Causal relationship between hyperfibrinogenemia, thrombosis, and resistance to thrombolysis in mice

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Epidemiologic studies have correlated elevated plasma fibrinogen (hyperfibrinogenemia) with risk of cardiovascular disease and arterial and venous thrombosis. However, it is unknown whether hyperfibrinogenemia is merely a biomarker of the proinflammatory disease state or is a causative mechanism in the etiology. We raised plasma fibrinogen levels in subjects with the highest plasma fibrinogen concentrations. The Framingham and Fragmin During Instability in Coronary Artery Disease studies positively correlated fibrinogen with risk of cardiovascular disease and incidence of death and/or myocardial infarction, respectively. The Leiden Thrombophilia Study showed that persons with elevated fibrinogen levels (4.0-4.9 vs < 3.0 mg/mL, 130%-160% of normal) have an adjusted odds ratio for venous thrombosis of 1.6, whereas persons with ≥ 5 mg/mL fibrinogen (≥ 170% of normal) have a 4-fold higher thrombotic risk, even after adjusting for C-reactive protein levels. These epidemiologic studies suggest that elevated fibrinogen is an independent risk factor for both arterial and venous thrombosis and therefore a potential diagnostic and therapeutic target for predicting and reducing thrombosis.

Importantly, however, epidemiologic studies have not and cannot show a causal relationship between fibrinogen and disease etiology. Fibrinogen levels increase with age, inflammatory processes, hematocrit, hypertension, glucose intolerance, cigarette smoking, and adiposity, and high fibrinogen levels increase plasma viscosity, a demonstrated risk factor for coronary heart disease. These potential confounders have not permitted distinction between fibrinogen’s role as a biomarker of inflammation or coexistent comorbidity and a direct, causative role in the etiology of cardiovascular disease.

Prior studies using animal models to clarify the role of hyperfibrinogenemia in thrombosis have been equivocal and controversial. Transgenic mice overexpressing murine fibrinogen (~ 45% higher than wild-type) demonstrate elevated D-dimer and spontaneous fibrin deposition in the spleen, suggesting that hyperfibrinogenemia is mildly prothrombotic. However, these mice demonstrate only marginal shortening of the time to 75% occlusion after 20% ferric chloride (FeCl3) application to the carotid artery, indicating that hyperfibrinogenemia is not important in arterial thrombosis. In contrast, rabbits treated with turpentine to elevate fibrinogen before stasis- or mechanical injury-induced venous thrombosis demonstrate a positive correlation between thrombus size, weight, and fibrin content. However, because turpentine also increases factor VIII, another thrombosis risk factor, the specific prothrombotic contribution of elevated fibrinogen is difficult to discern. One recent study in which the human γ'-chain of fibrinogen was expressed in transgenic mice suggested that fibrinogen’s thrombin-binding properties are antithrombotic, further questioning a pathologic mechanism relating hyperfibrinogenemia to disease.

The aim of the current study was to determine whether elevated fibrinogen directly contributes to thrombosis and identify the operative mechanism(s). We used in vivo models to assess fibrinogen’s effects on thrombus formation and stability, and cell and tissue factor (TF)-based ex vivo and in vitro methods to identify biochemical and biomechanical mechanisms by which fibrinogen modulates fibrin formation, structure, and function. Our data indicate that hyperfibrinogenemia directly promotes thrombosis and thrombolysis resistance and does so via enhanced fibrin formation and stability. These findings strongly suggest a causative role for hyperfibrinogenemia in acute thrombosis and have significant implications for thrombolytic therapy. Plasma fibrinogen levels may be used to identify patients at risk for thrombosis and inform thrombolytic administration for treating acute thrombosis/thromboembolism.

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Introduction

Elevated plasma fibrinogen is associated with risk of cardiovascular disease and arterial and venous thrombosis. Several studies have detected dose effects, with increased risk of death or thrombosis in subjects with the highest plasma fibrinogen concentrations. The Framingham and Fragmin During Instability in Coronary Artery Disease studies positively correlated fibrinogen levels with risk of cardiovascular disease and incidence of death and/or myocardial infarction, respectively. The Leiden Thrombophilia Study showed that persons with elevated fibrinogen levels (4.0-4.9 vs < 3.0 mg/mL, 130%-160% of normal) have an adjusted odds ratio for venous thrombosis of 1.6, whereas persons with ≥ 5 mg/mL fibrinogen (≥ 170% of normal) have a 4-fold higher thrombotic risk, even after adjusting for C-reactive protein levels. These epidemiologic studies suggest that elevated fibrinogen is an independent risk factor for both arterial and venous thrombosis and therefore a potential diagnostic and therapeutic target for predicting and reducing thrombosis.

Importantly, however, epidemiologic studies have not and cannot show a causal relationship between fibrinogen and disease etiology. Fibrinogen levels increase with age, inflammatory processes, hematocrit, hypertension, glucose intolerance, cigarette smoking, and adiposity, and high fibrinogen levels increase plasma viscosity, a demonstrated risk factor for coronary heart disease. These potential confounders have not permitted distinction between fibrinogen’s role as a biomarker of inflammation or coexistent comorbidity and a direct, causative role in the etiology of cardiovascular disease.


An Inside Blood analysis of this article appears at the front of this issue.

The online version of this article contains a data supplement.

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indicate that hyperfibrinogenemia directly and independently shortened the time to occlusion (TTO) and increased thrombus resistance to thrombolysis. These effects were mediated through enhanced fibrin formation and increased fibrin network density and mechanical and fibrinolytic stability. Together, these findings strongly suggest a causative role for hyperfibrinogenemia in the pathology of thrombosis. Information on plasma fibrinogen levels may be used to identify patients at risk for thrombosis and inform thrombolytic administration for treating arterial and venous thrombosis.

METHODS

Proteins and materials

Dulbecco modified Eagle medium with high glucose/2 mM l-glutamine, 0.05% trypsin and ethylenediaminetetraacetic acid, and phosphate-buffered saline (10 mM phosphate, pH 7.1, 150 mM NaCl, phosphate-buffered saline) were from Invitrogen. Thrombin fluorogenic substrate (Z-Gly-Gly-Arg-AMC) and calibrator (o2-macroglobulin/thrombin) were from Diagnostica Stago. Factor Xa chromogenic substrate (Pefachrome FXa) was from Pentapharm. Mouse anti-human TF antibody (HTF-1) was the kind gift of Dr Ronald Bach (University of Minnesota). Tissue-type plasminogen activator (tPA) and goat anti–mouse and anti–rabbit peroxidase-conjugated antibodies were from Calbiochem. Monoclonal anti-fibrin(ogen) antibody (59D8) was the generous gift of Drs Marschall Runge (University of North Carolina [UNC] Department of Medicine) and Charles Esmon (Oklahoma College of Medicine). Biotinylated secondary antibodies were from Vector Laboratories. Target Retrieval Solution was from Dako North America. Nonimmune mouse IgG antibody (MOPC-1), bovine serum albumin (BSA), and adenosine diphosphate were from Sigma-Aldrich. Recombinant human tumor necrosis factor α (TNF-α) was from Millipore. Corn trypsin inhibitor (CTI) and factor X were from Haematologic Technologies. Fibronectin, plasminogen-, and von Willebrand factor (VWF)–depleted fibrinogen was from Enzyme Research Laboratories. Fibrinogen was further depleted of factor XIII by immunoadfinity chromatography. The AlexaFluor-488 protein labeling kit was from Invitrogen. AlexaFluor-488-labeled fibrinogen (~8 mol fluorophore/mol fibrinogen) was prepared as described.18 Thrombin receptor activation peptide (serine-phenylalanine-leucine-arginine-asparagine, TRAP) was from Bachem. Collagen was from Chrono-Log. Tenecteplase (TNKase) was the generous gift of Genentech. Contact-inhibited normal pooled plasma (NPP) was prepared from whole blood from 40 healthy subjects (50% female, 68% nonwhite) in a protocol approved by the UNC Institutional Review Board.19 The fibrinogen concentration in human NPP (3 mg/mL) was determined by enzyme-linked immunoabsorbent assay.

Murine thrombosis and thrombolysis models

Procedures were approved by the UNC Institutional Animal Care and Use Committee. Mice (6- to 8-week-old male C57BL/6, Charles River Laboratories) were anesthetized with 1.5% to 2% isoflurane in 2% oxygen, and the left saphenous vein was exposed under a SZX12 dissecting microscope (Olympus) using a catheter constructed of pulled PE-10 tubing (Braintree Scientific) with a 3.0-ml (0.076-mm diameter) cleaning wire (Hamilton) placed into the lumen as a stylet, as described.20 Human fibrinogen or vehicle (20 mM N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic acid, pH 7.4, 150 mM NaCl [HBS] or HBS/BSA) was administered through the cannula on a per-weight basis (blood volume [milliliters] is 7% of body weight [grams]; plasma is 50% of blood volume) to achieve 135% or 170% of normal (endogenous murine fibrinogen plus infused human fibrinogen) in murine circulation 5 minutes before injury. Final plasma fibrinogen levels were measured as described21 in mice not subject to FeCl3 injury. Nanobead diffusion experiments showed that these fibrinogen concentrations did not alter plasma viscosity (R. Spero, unpublished observation, September 2010).

For carotid artery thrombosis, the right common carotid artery was exposed after midline cervical incision. A Doppler transonic flow probe (Transonic Systems) was applied and connected to a flow meter (model T206; Transonic Systems) supplying a data acquisition system (PowerLab 4/30 model ML866, AD Instruments). The carotid artery was dried and 10% FeCl3 (0.62 M FeCl3 on 0.5 × 0.5-mm filter paper) placed on the vein for 3 minutes, removed, and tissues washed 3 times with warm saline. After injury, blood flow was continuously monitored. For saphenous vein thrombosis, the saphenous vein of the right leg was dissected and exposed, 5% FeCl3 (0.31 M FeCl3 on 0.5 × 2-mm filter paper) placed on the vein for 3 minutes, removed, and tissues washed 3 times with warm saline. Blood flow was monitored auditorily by Doppler ultrasonic flow probe. In both models, the TTO was the time between FeCl3 administration and lack of flow for 60 consecutive seconds. Experiments were stopped at 45 minutes if no occlusion occurred. Occluded vessels were excised and fixed in 10% formalin.

Thrombolysis was assessed in mice subject to FeCl3 carotid artery thrombosis. After 5 consecutive minutes of blood flow less than 0.1 mL/min, mice were infused with TNKase (0.5-5 mg/kg) through the saphenous vein intravenous catheter while continuously monitoring carotid blood flow.

Hematoxylin and eosin staining and immunohistochemistry

Fixed tissues were dehydrated and paraffin-embedded, and consecutive, 5-µm sections cut and mounted with vectamount (UNC Lineberger Comprehensive Cancer Center Animal Histopathology Core). Slides were stained with hematoxylin and eosin to visualize the thrombus, and imaged with a Retiga 400R camera (Qimaging) linked to an Olympus Bx 61 microscope with a 20× U Plan FL N, 0.5 NA objective lens. A computer equipped with Velocity software (v5.5) was used to operate the system. Images were analyzed with Adobe Photoshop CS (v8.0). For immunohistochemistry, antigen retrieval was performed in Target Retrieval Solution in a 95°C water bath. Slides were stained with anti-fibrin antibody (59D8, 1:1000), which detects both human and mouse fibrinogen15,22 (supplemental Figure 1, available on the Blood Web site; see the Supplemental Materials link at the top of the online article) for 1 hour at room temperature in a humidity-controlled chamber and developed using avidin-biotin complex (Dako North America). Negative controls were stained simultaneously in the absence of primary antibody. Staining intensity of thrombi from at least 4 representative sections per mouse were analyzed by 3 independent, blinded observers on a scale of 0 to 3.

Platelet aggregation

Human platelet-rich plasma (PRP) obtained by centrifugation (150g, 15 minutes) of citrated blood was adjusted to 250 × 10^3 platelets/mL with autologous platelet-poor plasma (PPP). Aggregation was triggered by TRAP (50 µg/mL, final), collagen (2 µg/mL, final), or adenosine diphosphate (2.5 µM, final). Light transmission was recorded on a Chrono-Log Optical Aggregometer, model 470.

Cell culture

Primary human saphenous vein endothelial cells (HSVECs, PromoCell) and human smooth muscle cells (SMCs; Lonza Walkersville) were cultured as directed to 80% to 95% confluence in 5% CO2 at 37°C. Cells were used between passages 3 to 6 to reduce phenotypic drift.

Cellular activity assays

TF activity was measured by chromogenic substrate cleavage on a SpectraMax Plus340 plate reader (Molecular Devices) in the presence and absence of inhibitory anti-TF antibody (HTF-1, 10 µg/mL) or isotype control (MOPC-1, 10 µg/mL), as described.19 Thrombin was measured by calibrated automated thrombography using a Fluoroskan Ascent fluorometer (ThermoLabsystem) as described.19 Thrombin generation was calculated using Thrombinscope software Version 3.0.0.29 (Thrombinscope BV). The thrombin generation rate was calculated by dividing peak height by the difference from time to peak to lag time. The endogenous thrombin
potential is not reported because thrombin generation curves did not always return to baseline.

**Phospholipid vesicles**

Phosphatidylcholine (egg), phosphatidylethanolamine (soy), and phosphatidylserine (porcine brain) were from Avanti Polar Lipids. Large unilamellar vesicles (41% phosphatidylcholine/44% phosphatidylethanolamine/15% phosphatidylserine) were made as described. Briefly, lipids were combined, dried under nitrogen gas, and resuspended in cyclohexane. Resuspended lipids were lyophilized, resuspended in HBS containing 1mM ethylenediaminetetraacetic acid, and extruded through a 0.2-μm filter 10 times.

**Clot formation and lysis by turbidity**

For human plasma experiments, recalciﬁed (16mM, ﬁnal), lipidated (4μM, ﬁnal) NPP was spiked with ﬁbrinogen to 4.5, 6, or 7.5 mg/mL (150%, 200%, and 250% of normal, respectively, and ﬁnal) immediately added to washed cell monolayers (67.7% plasma, ﬁnal). For mouse plasma experiments, recalciﬁed (16mM, ﬁnal) murine PPP containing 2.4 ± 0.2 mg/mL ﬁbrinogen (100%) was spiked with 2 or 4 mg/mL human ﬁbrinogen to approximately 180% and 270% of normal, respectively, diluted 1:3 in HBS, and clotted with TF (Innovin 1:30 000 ﬁnal). Fibrinolysis assays included tPA (250 μg/mL, ﬁnal) or TNKase (concentrations indicated) at the reaction start. Clotting and lysis were detected by turbidity at 405 nm in a SpectraMax Plus340 plate reader.18,19

**Laser scanning confocal microscopy**

Clots were formed over washed cells in Lab-Tek II Chamber #1.5 coverglasses (Nalge Nunc International) with addition of AlexaFluor-488-labeled ﬁbrinogen as described. Clotting proceeded until a constant ﬁnal turbidity was reached in separate, parallel reactions. Clots were imaged on a Zeiss LSM5 Pascal laser scanning confocal microscope (Carl Zeiss Inc.) linked to a Zeiss Axiovert 200M microscope equipped with a Zeiss 60×/1.4 NA oil immersion plan apo-chromatic lens, as described. The 488nm line of a medium power multi-line argon ion laser was used for excitation and a 505-530-nm band-pass ﬁlter for emission. A computer equipped with Carl Zeiss software (v1.5) was used to operate the system. Optical sectioning was achieved by closing the pinhole in the front of the detector to one airy unit. The zoom factor was 1. Thirty optical sections (1024 × 1024 pixels) in 3 randomly chosen locations were collected at 0.36-μm intervals in the z-axis at the cell surface. Image volumes were 146 × 146 × 10 μm. Single images were collected in 15.47 seconds. Optical resolution was −0.14 μm in the xy-plane and −0.5 μm on the z-axis. The sectioning interval was smaller than the calculated z-axis optical section resolution to achieve Nyquist sampling in z based on the Zeiss software calculation. Images were deconvolved using 3-dimensional deconvolution algorithms in AutoQuant Autodeblur (Version x1.4.1; Media Cybernetics). Fibrin network density was analyzed using ImageJ (1.37V; National Institutes of Health) by placing random grids of 2 pixel crosses on individual slices (121-144 crosses/slice) and counting ﬁbers intersecting the middle of the crosses divided by the total number of crosses, less crosses in the volume occupied by the cells, as described.18,19

**Clot viscoelastometry**

Human PRP and PPP from blood drawn into 3.2% sodium citrate/18.3 μg/mL CTI was recalciﬁed (16mM, ﬁnal) and spiked with ﬁbrinogen or BSA in HBS. PRP and PPP had more than 300 × 107 and less than 8 × 107 platelets/mL, respectively. The baseline PRP and PPP ﬁbrinogen concentration was estimated in accord with NPP determinations. PRP was clotted with TF (1pM, ﬁnal), which corresponds to the TF activity of TNF-α–stimulated HSVECs (TNF-α-HSVECs). PPP was clotted with 1pM TF and 4μM phospholipid. Lysis assays were performed in the presence of tPA (500 ng/mL, ﬁnal). Clot elastic modulus (CEM) was measured by Hemodyne HAS.24

**Statistical methods**

For TTO and immunohistochemistry, normal and hyperfibrinogenemic conditions were compared by Mann-Whitney test. For CEM, normal and hyperfibrinogenemic conditions were compared by unpaired Student t-tests. For clotting assays, ﬁbrin density, and ﬁbrinolysis parameters, signiﬁcant differences between groups were identiﬁed by one-way analysis of variance and analyzed by Dunnett post-hoc test using 3 mg/mL ﬁbrinogen as the index group (on unstimulated HSVECs, SMCs, or TNF-α-HSVECs as indicated) to limit type I error. Statistical analyses were performed using Kaleidagraph, Version 4.1 (Synergy Software).

**Results**

**Hyperfibrinogenemia shortens the time to vessel occlusion after FeCl3 injury**

To determine the contribution of elevated ﬁbrinogen to intravascular thrombus formation, we used 2 murine thrombosis models based on FeCl3 application to the carotid artery (high shear) or saphenous vein (lower shear) after intravenous infusion of human ﬁbrinogen. This infusion-based approach enabled us to precisely control the plasma ﬁbrinogen level during thrombus formation. Control experiments and published studies demonstrate that human ﬁbrinogen is incorporated into murine clots, supports murine platelet aggregation, and has appropriate half-life in mouse circulation. The endogenous ﬁbrinogen concentration in mice was 2.4 ± 0.2 mg/mL (100%), and levels were raised to 3.2 ± 0.2 or 4.0 ± 0.1 mg/mL, ﬁnal (135% or 170% of normal, respectively), consistent with levels associated with thrombosis in humans.

Consistent with transgenic hyperfibrinogenemic mice13,15 and a prior study in which human ﬁbrinogen was injected into BALB/c mice,26 elevated ﬁbrinogen did not trigger spontaneous thrombosis (Figure 1). FeCl3 application to the carotid artery or saphenous vein of HBS-infused mice produced occlusive thrombi in 9.3 and 17 minutes (median values), respectively, conﬁrming the prothrombotic effect of vascular disruption. Infusion of control protein (BSA) did not further shorten the TTO in either vessel (data not shown). Mice infused with ﬁbrinogen to 3.2 ± 0.2 mg/mL (135%) demonstrated a nonsigniﬁcant trend to shorter TTO in the carotid artery model (data not shown); we did not test this ﬁbrinogen concentration in the saphenous vein model. Interestingly, compared with control (HBS or BSA infusion), ﬁbrinogen infusion to 4.0 ± 0.1 mg/mL ﬁnal (170%) before FeCl3 injury signiﬁcantly (P < .005) shortened the TTO in both carotid artery and saphenous vein models (median, 6.4 and 14.2 minutes, respectively, Figure 1). The shortened TTOs were not the result of factor XIII in the ﬁbrinogen preparation; infusion of factor XIII-depleted ﬁbrinogen also signiﬁcantly (P < .05) shortened the TTO versus control mice (Figure 1). These data demonstrate a direct contribution of hyperfibrinogenemia to thrombus formation after vascular injury.

**Elevated ﬁbrinogen increases the ﬁbrin(ogen) content of thrombi**

To probe the mechanism for the shortened TTO, we ﬁrst examined the morphology of thrombi formed in the murine carotid artery and saphenous vein. Hematoxylin and eosin staining showed extensive, occlusive thrombi containing distinct regions of proteinaceous material and erythrocytes in injured vessels from both control and ﬁbrinogen-injected mice (Figure 2A-B). Immunohistochemistry of thrombi in carotid artery and saphenous vein thrombi from control (HBS or BSA-infused) mice demonstrated weak ﬁbrin staining concentrated primarily at the luminal edge of proteinaceous
regions. No staining was detected in the absence of primary antibody, confirming that the secondary antibody did not bind mouse tissue nonspecifically (Figure 2A-B). Staining was slightly more intense in thrombi from saphenous vein than carotid artery, consistent with higher fibrin production in lower shear conditions.27 Both carotid artery and saphenous vein thrombi from mice infused with human fibrinogen demonstrated more intense fibrin staining at the periphery of proteinaceous regions and intense, diffuse staining in regions containing erythrocytes (Figure 2). These findings suggest that hyperfibrinogenemia increased thrombus fibrin content in both high and low shear vessels.

In vitro platelet aggregation is not increased by elevated fibrinogen levels

We then tested whether hyperfibrinogenemia increased platelet aggregation, a process dependent on fibrinogen binding to platelet αIIbβ3. The endogenous fibrinogen concentration of human plasma was 3 mg/mL (100%), and fibrinogen levels were raised with additional human fibrinogen as indicated. As shown in supplemental Figure 2 and supplemental Table 1, elevated fibrinogen did not increase, and even slightly decreased, platelet aggregation induced by TRAP, collagen, or adenosine diphosphate, consistent with a mechanism where high fibrinogen promotes full occupancy of platelet αIIbβ3 and inhibits interplatelet bridging.28 These data suggest that the mechanism by which hyperfibrinogenemia shortened the TTO was not via enhanced platelet aggregation.

Both cellular PCA and elevated fibrinogen increase fibrin formation

We next determined the effect of fibrinogen level on fibrin formation in models of vasculature by incubating cultured cell monolayers with recalcified NPP spiked with fibrinogen. We used unstimulated HSVECs to model unperturbed endothelium and SMCs to model FeCl3-injured vessels. We also used TNF-α-stimulated HSVECs (100 ng/mL, TNF-α for 6 hours, TNF-α-HSVECs) to model intact cytokine-stimulated endothelium thought to promote clotting in venous thrombosis (supplemental Figure 3). Whereas quiescent endothelial cells in vivo do not express significant TF, cultured unstimulated endothelial cells exhibit low TF activity.19 However, the net procoagulant activity (PCA) of cultured HSVECs was significantly lower than cultured SMCs or TNF-α-HSVECs (supplemental Figure 3; Table 1).

These experiments demonstrated unique effects of cellular PCA and fibrinogen level on fibrin formation (Figure 3A-C). Relative to unstimulated HSVECs, increased PCA of SMCs and TNF-α-HSVECs (supplemental Figure 3; Table 1) significantly (P < .002) shortened the fibrin formation onset (Figure 3D), consistent with the premise that exposure of procoagulant cells to blood triggers clotting. Elevated fibrinogen trended toward a shortened clotting onset in reactions induced by unstimulated HSVECs but did not reach statistical significance because of large variability. In addition, fibrinogen did not further shorten the onset in reactions triggered by the more procoagulant SMCs or TNF-α-HSVECs. Elevated fibrinogen did, however, significantly (P < .001) increase peak turbidity (Figure 3E), indicating increased fibrinogen incorporation into clots. Interestingly, relative to NPP clots produced by unstimulated HSVECs, both cellular PCA (SMCs, P = .08; and TNF-α-HSVECs, P < .02) and hyperfibrinogenemia (P < .001) increased the fibrin formation rate (Figure 3F). We observed a similar increase in the rate and final turbidity of TF-initiated clotting of murine plasma spiked with human fibrinogen (Figure 3G). This increase in the rate and amount of fibrin production is consistent with the shortened TTO observed in the murine thrombosis model (Figure 1).

Both cellular PCA and elevated fibrinogen increase fibrin network density

We and others have previously correlated fibrin formation parameters with fibrin network structure.18,19,29,30 To assess the impact of elevated fibrinogen on fibrin structure, we used laser scanning confocal microscopy to examine NPP clots produced by unstimulated HSVECs, SMCs, and TNF-α-HSVECs with increasing fibrinogen concentrations. Clots produced by SMCs and TNF-α-HSVECs were composed of more densely packed fibers (P < .05) than those produced by unstimulated HSVECs (Figure 4), consistent with observations that high thrombin generation promotes formation of dense networks of thin fibers.18,19,20 Notably, fibrin network density also correlated positively and significantly (P < .001) with fibrinogen concentration in both human (Figure 4A-B) and murine (Figure 4C) plasmas, suggesting that elevated fibrinogen levels promote formation of abnormally dense fibrin networks.

Elevated fibrinogen increases clot mechanical strength

Fibrin network structure determines clot viscoelasticity,24,30 suggesting that elevated fibrinogen alters clot mechanical properties. To
assess the impact of fibrinogen level on clot elasticity, we measured the CEM in clots forming in human PRP or PPP spiked with fibrinogen to 6 mg/mL, final (200%) or BSA (control). Because these assays cannot be initiated by cell monolayers, we initiated reactions with 4 μM phospholipid and 0 (to model unstimulated endothelium) or 1pM (to model vascular disruption) TF. No CEM developed in the absence of added TF, reflecting the requirement for a procoagulant stimulus (ie, vascular disruption) to initiate clotting, even in hyperfibrinogenemic conditions. Initiation of clotting by TF showed a significant (P < .05) fibrinogen-dependent increase in peak CEM in both PRP and PPP (Figure 5A), suggesting that elevated fibrinogen increases the structural integrity (mechanical strength) of the clot.

Elevated fibrinogen increases plasma clot resistance to fibrinolysis

Fibrin network density also determines a clot’s resistance to fibrinolysis.18,19,32 To evaluate the effect of elevated fibrinogen on clot resistance to fibrinolysis, we used both turbidimetric and mechanical (elastometry) lysis assays. We first initiated clotting by incubating human NPP spiked with fibrinogen or HBS with HSVECs, SMCs, and TNF-α-HSVECs in the presence of tPA and monitored clotting and lysis by turbidity (Figure 5B-D). Compared with unstimulated HSVECs, SMCs and TNF-α-HSVECs significantly (P < .05) shortened the time to peak turbidity of NPP clots (Figure 5E). In contrast, elevated fibrinogen significantly (P < .05) shortened the time to peak turbidity only on unstimulated HSVECs but increased peak turbidity on all cell types. We observed a similar pattern in mouse PPP spiked with human fibrinogen (Figure 5G). These data indicate that cellular PCA triggers fibrin formation, but fibrinogen concentration dictates fibrin incorporation into the clot and its resistance to lysis.

We then evaluated the effects of elevated fibrinogen on mechanical stability during tPA-mediated lysis of human PRP and PPP...
clots. Compared with control, addition of fibrinogen to 6 mg/mL, final (200%) increased peak CEM, area under the lysis curve, and half-lysis time (Figure 5H; Table 2). Because fibrinogen produced similar effects in both PRP and PPP, these assays support findings that elevated fibrinogen promotes resistance of clots to fibrinolytic and mechanical disruption by a fibrin-dependent, not platelet-dependent, mechanism.

Elevated fibrinogen increases resistance to thrombolysis in vivo

Finally, because in vitro experiments suggested that elevated fibrinogen increased clot stability, we assessed thrombolysis in vivo in mice with normal and elevated fibrinogen. Because of the small size of the saphenous vein and its proximity to the saphenous artery, it was not feasible to continuously monitor blood flow during thrombolysis in the saphenous vein. Therefore, thrombolysis was only performed in the carotid artery model. After occlusive thrombus formation, we initiated lysis via bolus infusion of the tPA analog TNKase. TNKase’s increased fibrin specificity and enhanced resistance to inhibition by plasminogen activator inhibitor-1 provide a longer plasma half-life and facilitate dosing regimens (single IV bolus vs continuous infusion required for tPA/alteplase). Lysis of murine PPP clots was TNKase-dose-dependent (supplemental Figure 4A). Importantly, human fibrin and murine fibrin are similarly cleaved by murine plasmin, and TNKase-mediated lysis was similarly dose-dependent with regard to fibrinogen (human, supplemental Figure 4B; or mouse, supplemental Figure 4C) spiked into murine PPP.

TNKase infusion produced dose-dependent lysis of in vivo thrombi; 0.5, 2.5, and 5 mg/kg TNKase produced no (2 of 2 mice), partial (30%-70% in 2 of 2 mice), or complete (5 of 5 mice) return of flow, respectively, in vehicle-infused mice within 1 hour of infusion (Figure 6A-F). Mice infused to 3.2 ± 0.2 mg/mL final plasma fibrinogen (135%) showed only partial (0%-30%) return of
Discussion

Although epidemiologic studies have implicated elevated plasma fibrinogen as an independent risk factor for both arterial and venous thrombosis, it remains highly controversial whether fibrinogen is merely a biomarker of a coincident inflammatory state or is also a causative mechanism in the etiology. We addressed this question in vivo by acutely elevating fibrinogen in mice and examining their susceptibility to thrombosis, and used ex vivo and in vitro experiments to elucidate biochemical and biophysical mechanisms by which fibrinogen contributes to thrombus formation and stability. We also used a murine thrombolysis model to test thrombus stability in vivo. Our findings show that elevated fibrinogen levels: (1) specifically and independently shortened the TTO in both high (arterial) and lower (venous) shear injury models, (2) increased thrombus fibrin content, (3) promoted faster fibrin formation, higher network density, and increased clot strength and stability in TF-initiated in vitro models that recapitulate in vivo thrombosis, and (4) increased thrombus resistance to lysis in vivo. Together, these findings show that hyperfibrinogenemia independently promotes thrombosis, identify biochemical and biomechanical mechanisms by which elevated fibrinogen levels contribute to thrombus formation, and demonstrate a critical role for plasma fibrinogen levels in resistance to thrombolytic therapy.

Our study is the first to show elevated fibrinogen significantly and independently shortens the TTO after acute injury. These findings differ somewhat from observations that transgenic hyperfibrinogenemic mice demonstrate a nonsignificant trend to shorter time to 75% occlusion in a FeCl₃-carotid model. Differences in the 2 studies include the method of elevating fibrinogen levels (infusion vs transgenic expression) and final fibrinogen levels (170% vs 145%). In our study, mice infused with lower fibrinogen levels (to 135%) showed a TTO of 9 minutes (median) after carotid artery injury, which was not significantly different from control mice (data not shown). In addition, Kerlin et al used mice with elevated murine fibrinogen, whereas the mice in our study had a mixture of human and mouse fibrinogen. Differences between these 2 molecules may make human fibrinogen more prothrombotic. Finally, the higher FeCl₃ concentrations (20%) used in the earlier study may have dampened differences between normal and hyperfibrinogenemic mice. Our findings also differ somewhat from observations that human fibrinogen ϴ-chain expression in mice paradoxically decreases thrombus weight after electrolytic femoral injury. However, the total fibrinogen levels achieved in human ϴ-expressing mice were lower than in controls, so any role of hyperfibrinogenemia was not examined in that study. Our in vivo observations are supported by ex vivo and in vitro cell-based experiments that recapitulate aspects of the murine thrombosis models. Notably, these findings are consistent with studies demonstrating fibrin network density, mechanical stability, and resistance to lysis are positively correlated with fibrinogen concentration in clotting experiments that trigger thrombus formation and stability.

flow in 3 of 3 mice with subsequent reclosure after 5 mg/kg TNKase infusion (Figure 6G-H). In contrast, mice infused to 4.0 ± 0.1 mg/mL final plasma fibrinogen (170%) did not reacquire flow for up to 120 minutes (3 of 3 mice), even at the highest TNKase dose or with repeat dosing (Figure 6I-J). Together with in vitro lysis data (Figure 5), these in vivo data demonstrate that hyperfibrinogenemia promotes resistance to thrombolysis.
thrombotic disease and suggest specific mechanisms by which elevated fibrinogen is pathogenic.

Table 2. Elevated fibrinogen increased clot elastic modulus during lysis

<table>
<thead>
<tr>
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<th>Peak CEM</th>
<th>Area under the curve</th>
<th>Half-lysis</th>
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<tbody>
<tr>
<td>PPP</td>
<td>1</td>
<td>1</td>
<td>1</td>
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<tr>
<td>PPP + fibrinogen</td>
<td>2.7 ± 0.6</td>
<td>3.9 ± 1.5</td>
<td>1.3 ± 0.1</td>
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<tr>
<td>PRP</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>PRP + fibrinogen</td>
<td>2.2 ± 0.6</td>
<td>2.6 ± 1.3</td>
<td>1.3 ± 0.5</td>
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Data are presented as fold increase over PPP or PRP with no additional fibrinogen.

\*P < .05 versus no additional fibrinogen.

Our study correlating abnormal fibrin quality with thrombosis in hyperfibrinogenemia informs several patient cohort studies. Of note, both hyperfibrinogenemia and denser, fibrinolysis-resistant clots were observed in plasma from patients with coronary artery disease, unifying observations of elevated fibrinogen, abnormal clot quality, and cardiovascular disease in a single patient cohort. Abnormal fibrin network structure and/or function (high mechanical strength and/or increased resistance to lysis) have also been detected in plasma clots from patients with diabetes, ischemic stroke, pulmonary hypertension, myocardial infarction, venous thromboembolism, and healthy relatives of patients with premature coronary artery disease. Overall, these studies suggest that abnormal fibrin quality is a pathologic mechanism operant not
only in hyperfibrinogenemia, but also in other prothrombotic pathologies.

A particularly interesting outcome of our study is the observation that hyperfibrinogenemia did not cause spontaneous thrombosis in vivo. This observation is consistent with reports that injecting human fibrinogen into mice does not cause spontaneous fibrin deposition and, importantly, with findings that, although hyperfibrinogenemia is a thrombosis risk factor, it does not cause thrombosis \textit{ipso facto}. Virchow proposed that multiple “hits” from abnormalities in plasma composition, vascular cell function, and blood flow are required to trigger thrombosis. This paradigm suggests that an initiating trigger, probably exposure of cellular PCA during plaque rupture or vasculitis, is required to initiate thrombus formation. Indeed, markers of vascular disruption, including circulating leukocyte and endothelial-derived microparticles, are elevated 1.3-fold in venous thromboembolism patients. Consistent with this hypothesis, the prognostic importance of elevated fibrinogen levels appears to be independent of, and additive to, myocardial damage (troponin-T levels) in patients with unstable CAD. To our knowledge, few diagnostic algorithms simultaneously consider markers of tissue damage and plasma hypercoagulability when assessing thrombosis risk. Of note is one

Figure 6. Hyperfibrinogenemia increases resistance to thrombolysis in vivo. Thrombosis was triggered in the carotid artery of wild-type mice infused with fibrinogen (plasminogen-, fibronectin-, VWF-, and factor XII-depleted, concentrations indicated in the figure) or vehicle control. After stable occlusion for 5 minutes, mice were infused with TNKase (concentrations indicated in the figure). Blood flow was monitored by flow probe throughout the experiment. Each panel shows data from an individual mouse, representative of at least 2 mice for each condition. Shaded grey area represents the time of FeCl₃ treatment plus time to reacquire flow.
study\(^1\) that examined independent and combined effects of elevated fibrinogen and homocysteine (a potential initiator of endothelial damage) on mortality of patients from a high-risk cardiology clinic; elevated levels of both homocysteine and fibrinogen contributed to an increased hazard ratio, consistent with our findings that endothelial dysfunction (TF expression) and hyperfibrinogenemia independently promoted thrombus formation. Reduced blood flow (stasis), the third component of the Virchow triad, is thought to explain differences in arterial and venous thrombosis. The shear rates between arteries and veins differ significantly; in humans, large arteries, such as the carotid, have wall shear rates of 300 to 800 inverse seconds, whereas venous shear rates are in the range of 20 to 200 inverse seconds.\(^4\) Interestingly, hyperfibrinogenemia has been correlated with both arterial and venous thrombosis, suggesting that hyperfibrinogenemia contributes to thrombus formation independently of the shear rate. We addressed the role of shear by applying the FeCl\(_3\) injury model to both high (artery) and lower (vein) shear vessels. Thrombi from saphenous veins appeared more fibrin-rich than thrombi from carotid arteries (Figure 2), consistent with observations that lower shear promotes fibrin deposition.\(^27\) Notably, however, hyperfibrinogenemnic mice demonstrated shortened TTDs and increased fibrin content in thrombi in both high and lower shear vessels (Figures 1, 2). These findings do not diminish a role for stasis in thrombosis but rather suggest that the contributions of shear are eclipsed in the setting of hyperfibrinogenemia and potentially other pathologies as well.

Thrombolytic therapy has met limited success. The GUSTO-I trial demonstrated that complete coronary artery perfusion determines 30-day survival after myocardial infarction; however, tPA therapy achieved perfusion in only 54% of patients.\(^47\) Similarly, in acute stroke, proximal arterial patency after tPA infusion correlated with positive long-term outcome but was achieved in only 27% of patients.\(^48\) Interestingly, although prior studies associated thrombus platelet content with decreased tPA efficacy,\(^49\) our study shows that, within a given thrombosis model, increased fibrin content also decreases thrombolytic efficacy. This novel finding suggests that the plasma fibrinogen level present during thrombus formation is an independent predictor not only of thrombotic risk, but also of the potential efficacy of thrombolysis. Of note is the experiment in which we saw no return of flow in the occluded carotid artery despite 2 doses of TNKase (Figure 6J); postmortem dissection revealed diffuse bleeding from microvasculature within the neck and abdominal cavity. Given the risk and devastating consequences of intracerebral hemorrhage with thrombolytic therapy, our data suggest that screening patients for fibrinogen levels may inform risk/benefit analysis before initiating thrombolysis. Confirming a role for fibrinogen level in thrombolysis will require a study of patients who received thrombolytic therapy and for whom both the fibrinogen level and degree of reperfusion are known. Our data provide strong justification for such an investigation.

This study has potential limitations. First, we used human fibrinogen to increase circulating levels in the mouse. However, published studies\(^17,25,26\) as well as our data demonstrate that human fibrinogen is stable in murine circulation and incorporated into murine clots. Moreover, elevating either human or mouse fibrinogen in mouse plasma increased peak turbidity and prolonged tPA- or TNKase-mediated fibrinolysis similar to that seen with tPA- or TNKase-mediated lysis of human fibrinogen in human clots (Figure 5; supplemental Figure 4). Second, the FeCl\(_3\) model may not fully recapitulate thrombosis/thromboembolism; it will be interesting to examine the effects of hyperfibrinogenemia in other models (stasis- or electrolytic-based) in future studies. Finally, we evaluated the immediate effects of fibrinogen on thrombosis; however, elevated fibrinogen may have additional prothrombotic effects in vivo. Previous studies\(^50\) demonstrated fibrin induction of TF in human vascular ECs, suggesting that prolonged exposure of vasculature to hyperfibrinogenemia may feedback on additional cellular mechanisms.

In conclusion, our results show that hyperfibrinogenemia independently promotes thrombus formation and stability via increased fibrin network density and resistance to dissolution. These findings establish hyperfibrinogenemia in the etiology of both arterial and venous thrombosis/thromboembolism and suggest that fibrin is a potential therapeutic target in the management of these pathologies. Furthermore, our study establishes a model for future investigations of plasma hypercoagulability and vascular dysfunction; modulating plasma composition via intravenous procoagulant infusion will allow examination of the specific roles of additional plasma proteins in thrombosis and thrombolysis.

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Authorship

Contribution: K.R.M. performed experiments, analyzed data, and wrote the manuscript; J.C.C. performed experiments; F.C.C. provided reagents and reviewed the manuscript; and A.S.W. designed and supervised the study, analyzed data, and wrote the manuscript.

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References

Causal relationship between hyperfibrinogenemia, thrombosis, and resistance to thrombolysis in mice

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