPES, pH 7.0, 0.15 M NaCl, and 0.01% Tween 80 (vol/vol). The mixture was immediately added to another well containing 4 µl of 75 NIH U/ml thrombin, 2 µl of 1 M CaCl2, and 4 µl of 0.5 µg/ml t-PA in separate aliquots. The total volume of the mixture was made up to 120 µl with water. After mixing, the change in absorbance at 405 nm was monitored at 37°C every minute for 60 minutes using a SpectraMAX 250 Microplate Spectrophotometer (Molecular Device Corp., Sunnyvale, California, USA). Lysis time was defined as the time at which the absorbance is one-half of the difference between the plateau reached after clotting and the base line value achieved at complete lysis.

**Hematological analysis, coagulation assays, and tail bleeding.** Blood samples were collected from the inferior vena cava of animals into one-tenth volume of 3.8% sodium citrate. Blood cell counts were analyzed using a Bayer 9120+CP Cell Counter (ABX Diagnostics Inc., Irvine, California, USA). Fibrinogen level was determined according to the method described by Macart (38), and the results are expressed as a percentage of the initial value. The activated partial thromboplastin time (aPTT) and prothrombin time (PT) were performed with platelet-poor plasma using an Electra 900C coagulometer (Beckman Coulter, Brea, California, USA). For blood chemistry analysis, serum was obtained from clotted whole blood and analyzed by IDEXX Veterinary Services Inc. (West Sacramento, California, USA). For measurement of tail bleeding, mice were anesthetized with an intraperitoneal injection of pentobarbital (62.5 mg/kg body weight), and a 1-mm segment was amputated from the tip using a device that cuts with a razor blade. The tail was immediately immersed in 25 ml of saline prewarmed to 37°C, with constant mixing. After 30 minutes, 5 ml of 5% EDTA in PBS was added, and the number of red blood cells released from the tail wound into saline was determined with a Baker 9120+CP Cell Counter. For studying the effect of heparin on tail bleeding, mice were preinjected in the tail vein with Lovenox, a low-molecular-weight heparin (3 mg/kg body weight) 2 minutes before tail amputation.

**Mouse thrombosis models.** A murine model of vascular injury induced by a photochemical reaction either in the artery or in the vein was performed as described previously (39). Briefly, either the left common carotid artery or the left jugular vein was isolated, and a vascular flow probe (Transonic Systems, Ithaca, New York, USA) was applied to monitor blood flow continuously. Rose bengal (Fisher Scientific Co., Pittsburgh, Pennsylvania, USA) at a concentration of 10 mg/ml was mixed with 3

In the thrombin-induced acute thromboembolism paradigm, human thrombin was injected over 2 to 3 seconds into the tail vein of conscious mice at a dose of 1,600, 2,400, or 3,200 NIH U/kg body weight. The survival rate at 10 minutes after intravenous administration of thrombin was measured as an index of the severity of thromboembolism. In the endotoxin-induced septic shock model, mice were injected intraperitoneally with either 30 mg/kg or 40 mg/kg body weight of LPS dissolved in saline. The mice were observed for signs of endotoxemia daily, such as lethargy, scruffy fur, and some exudate from eyes, and survival time was recorded. In separate experiments, an effect of LPS on fibrin deposition in the kidney was investigated. Three hours after injection of LPS, mice were anesthetized and citrated blood was collected by cardiac puncture. Kidneys were perfused immediately with PBS containing 3.8% sodium citrate, 100 µg/ml Lovenox, and 1 µM t-Val-Phe-Lys chloromethylketone, followed by 4% paraformaldehyde (in PBS, sucrose 1% wt/vol). The kidneys were harvested and fixed overnight with fresh 4% paraformaldehyde. Kidneys were cut in half and embedded in paraffin. Tissue sections (5 µM-thickness) were cut, hydrated, and incubated sequentially first with a solution of 3% hydrogen peroxide in PBS for 30 minutes and then a 10% normal goat serum in PBS at room temperature. Sections were immunostained for fibrin/fibrinogen using a rabbit antifibrinogen antiserum (kindly provided by J.L. Degen; University of Cincinnati, Ohio, USA; diluted 1:1000, vol/vol) followed by biotinylated goat anti-rabbit IgG, and detected with streptavidin peroxidase conjugate and diaminobenzidine (Zymed Laboratories, Inc., South San Francisco, California, USA). Following immunostaining, the extent of fibrin deposition was analyzed by a pathologist without prior knowledge of the genotype of the mice. In the factor X coagulant protein–induced (XCP-induced) subacute thrombosis model, mice were injected intraperitoneally with...
150 μg/kg body weight XCP from Russell viper venom. The survival rate, platelet counts, and fibrinogen level were determined at 2 and 16 hours.

**Kaolin writhing test.** Mice were injected intraperitoneally with 0.5 ml kaolin suspended in saline (5 mg/ml) to introduce the writhing stretching response (40, 41). Subsequent writhing responses were counted over the next 15 minutes.

**Statistical analysis.** All data are presented as mean plus or minus SEM. The significance of differences in tail bleeding between groups was determined by ANOVA. For occlusion time, the nonparametric Kruskal-Wallis one-way ANOVA followed by the Mann-Whitney U test was used. \( P \) values less than 0.05 were considered significant.

**Results**

**Generation of TAFI-deficient mice.** To investigate in vivo consequences of TAFI deficiency, the TAFI gene was disrupted in mice as described in Methods. Successful gene targeting resulted in the replacement of a portion of exon 8 and 9 with a neo cassette (Figure 1a), thereby removing the DNA segment encoding residues critical for substrate and zinc binding essential for catalytic activity. The expected structure of the targeted TAFI locus was confirmed by Southern blot analysis. Thus, after SacI digestion of genomic DNA, the probe that encompasses exon 6 recognized a band of approximately 9.2 kb in the wild-type allele and a band of approximately 7.0 kb in the disrupted allele (Figure 1b). Germline transmission of the TAFI-null allele was achieved in a number of chimeric mice derived from two independent embryonic stem cell clones. Mice generated from both embryonic stem cell clones were characterized independently and were shown to be indistinguishable phenotypically. Consequently, most of the work described in the current study was performed with one line.

The targeted disruption of the **TAFI** gene resulted in the truncation of the TAFI transcript as shown by RT-PCR of mRNA isolated from the liver of homozygous TAFI-null mice (Figure 1d). The liver was the main tissue in which TAFI mRNA was detected by Northern blot analysis in the mouse (42, 43). Using primers that correspond to the 5’ and 3’ end of TAFI cDNA, the RT-PCR produced a fragment of approximately 1,000 bp in TAFI-deficient mice compared with a fragment of approximately 1,350 bp in the wild-type littermates. The RT-PCR amplified both fragments when mRNA from heterozygous littermates was tested. In the absence of reverse transcriptase, no products were obtained.

To determine if the truncated TAFI transcript in the homozygous null mice leads to a synthesis of a truncated TAFI protein, Western blot analysis of plasma samples from different genotypes was performed. Polyclonal Ab’s raised against a TAFI peptide corresponding to a region in the TAFI molecule that is encoded by exon 5 was used (Figure 1c). Therefore, if a truncated form of TAFI protein existed in circulation, it should be detected using this Ab. In the plasma from the wild-type and heterozygous TAFI mice, polyclonal Ab’s bound to a band with a molecular weight slightly higher than that of albumin. This is consistent with the predicted molecular weight for mouse TAFI. On the other hand, no band was detected in the plasma from TAFI-deficient mice. The lack of TAFI protein in the plasma of TAFI-deficient mice was also demonstrated using a commercially available ELISA kit for detection of human TAFI, with purified human TAFI as standard. The plasma level of TAFI in heterozygous mice was found to be about half of that detected in the wild-type mice (Table 1). Only the background value in the ELISA was found with plasma samples from homozygous mice, which was similar to the value obtained with human plasma samples depleted of TAFI.

**Viability and fertility of TAFI-deficient mice.** DNA analysis of 259 progeny derived from TAFI+/− intercrosses showed that TAFI mice were born in the expected Mendelian ratio of one TAFI+/−/two TAFI+/+/one TAFI−/− (Table 2). This indicated that TAFI deficiency did not lead to embryonic lethality. TAFI-deficient mice developed normally and reached adulthood. No physical abnormalities were observed in these mice up to 24 months of age. Matings of homozygous males with homozygous females produced viable offspring of normal litter size at normal frequency indicating that TAFI-deficient mice were fertile and pregnancies were carried to full term. Hematological analysis of TAFI−/− mice did not show any major differences in blood cell counts, plasma fibrinogen level, PT, and aPTT, compared with those of TAFI+/− mice (Table 3). There was no abnormality in liver and kidney functions as shown by blood chemistry analysis (Table 3).

**Table 2**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>TAFI+/+</th>
<th>TAFI+/−</th>
<th>TAFI−/−</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of pups</td>
<td>64</td>
<td>127</td>
<td>68</td>
</tr>
<tr>
<td>Percentage of total</td>
<td>25%</td>
<td>49%</td>
<td>26%</td>
</tr>
</tbody>
</table>

*Genotype of offspring from intercrossing of heterozygous TAFI mice was determined by PCR.

**Table 3**

<table>
<thead>
<tr>
<th>TAFI+/+</th>
<th>TAFI+/−</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin (g/%)</td>
<td>14.7 ± 1.4 (n = 15)</td>
</tr>
<tr>
<td>Platelet count (10^9/l)</td>
<td>1013 ± 176 (n = 15)</td>
</tr>
<tr>
<td>Fibrinogen (%)</td>
<td>100 ± 35 (n = 15)</td>
</tr>
<tr>
<td>Alkaline phosphatase (IU/l)</td>
<td>85.4 ± 22 (n = 5)</td>
</tr>
<tr>
<td>Blood urea nitrogen (mg/dl)</td>
<td>25.8 ± 4.7 (n = 5)</td>
</tr>
<tr>
<td>Creatine (mg/dl)</td>
<td>0.44 ± 0.06 (n = 5)</td>
</tr>
<tr>
<td>PT (s)</td>
<td>10.9 ± 0.41 (n = 6)</td>
</tr>
<tr>
<td>aPTT (s)</td>
<td>19.3 ± 0.97 (n = 6)</td>
</tr>
</tbody>
</table>

For blood chemistry analysis, serum was obtained from clotted whole blood and analyzed by IDEXX Veterinary Services Inc. Fibrinogen level was measured according to the method described by Macart (38). The aPTT and PT were performed with platelet-poor plasma using a MLA Electra 3000C coagulometer. The number of mice used in each group is indicated in parentheses.
TAFI-deficient mice did not suffer from any apparent untoward effects of tail transection when samples were taken for genotyping. To evaluate the effect of TAFI deficiency on hemostasis, blood loss was determined during the 30 minutes following tail transection of a 1-mm segment. The mice used in the study had intact tails prior to the amputation because routine genotyping was performed with ear biopsies instead of tail clips. There was no statistically significant difference in the amount of blood loss between TAFI-deficient mice and the control mice (Figure 2a). It has been shown that the 129/Sv mice have significantly longer bleeding times than the C57Bl/6 mice do, indicating that genetic backgrounds influence hemostasis (44). In this study, we compared offspring of TAFI-deficient mice with offspring of their wild-type littermates, rather than comparing among the same littermates. However, the present results were similar to those obtained from our previous study in which tail-bleeding time in TAFI-deficient mice was compared with their littermate controls and no differences were found (44). The effect of TAFI deficiency on bleeding was also studied in the presence of an antithrombotic agent that promoted exaggerated bleeding. In the presence of low-molecular-weight heparin, there were wide individual differences in the extent of bleeding. However, the blood loss in TAFI-deficient mice was not significantly different from that in the wild-type mice, implying that TAFI deficiency did not compromise hemostasis severely (Figure 2b).

**Effect of TAFI deficiency on plasma carboxypeptidase activity and in vitro clot lysis.** To demonstrate a role of activated TAFI in fibrinolysis, the effect of TAFI deficiency was tested in a plasma clot lysis assay. Coagulation and fibrinolysis were initiated in mouse platelet-poor plasma with thrombin and t-PA, respectively. In the presence of 50 nM thrombomodulin, clot lysis time was prolonged at least twofold in plasma from wild-type mice (Figure 3a). This thrombomodulin-dependent prolongation was likely due to the activation of TAFI by a thrombin-thrombomodulin complex since an inhibitor of CPI reversed it (data not shown). A similar extent of prolongation was observed in plasma from heterozygous mice, indicating that while their plasma TAFI antigen level is about half of that in wild-type, enough TAFI is activated to perturb fibrinolysis. There was no prolongation of clot lysis time in plasma from TAFI-deficient mice upon the addition of thrombomodulin (Figure 3b). On the other hand, an addition of purified human TAFI to the same plasma resulted in a similar prolongation of clot lysis time as seen with plasma from the wild-type and heterozygous mice (data not shown). Therefore, activated TAFI is responsible for the retardation of fibrinolysis in the wild-type and heterozygous mice. These results are in agreement with TAFIa activity measured directly in these plasma samples with a small molecule substrate, FA-Ala-Arg, following in vitro activation with a thrombin/thrombomodulin complex (Table 1). Furthermore, it was evident that the activity of the second basic carboxypeptidase in plasma, CPN, remained relatively constant in the three groups of mice and that it did not influence fibrinolysis.

---

**Figure 2**
Tail bleeding. Wild-type mice and TAFI-deficient mice were anesthetized with an intraperitoneal injection of pentobarbital and a 1-mm segment was amputated from the tip of the tail, as described in Methods. The tail was immediately immersed in saline at 37 °C. The amount of red blood cells released from the tail wound into saline was determined with a Baker 9120+CP Cell Counter (a). For tail bleeding in the presence of low-molecular-weight heparin, mice were injected with Lovenox (3 mg/kg body weight) in the tail vein 2 minutes before surgery and in the presence of low-molecular-weight heparin, there were untoward effects of tail transection when samples were taken for genotyping. RBC, red blood cell; WT, wild-type; KO, knockout.

**Figure 3**
Plasma clot lysis profile. (a) represents wild-type mice and (b) represents TAFI-deficient mice. Clot formation and lysis were initiated in the recalcified (Ca²⁺, 17 mM) mouse plasma with human thrombin (2.5 NIH U/ml) and t-PA (0.017 µg/ml, respectively). TM, human recombinant soluble thrombomodulin, was added at a final concentration of 50 nM to enhance activation of endogenous TAFI. Absorbance at 405 nm was plotted against incubation time. Typical clot lysis profile obtained in the absence of TM and t-PA (solid line with open circle), in the absence of TM but presence of t-PA (dotted line with filled circle), in the presence of TM but absence of t-PA (dashed line with filled square), and in the presence of both TM and t-PA (dashed line with open square) is shown.
Table 4
Survival rate in thrombin-induced acute pulmonary thromboembolism

<table>
<thead>
<tr>
<th>Thrombin dose</th>
<th>Survival rate in TAFI+/+ (%)</th>
<th>Survival rate in TAFI−/− (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1,600 NIH U/kg</td>
<td>87 (n = 8)</td>
<td>62 (n = 8)</td>
</tr>
<tr>
<td>2,400 NIH U/kg</td>
<td>78 (n = 9)</td>
<td>64 (n = 11)</td>
</tr>
<tr>
<td>3,200 NIH U/kg</td>
<td>60 (n = 15)</td>
<td>50 (n = 12)</td>
</tr>
</tbody>
</table>

Human thrombin was injected intravenously over 2 to 3 seconds into the tail vein of conscious mice. The survival rate at 10 minutes after thrombin administration was taken as an index of the severity of pulmonary thromboembolism. The number of mice used in each group is indicated in parentheses.

Discussion
A correlation between fibrinolytic potential and the plasma level of TAFI antigen has been documented in the general population (46). In normal healthy subjects, plasma TAFI antigen level varies between 45% and 150% of the mean value (46–49). There are some reports that age influences the TAFI antigen level (47–49). The presence of polymorphisms in the TAFI gene has been observed (50–53), and it is now known that some of the polymorphisms are responsible for variations in the plasma TAFI antigen level (52, 53). The gene encoding human TAFI has been mapped to chromosome 13q14.11 (54, 55), but no known hemostatic disease is currently associated with this region; there has been no report of an individual lacking in plasma TAFI. One model, the significant decreases in the plasma fibrinogen level and platelet counts observed at 2 hours and 16 hours after the stimulus were similar in both genotypes (Table 5). Furthermore, the survival rate did not differ between the two groups.

Kaolin-induced writhing test. Since it has been shown that TAFI cleaves C-terminal arginine residue from bradykinin in vitro (12, 17, 18), the effect of TAFI deficiency on the writhing reaction induced by kaolin was studied in normal animals. The number of mice writhing during 15 minutes after the intraperitoneal injection of kaolin was 1.29 ± 1.78 (n = 14) in the wild-type compared with 1.21 ± 1.85 (n = 14) in TAFI-deficient mice. Kaolin is a known activator of factor XII and induces formation of bradykinin, which in turn induces writhing response. The result with TAFI-deficient mice demonstrated that TAFI does not play a major role in the degradation of bradykinin under normal conditions in this model.
study indicated an increased TAFI level was a mild risk factor for venous thrombosis (49). On the other hand, surprisingly, there has been a recent report that indicated lower levels of plasma TAFI as a risk factor for coronary heart diseases (56). Thus, we generated and characterized TAFI-deficient mice to study its pathophysiological role in various thrombotic diseases in the light of these current conflicting reports.

Mice deficient in TAFI were born without any abnormalities and developed normally to reach adulthood. They were fertile, producing viable offspring, which showed that TAFI deficiency was compatible with embryogenesis and development. Hematological and hemostatic analysis revealed no major differences between TAFI-deficient and wild-type mice, including tail bleeding. Previously, it was shown that in vitro clots formed from patient plasma deficient in coagulation factor VIII, IX, X, or XI lysed prematurely when induced by exogenous t-PA (36). When TAFI was activated upon addition of either the missing coagulation factor or thrombomodulin to such plasma, normal lysis was restored. Such data implicated a physiological role of TAFI in stabilization of newly formed fibrin clots. However, the results of tail bleeding in this study demonstrated that loss of TAFI did not lead to grossly impaired hemostasis.

The effect of TAFI deficiency was also studied in arterial as well as venous thrombosis models induced by a photochemical injury. In these models, initial formation of an occlusive thrombus depends on the balance between coagulation and fibrinolysis. TAFI deficiency did not have a significant impact on the rate of thrombus formation in either vessel. To test the possibility that enhanced endogenous fibrinolysis due to the absence of activated TAFI may facilitate spontaneous lysis of the stabilized clot, animals were allowed to recover, and the size of the residual thrombus was measured 24 hours later. We did not observe any statistically significant difference in the cross-sectional area of the thrombus from TAFI-deficient mice compared with that of the wild-type littermates (data not shown). Such results indicate that TAFI does not greatly influence the rate of spontaneous reperfusion. It is possible that PAI-1 release from the vessel wall in response to a photochemical injury may well mask any effect of TAFI. In the same arterial model, PAI-1 has been shown to affect both occlusion and spontaneous reperfusion rate (57, 58).

TAFI-deficient mice were challenged with other stimuli that are known to induce systemic intravascular coagulation. In a pulmonary thromboembolism model reported previously, intravenous injection of human thrombin into mice caused quick death (59–61). Fibrin deposition was detected in both large and small vessels in the lung (59, 61). In one of the papers, the authors demonstrated that thrombin-induced activation of clotting factors, and subsequent additional formation of thrombin, and hence fibrin formation, contributed significantly to organ failures (61). Furthermore, it was proposed that thrombin-induced activation of factor XIII and/or TAFI, which render newly formed fibrin clots more resistant to plasmin degradation, as additional pathological mechanisms. In our study, survival rate determined at 10 minutes after thrombin injection was similar among TAFI-deficient mice compared with the wild-type. To investigate any subtle effect of TAFI deficiency in response to thrombin injection, tissue fibrin deposition was measured using 125I-labeled human fibrinogen in separate experiments. There was no significant accumulation of radioactivity in the lung and kidney of the control mice at any time point up to 6 hours after thrombin injection (our unpublished observations). Intravenous injection of thrombin does not lead to substantial accumulation of fibrin in the kidney or lung in mice.

Following endotoxin or XCP injection, there were no differences in the survival rate between the wild-type mice and TAFI-deficient mice. Yamamoto and Loskutoff showed that LPS-induced transient fibrin deposition in tissue-specific vasculature was likely due to activation of the coagulation cascade through induction of tissue factor expression, and concomitant suppression of the fibrinolytic system through increased PAI-1 and decreased urokinase-type tissue plasminogen activator (45). In numerous studies, agents that directly inhibit tissue factor or thrombin as well as agents that inhibit amplification of coagulation cascade such as activated protein C have been effective in reducing endotoxin-induced lethality (for review, see refs. 62, 63). On the other hand, the effectiveness of an enhanced endogenous fibrinolytic system against a lethal dose of LPS has not been reported. However, reduced renal fibrin deposition in response to a nonlethal dose of endotoxin has been demonstrated in α2-antiplasmin–deficient mice (34, 64), but not in PAI-1–deficient mice (64). In the current study, we did not detect any difference in the extent of fibrin deposition in the kidney of wild-type and their TAFI-deficient littermates in response to LPS.

### Table 5
Survival rate, fibrinogen level, and platelet counts following XCP challenge

<table>
<thead>
<tr>
<th></th>
<th>TAFI+/+ (n = 15)</th>
<th>TAFI–/– (n = 15)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Survival rate (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>at 2 hours</td>
<td>93</td>
<td>87</td>
</tr>
<tr>
<td>at 16 hours</td>
<td>53</td>
<td>60</td>
</tr>
<tr>
<td>Plasma fibrinogen level (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>at 0 hours</td>
<td>100 ± 35</td>
<td>96 ± 26</td>
</tr>
<tr>
<td>at 2 hours</td>
<td>52 ± 23</td>
<td>56 ± 16</td>
</tr>
<tr>
<td>at 16 hours</td>
<td>34 ± 34</td>
<td>106 ± 56</td>
</tr>
<tr>
<td>Platelet counts (10^9/l)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>at 0 hours</td>
<td>1013 ± 176</td>
<td>1053 ± 288</td>
</tr>
<tr>
<td>at 2 hours</td>
<td>328 ± 252</td>
<td>416 ± 239</td>
</tr>
<tr>
<td>at 16 hours</td>
<td>319 ± 285</td>
<td>353 ± 142</td>
</tr>
</tbody>
</table>

Mice were injected intraperitoneally with 150 µg/kg body weight factor X coagulant protein from Russell viper venom. The survival rate, fibrinogen level, and platelet counts were determined 2 and 16 hours later. The number of mice used in each group is indicated in parentheses.
A potential involvement of TAFI in degradation of bradykinin was tested in TAFI-deficient mice using the kaolin-induced writhing model. Previous work has shown that kaolin induced writhing responses by way of bradykinin formation and that inhibition of the degradation of bradykinin by pretreatment of mice with a nonselective inhibitor of basic carboxypeptidases poten-
tiated the writhing response in this model (40, 41). In the absence of circulating TAFI antigen, however, there was no poten-
tiation of the writhing response, indicating that TAFI does not contribute significantly to the degradation of bradykinin under normal conditions.

In summary, we have generated and characterized TAFI-deficient mice. The current study demonstrated that TAFI deficiency is compatible with murine life. TAFI-deficient mice did not exhibit any abnormal phe-
notypes including hemostasis. Furthermore, the dis-
ruption of TAFI gene did not appear to participate in the acute challenges tested here. While it is evident from previous studies using TAFI inhibitors that TAFI plays a regulatory role in t-PA–induced thrombolyis (23–26), its effect on endogenous fibrinolysis may be more subtle. Our study also indicated that TAFI does not play a significant role in bradykinin turnover under normal conditions. Any potential role(s) of TAFI in the degradation of other biologically active peptides, es-
pecially anaphylatoxins, can now be tested unequivocally in TAFI-deficient mice instead of using nonselective inhibitors of basic carboxypeptidases. Currently, studies are underway to address these issues. During the preparation of our manuscript, Wagenaar et al. inde-
pendently reported generation of TAFI-deficient mice and observed a lack of abnormal phenotypes in their TAFI-deficient mice, consistent with our result (65).

Acknowledgments
We thank Joseph Post for determination of APTT and PT, Ron Vergona for helpful discussion, James Wang for statistical analyses, Bill Lillis and staff members of Animal Facilities at Berlex for animal care. We would also like to thank Bill Dole, Raju Mohan, Brad Buck-
man, and Katalin Kauser for their support.

References
1. Vasalli, J.-D., Sappino, A.-P., and Belin, D. 1991. The plasminogen acti-
stasis. In The molecular basis of blood diseases. 2nd edition. G. Stamatoy-
4. Verstraete, M. 1995. The fibrinolytic system. From petri dishes to genet-
6. Thorsen, S. 1992. The mechanism of plasminogen activation and vari-
ability of the fibrin effector during tissue-type plasminogen activator-
mediated stimulation of plasminogen activation by reducing the affin-
able fibrinolysis inhibitor potentiates tissue-type plasminogen activator-
lolysis in tissue factor-induced rat DIC model. Fibrinolysis and Proteoly-
sis 14:70 (Abstr.).
boxypeptidases. In Zinc metallopeptidases in health and disease. N.M. Hoop-
17. Bugge, T.H., Flick, M.J., Duhaucher, C., and Degen, J.L. 1995. Plas-
mogen deficiency causes severe thrombosis but compatible with devel-
19. Lijnen, H.R., Okada, K., Matsuo, O., Collen, D., and Dewerchin, M. 1999. 62-antiplasmin gene deficiency in mice is associated with enhanced fib-

The Journal of Clinical Investigation | January 2002 | Volume 109 | Number 1