Neutrophil histone modification by peptidylarginine deiminase 4 is critical for deep vein thrombosis in mice

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Deep vein thrombosis and pulmonary embolism are major health problems associated with high mortality. Recently, DNA-based neutrophil extracellular traps (NETs) resulting from the release of decondensed chromatin, were found to be part of the thrombus scaffold and to promote coagulation. However, the significance of nuclear decondensation and NET generation in thrombosis is largely unknown. To address this, we adopted a stenosis model of deep vein thrombosis and analyzed venous thrombi in peptidylarginine deiminase 4 (PAD4)-deficient mice that cannot citrullinate histones, a process required for chromatin decondensation and NET formation. Intriguingly, less than 10% of PAD4−/− mice produced a thrombus 48 h after inferior vena cava stenosis whereas 90% of wild-type mice did. Neutrophils were abundantly present in thrombi formed in both groups, whereas extracellular citrullinated histones were seen only in thrombi from wild-type mice. Bone marrow chimera experiments indicated that PAD4 in hematopoietic cells was the source of the prothrombotic effect in deep vein thrombosis. Thrombosis could be rescued by infusion of wild-type neutrophils, suggesting that neutrophil PAD4 was important and sufficient. Endothelial activation and platelet aggregation were normal in PAD4−/− mice, as was hemostatic potential determined by bleeding time and platelet plug formation after venous injury. Our results show that PAD4-mediated chromatin decondensation in the neutrophil is crucial for pathological venous thrombosis and present neutrophil activation and PAD4 as potential drug targets for deep vein thrombosis.

Histone citrullination | venous thromboembolism | inflammation | programmed cell death | nuclear swelling

Venous thromboembolism is associated with high mortality, with ~300,000 deaths resulting from an estimated 900,000 cases in the United States annually (1). Neutrophil extracellular traps (NETs) are produced by a cell death pathway shown to be important in innate immunity against microbes (2–5). Recently, NETs were also shown to be involved in thrombosis by binding erythrocytes and platelets (6) and in promoting coagulation by degrading tissue factor pathway inhibitor (7). Coagulation, platelets, neutrophils, and NETs have all been implicated in thrombosis generation in mouse models of deep vein thrombosis (DVT) (8–10). Also, certain cancers associated with neutrophilia result in an increased propensity of circulating neutrophils to form NETs and spontaneous thrombosis (11). NETs are composed of decondensed chromatin fibers lined with antimicrobial proteins such as neutrophil elastase and myeloperoxidase (2). A proteomic analysis identified histones as the major protein component of NETs (12). Histones are highly cationic to the surrounding environment into which NETs are released (13, 14) and can enhance thrombin generation (15). Histone infusion leads to rapid Weibel–Palade body (WPB) secretion from endothelial cells, promoting platelet and leukocyte adhesion and leading to accelerated thrombus generation (9). Pretreatment with DNase 1, which can degrade NETs, reduces thrombus incidence in wild-type (WT) mice (9, 10). Neutrophils play an important role, as neutrophil depletion greatly reduces thrombus weight in this model (10). Whether NETs are involved in the pathogenesis of DVT or whether they are merely a consequence of leukocyte recruitment to the thrombus is unknown. Similarly, the role of NETs in normal platelet plug formation in response to injury, where their presence was also observed (7), remains to be addressed.

Although the cellular changes occurring in the neutrophil during the process of generating neutrophil extracellular traps (NETosis) have been well described, the molecular players and processes leading to NET formation are incompletely defined. A critical step in NET formation is the decondensation of chromatin that occurs in the nucleus. Peptidylarginine deiminase 4 (PAD4) is a nuclear enzyme that converts specific arginine residues to citrulline on histone tails. Upon PAD4 activation, histones can become hypercitrullinated at histone H3Arg8 and -17 or at histone H4Arg3 residues, resulting in the extensive chromatin decondensation that leads to nuclear delobulation and swelling during NETosis (16, 17). PAD4-deficient mice are incapable of decondensing chromatin or forming NETs (18), whereas overexpression of PAD4 is sufficient to drive chromatin decondensation to form NET-like structures in cells that normally do not form NETs (19). Because NETs are present in pathological thrombi and their dissolution by DNase 1 reduced the frequency of DVT, it was important to address the role of histone modification by PAD4 in the process of NET formation and its involvement in thrombosis.

Results

PAD4−/− Neutrophils Fail to Produce NETs in Response to Lipopolysaccharide or Lysosomal Vacuoles

PAD4 is a calcium-dependent enzyme with a nuclear localization signal that distinguishes it from other PAD family members (20, 21). The enzyme is activated by Ca2+ (17), and therefore the calcium ionophore ionomycin was used to induce histone hypercitrullination in isolated peripheral blood murine neutrophils (Fig. 1A). This led to NET formation in WT and PAD4+/− neutrophils, whereas PAD4−/− neutrophils did not citrullinate histone H3 or form NETs (Fig. 1A and B). Lipopolysaccharide (LPS) from Klebsiella pneumoniae was also unable to induce histone citrullination in isolated peripheral blood murine neutrophils (Fig. 1A and B). Moreover, immunostaining showed citrullinated histone H3 (H3Cit)-positive NETs being released from WT neutrophils, whereas in PAD4+/− neutrophils H3Cit staining was absent (Fig. 1C). Although PAD4−/− nuclei lost their lobulated shape characteristic of neutrophils, they did not swell, indicating that nuclear decondensation is not occurring.
in PAD4<sup>−/−</sup> mice and confirming that other PAD enzymes do not compensate for PAD4 deficiency in vitro. Ionomycin-induced NETosis has been shown to be independent of reactive oxygen species (22), and here we show that it is dependent on PAD4. Thus, PAD4<sup>−/−</sup> neutrophils have shown a complete inability to produce NETs (18) (Fig. 1B).

**PAD4<sup>−/−</sup> Mice Are Protected in the Venous Stenosis Model of DVT.** Biomarkers of NETs are abundant in baboon (6) and mouse DVT (9). We performed the venous stenosis model of DVT in mice (8). Briefly, the inferior vena cava (IVC) is ligated to induce a 90% restriction in blood flow. This results in the formation of a thrombus that is macroscopically similar to human deep vein thrombosis, containing a platelet-rich white portion (distal to the ligation site) and an erythrocyte-rich red portion (proximal to the ligation site). Because PAD4-deficient mice do not form NETs in vivo, we hypothesized that NET formation would be greatly impaired in these mice during thrombosis and that this may affect thrombus formation and/or stability. PAD4<sup>−/−</sup> mice were partially protected from producing venous thrombi early after stenosis, with only 28.6% of PAD4<sup>−/−</sup> mice forming thrombi at 6 h compared with 66.7% of WT mice (P = 0.04) (Fig. 2A). Thrombi that formed in PAD4<sup>−/−</sup> mice were similar in length to WT thrombi (Fig. 2B). PAD4<sup>−/−</sup> mice also maintained normal platelet counts compared with control sham-operated animals and after stenosis had significantly higher platelet counts than WT mice (Fig. 2C). In contrast to the diffuse extracellular H3Cit pattern in thrombi previously described at 48 h (9), immunostaining revealed that the majority of H3Cit was nuclear rather than extracellular in the WT thrombi at this early time point (Fig. S1A), whereas H3Cit staining was not detected in PAD4<sup>−/−</sup> thrombi (Fig. S1B) (WT, 28.3 ± 0.75 H3Cit<sup>−</sup> cells per mm<sup>2</sup>; PAD4<sup>−/−</sup>, no H3Cit<sup>−</sup> cells detected; n = 4). The thrombi presented similar histological morphologies with comparable density of leukocytes as seen by H&E staining (Fig. S1C). To see if this early protection could be a result of delayed thrombus formation in PAD4<sup>−/−</sup> mice, we next maintained stenosis for 48 h, when the great majority of WT mice form a thrombus (8), and found that, whereas 90% of WT stenotic vessels thrombozed, fewer than 10% of PAD4<sup>−/−</sup> mice had a thrombus at this time point (P = 0.0002) (Fig. 2 D and E). Thrombocytopenia was associated with thrombus formation in WT mice, whereas PAD4<sup>−/−</sup> mice retained normal platelet levels (Fig. 2F). Thus, platelets are likely consumed by the thrombus. It is important to note that at baseline WT and PAD4<sup>−/−</sup> mice had similar platelet counts (Fig. 2F). The extracellular H3Cit meshwork seen in WT thrombi at 48 h postligation was again completely absent in the sole PAD4<sup>−/−</sup> thrombus, indicating also that in vivo only PAD4 modifies histone H3 to produce H3Cit (Fig. 2 G and H). Hematoxylin and eosin staining revealed many swollen neutrophil nuclei in WT thrombi, whereas the PAD4<sup>−/−</sup> thrombus contained smaller and more dense neutrophil nuclei (Fig. 2I). Neutrophils are abundantly present within WT thrombi at 48 h (9) and also in the only thrombus in the PAD4<sup>−/−</sup> mouse (Fig. 2D), suggesting that neutrophils are recruited to the PAD4<sup>−/−</sup> thrombus. NETs were absent in the single small PAD4<sup>−/−</sup> thrombus that we obtained and analyzed by immunofluorescence staining. Thus, PAD4 deficiency was highly protective in the DVT mouse model at 48 h when, in WT mice, extracellular chromatin is prominently displayed (9) (Fig. 2G).

**Endothelial Activation and Platelet Function Are Not Affected by PAD4 Deficiency.** Although PAD4 is highly expressed in neutrophils, it is also present at lower levels in many cell types; therefore, citrullination of histones may regulate gene expression (17, 23). Therefore, it was important to establish in which cell type PAD4 was responsible for this prominent role in pathological thrombosis. First, we wanted to investigate whether endothelial release of adhesion molecules for platelets and leukocytes, von Willebrand factor (VWF) and P-selectin, from WPBs occurs normally in PAD4<sup>−/−</sup> mice. Both VWF and P-selectin were determined to be important in the stenosis model of DVT (8, 10). We turned to a time point (6 h) by which WPBs have been released, resulting in massive platelet and neutrophil adhesion to the endothelium (8). Platelet-leukocyte adhesion to endothelial cells at 6 h after stenosis (Fig. 3A) (WT, 8.97 ± 0.50; PAD4<sup>−/−</sup>, 13.56 ± 2.11 leukocytes per 200 μm of vessel wall; P = 0.10; n = 3). Another way to examine the capacity of mice to release WPBs is by infusion of activated platelets, which results in systemic WPB release from endothelial cells and consequently increases leukocyte rolling in veins (24). Leukocyte rolling occurred in both WT and PAD4<sup>−/−</sup> mice to a similar extent over baseline 2 h after active platelet infusion (Fig. 3A and B). This suggests that the protection of WT and PAD4<sup>−/−</sup> ICVs 6 h after stenosis (Fig. 3A) (WT, 8.97 ± 0.50; PAD4<sup>−/−</sup>, 13.56 ± 2.11 leukocytes per 200 μm of vessel wall; P = 0.10; n = 3) is not due to a defect in endothelial activation leading to VWF and P-selectin release, nor is it due to a defect in the initial leukocyte and platelet adhesion to the vessel wall that is essential for thrombus initiation in venous stenosis (8, 10). Because PAD4 could be expressed in megakaryocytes and thus affect the properties of platelets, we next evaluated the platelets from PAD4<sup>−/−</sup> mice. As Fig. 2F shows, at baseline the platelet count in PAD4<sup>−/−</sup> mice was normal. Platelets isolated from PAD4<sup>−/−</sup> mice aggregated normally in response to low and high doses of thrombin (Fig. 3 D and E). This shows that, in general, platelet activation, granule secretion, and integrin activation are not affected in these mice. We then examined in vivo models in which platelets respond to injury by forming a platelet plug; tail bleeding time and the ferric chloride model of thrombosis. Tail-tip transection results in severing of both veins and artery, and the rapid cessation of blood flow depends on platelet function and coagulation. Time to cessation of bleeding was similar in WT and PAD4<sup>−/−</sup> mice (Fig. 3F), indicating that these mice have normal hemostatic potential. In the venous stenosis model, flow restriction leads to WPB secretion potentially due to local hypoxia (25), resulting in platelet-leukocyte interactions with the
endothelium. In the ferric chloride model, by contrast, oxidative injury removes the endothelium and exposes circulating blood to the subendothelial matrix (26). Platelets bind to the subendothelium via glycoprotein Ib (GPIb), integrins, and glycoprotein VI (GPVI) and then become activated and aggregate via integrin receptors (27). We visualized thrombus formation in FeCl3-injured mesenteric venules using intravital microscopy and found that PAD4−/− mice formed occlusive thrombi with similar kinetics (Fig. 3G) and thrombus morphologies (Fig. 3H) as WT mice. Thus, we have no evidence that PAD4 or NETs play a significant role in hemostasis in healthy mice. For platelet plug formation in the injured veins, coagulation is likely driven by vessel wall tissue factor. Here the enhancement of coagulation and thrombosis by NETs may not be necessary.

Neutrophil PAD4 Is a Key Player in Pathological Thrombus Formation.

To assess whether the influence of PAD4 was of hematopoietic origin, we generated bone marrow (BM) chimeras by infusing WT or PAD4−/− BM cells into lethally irradiated WT recipients. Both groups of mice recovered body mass with similar kinetics (Fig. S2A). To test the chimerism, we stimulated leukocytes from whole blood after red blood cell (RBC) lysis with ionomycin and found that the PAD4−/− BM chimeras failed to hypercitrullinate arginine residues at histone H3 (Fig. S2B and C). Similar to the thrombus formation frequencies in WT and PAD4−/− mice, a majority (six of seven) of WT BM chimeras produced thrombi after 48 h of stenosis, whereas only one of the seven PAD4−/− BM chimeras formed a small thrombus (Fig. 4A and B). The H3Cit pattern within thrombi paralleled the results seen in WT and PAD4−/− mice, with extracellular H3Cit present in the thrombi of WT BM chimeras but H3Cit staining absent in the PAD4−/− BM chimera thrombus (Fig. S2D and E). WT BM chimeras became thrombocytopenic 48 h poststenosis, whereas PAD4−/− BM chimeras maintained normal platelet counts (Fig. 4C).

Fig. 2. PAD4−/− mice are protected from DVT, especially at 48 h after stenosis. (A and B) PAD4−/− mice were less likely to form thrombi at 6 h than WT mice. (C) Platelet count is decreased in WT compared with PAD4−/− mice after 6 h in the venous stenosis model. (D and E) At 48 h, only one small thrombus was present in a PAD4−/− mouse, whereas all but one of the WT mice had thrombi. (F) WT and PAD4−/− mice have similar platelet counts before stenosis (“Baseline”). WT mice are thrombocytopenic at 48 h, whereas platelet counts remain stable in PAD4−/− mice. (G and H) Composite image of a section from an entire thrombus collected at 48 h showing diffuse, extracellular H3Cit (green) staining in a WT mouse (G), which is completely absent in the sole thrombus from a PAD4−/− mouse (H). Hoescht staining of DNA is shown in blue and shows extensive presence of leukocytes in both thrombi. (Scale bars, 100 μm.) (I) H&E staining of 48 h thrombi shows less dense nuclear staining and diffuse, likely extracellular, DNA (arrows) in WT thrombi compared with dense nuclei found in the PAD4−/− thrombus. (Scale bar, 100 μm.) (J) Thrombi collected from WT mice or the PAD4−/− mouse that formed a thrombus 48 h after IVC stenosis showed an abundance of neutrophils by Ly6G immunostaining (red). (Scale bar, 20 μm.) DNA is shown in blue. *P < 0.05, **P < 0.01, ***P < 0.001.
As PAD4 is expressed by other leukocytes in addition to neutrophils (28, 29) that may enter the thrombus, we investigated whether the infusion of WT neutrophils could rescue thrombosis in PAD4−/− mice at 48 h. Infusing 4–5 × 10^6 isolated WT BM neutrophils into PAD4−/− mice restored thrombus generation (Fig. 4D and E). Although a significant lowering of platelet count was not achieved by neutrophil infusion (Fig. 4F), mice that had formed a thrombus in either WT and PAD4−/− recipients did, however, have a significant decrease in platelet count compared with baseline levels (Fig. S3). Infused neutrophils were of >95% purity as assessed by Wright–Giemsa staining (Fig. S4C). The thrombi from neutrophil-infused PAD4−/− mice contained large numbers of Ly6G+ neutrophils, whereas F4/80+ monocytes/macrophages were rarely seen within all thrombi examined (Fig. S4A, B, and D). Thrombi were analyzed by immunofluorescence, and extracellular H3Cit staining was present within both WT and PAD4−/− recipient thrombi after neutrophil infusion (Fig. 4G and H), indicating that the WT neutrophils were incorporated and produced NETs in the thrombus scaffold in PAD4−/− recipient mice. Therefore, PAD4 within neutrophils is important for formation of stable thrombi in large veins, as the defect in thrombosis in PAD4−/− mice can be rescued by supplementation with WT neutrophils.

**Discussion**

For more than 150 y, the conditions of hypercoagulable state, hypoxia due to flow disturbance such as in valves or stasis, and vascular activation were recognized as the main players in DVT (30). Only recently has the active contribution of blood cells to DVT begun to be appreciated. First, the complex role of inflammatory cells was recognized in that they promote thrombosis, for example, through production of tissue factor, and also contribute to thrombus resolution (31). Second, the important role of platelets in DVT has been described in animal models and human trials (9, 32, 33). Venous pathological thrombi were thought to be formed by fibrin and trapped RBCs, and the main treatment in humans is still anticoagulation. It is now clear that pathological thrombus formation and stabilization are much more complex, with many platelets and leukocytes present, and even the RBCs may not be trapped but rather actively recruited to the thrombus by NETs (6). Thus, new treatment options are being revealed.

Here we reveal a different facet of DVT—the importance of neutrophil posttranslational modification and chromatin decondensation, factors dramatically influencing outcome of stenosis-induced DVT in mice. The enzyme PAD4 allows for dissociation of heterochromatin protein 1-β (19), affects linker histone H1 (16), and reduces the positive charge of histones by arginine citrullination, thus allowing nucleosomes to unravel and chromatin to be expelled from the swollen nuclei. Absence of NETosis resulted in the formation of fewer thrombi in PAD4−/− mice early after the flow disturbance/hypoxia onset with almost no thrombi by 48 h. Here we describe DVT as a noninfectious model in which the PAD4−/− mice are affected, and, surprisingly, to date this is the strongest phenotype described in these mice (18, 34, 35).

We have demonstrated that PAD4 deficiency does not perturb initial vessel wall activation and platelet–leukocyte adhesion. Also, platelets from PAD4 knockout mice are fully functional and able to produce platelet plugs like WT platelets. The low incidence of DVT in PAD4−/− mice is largely caused by a lack of PAD4 in neutrophils as WT neutrophils could rescue the DVT...
Platelet counts in BM chimeras 48 h after DVT surgery. WT +PAD4−/− lethally irradiated WT recipients and later subjected to 48 h of IVC stenosis. (A) Percentage of BM chimeras that produced thrombi. (B) Length of thrombi. (C) Platelet counts in BM chimeras 48 h after DVT surgery. WT→WT BM chimeras became thrombocytopenic, whereas PAD4+/−→WT BM chimeras maintained normal platelet counts. (D–H) WT or PAD4−/− mice received an i.v. infusion of 4×10^6 WT neutrophils immediately before DVT surgery and 2–2.5×10^6 neutrophils at 24 h. (D and E) A majority (80%) of both WT and PAD4−/− mice formed thrombi after the infusion of WT neutrophils. (F) Similar platelet counts were measured 48 h after IVC ligation in WT and PAD4−/− mice infused with WT neutrophils. (G and H) Composite images of thrombi collected from mice infused with WT neutrophils. H3Cit-positive nuclei and extracellular staining was detected in WT (G) and PAD4−/− (H) thrombi, indicating that the infused WT neutrophils integrated into the PAD4−/− thrombus. H3Cit, green; DNA, blue. Representative of n=4. (Scale bar, 100 μm.) *P < 0.05.

Neutrophil Isolation for in Vitro NET Assays. Peripheral blood was collected via the retroorbital venous plexus, and neutrophils were isolated from 6- to 10-wk-old male or female WT, PAD4−/−, or PAD4−/− mice as previously described (11). Cells were routinely assessed to be >90% pure by Wright–Giemsa stain. Neutrophils in RPMI/Hepes were allowed to adhere at 37 °C for 10 min at 4 °C. Samples were incubated for 1 h at room temperature in antibody dilution buffer containing either biotinylated rat anti-Ly6G (clone RB6-8C5; eBioscience), rat anti-Ly6G (clone RB6-8C5; Biolegend), or rat anti-F4/80 (1:250, ab16911; Abcam). After several washes, samples were incubated for 2 h at room temperature in antibody dilution buffer containing Alexa Fluor-conjugated secondary antibodies in 0.3% BSA, 0.1% Tween-20, and either rat anti-Ly6G (0.5 μg/mL, clone RB6-8C5; eBioscience), rat anti-Ly6G (0.5 μg/mL, clone 1A8; Biolegend), or rat anti-F4/80 (1:250, ab16911; Abcam). After several washes, samples were incubated for 2 h at room temperature in antibody dilution buffer containing Alexa Fluor-conjugated secondary antibodies in 0.3% BSA in PBS: goat anti-rat IgG (Alexa555, 2 μg/mL), donkey anti-rabbit IgG (Alexa488, 1.5 μg/mL), or donkey anti-sheep IgG (Alexa688, 2 μg/mL; Invitrogen). DNA was counterstained with 1 μg/mL Hoechst 33342 and slides were cover-slipped with Fluoromount gel (Electron Microscopy Sciences). Fluorescent images were acquired using an Axiocam 200i widefield fluorescence microscope (Zeiss) in conjunction with an Axiocam MMr monochromatic CCD camera (Zeiss) and analyzed with Zeiss Axiovision software. All channels were acquired in grayscale and pseudocolored using Zeiss Axiovision or ImageJ software (National Institutes of Health). Exposure times are identical between

Materials and Methods

Mice. Experimental procedures in this study were reviewed and approved by the Institutional Animal Care and Use Committee of Boston Children’s Hospital (protocol nos. 11–03-1919, 11–04-1848, 11–03-1941). WT mice were purchased from Jackson Laboratory. PAD4−/− mice have been recently backcrossed to C57BL/6J for seven or more generations.

Venous Stenosis Model. Venous stenosis experiments were performed as previously described (8). Please see Supporting Information for more detailed methods.

Immunostaining and Fluorescence Microscopy. Fixed cells or tissue sections were washed with PBS and permeabilized (0.1% Triton X-100, 0.1% sodium citrate) for 10 min at 4 °C. Samples were blocked with 3% (v/v) BSA for 90 min at 37 °C, rinsed, and then incubated overnight at 4 °C or for 1 h at 37 °C in antibody dilution buffer containing 0.3% BSA, 0.1% Tween-20, and either rabbit antihistone H3 (citrulline 2, 8, 17) (0.3 μg/mL, ab5103; Abcam), rat anti-Gr1 (0.5 μg/mL, clone RB6-8C5; eBioscience), rat anti-Ly6G (0.5 μg/mL, clone 1A8; Biolegend), or rat anti-F4/80 (1:250, ab16911; Abcam). After several washes, samples were incubated for 2 h at room temperature in antibody dilution buffer containing Alexa Fluor-conjugated secondary antibodies in 0.3% BSA in PBS: goat anti-rat IgG (Alexa555, 2 μg/mL), donkey anti-rabbit IgG (Alexa488, 1.5 μg/mL), or donkey anti-sheep IgG (Alexa688, 2 μg/mL; Invitrogen). DNA was counterstained with 1 μg/mL Hoechst 33342 and slides were cover-slipped with Fluoromount gel (Electron Microscopy Sciences). Fluorescent images were acquired using an Axiocam 200i widefield fluorescence microscope (Zeiss) in conjunction with an Axiocam MMr monochromatic CCD camera (Zeiss) and analyzed with Zeiss Axiovision software. All channels were acquired in grayscale and pseudocolored using Zeiss Axiovision or ImageJ software (National Institutes of Health). Exposure times are identical between
WT and PAD4−/− thrombi or neutrophils. Composite images were generated with the MosaicJ plugin (36) for ImageJ software.

**Platelet Counts.** Whole blood was collected via the retroorbital sinus into EDTA-coated capillary tubes. Twenty-five μL of blood was analyzed by a Hemavet 950FS (Drew Scientific) for complete blood counts.

**Platelet Aggregation.** Murine blood was collected in 10% (vol/vol) sodium citrate (3.2% [wt/vol]) and centrifuged at 800 × g for 10 min. Platelet-rich plasma was collected, and supplemented with prostacyclin in Tyrode’s buffer. Platelets were washed and resuspended in Tyrode’s buffer and incubated at 37 °C. Platelets (2.5 × 10^5/L) were analyzed using a Chronolog Platelet aggregometer. Thrombin (1 or 0.1 U/mL) in the presence of calcium chloride was used as an agonist for aggregation, and samples were analyzed for at least 10 min.

**Tail Bleeding Time.** Six- to 8-week-old male or female C57BL/6J or PAD4−/− mice were anesthetized using 2.5% (vol/vol) tribromoethanol (300 mg/kg) administered intraperitoneally. A 2-mm segment of the tail tip was transected using a single-edge razor blade, and the tail was immediately submerged in 37 °C PBS. Tail bleeding was monitored for 15 min, and the time to stop was recorded when bleeding had stopped for more than 30 s. All mice were euthanized immediately at the end of the observation period.

**FeCl3-Induced Mesenteric Venous Thrombosis.** Three- to 4-week-old C57BL/6J or PAD4−/− male mice were anesthetized using 2.5% (vol/vol) tribromoethanol (300 mg/kg) administered i.p., and the level of anesthesia was verified by lack of response to footpad squeeze. Rhodamine 6G was given via retroorbital injection to fluorescently label platelets and leukocytes. A midline incision was performed, and the intestines externalized to reveal mesenteric vessels. A single venule of between 200 to 300 μm in diameter was visualized for each mouse (WT mice, 222.2 ± 20.6 μm; PAD4−/− mice, 239.0 ± 44.6 μm). A 1- × 3-mm piece of filter paper was soaked in 10% ferric chloride, gently placed on the vessel for 5 min, and then removed. Occlusion time was noted for each mouse. Vessels were considered occluded when blood flow had stopped for at least 10 s.

**Bone Marrow Chimeras Preparation.** Four-week-old C57BL/6J males were lethally irradiated with 1,100 rad and immediately given an i.v. injection of at least 1 × 10^9 bone marrow cells from either WT or PAD4−/− mice. See Supporting Information for detailed methods.

**Neutrophil Infusion.** Neutrophil adoptive transfer studies were performed as previously described (37). BM neutrophils were isolated from 6- to 8-week-old C57BL/6J WT mice using a Percoll (GE Healthcare Life Sciences) gradient and rapid hypotonic lysis. Bone marrow was flushed out of tubis and femurs using a 30-G needle into cold RPMI. Cells were pelleted, resuspended in PBS, and layered on a Percoll gradient containing 78%/69%/52% (vol/vol) Percoll. Cells at the 78%/69% interface were collected and subjected to hypotonic lysis. Cells were counted using a hemocytometer and resuspended in RPMI. Purity was assessed by Wright–Giemsa staining, which was determined to be >95% neutrophils. Cells (4–5 × 10^9) were injected i.v. through the retroorbital sinus immediately before DVT surgery. A second i.v. neutrophil infusion of 2–2.5 × 10^9 neutrophils was given at 24 h. Mice were euthanized at 48 h, and any resulting thrombi were harvested for analysis.

**Statistics.** Data are presented as means ± SEM unless otherwise noted and were analyzed using a two-sided Student t test or Mann–Whitney U test. Thrombus frequencies were analyzed using χ^2 tests of contingency tables. All analyses were performed using GraphPad Prism software (Version 5.0). Results were considered significant at P < 0.05 (**P < 0.01, ***P < 0.001).

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