Novel evidence that crosstalk between the complement, coagulation and fibrinolysis proteolytic cascades is involved in mobilization of hematopoietic stem/progenitor cells (HSPCs)

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The role of blood proteinases in the mobilization of hematopoietic stem/progenitor cells (HSPCs) is still not well understood. As previously reported, activation of the complement cascade (ComC) and cleavage of C5 by C5 convertase are enabling events in the release of C5a that plays a crucial role in the egress of HSPCs from bone marrow (BM) into peripheral blood (PB) and explains why C5-deficient mice are poor mobilizers. Here we provide evidence that during granulocyte colony-stimulating factor- and AMD3100-induced mobilization, not only the ComC but also two other evolutionarily ancient proteolytic enzyme cascades, the coagulation cascade (CoaC) and the fibrinolytic cascade (FibC), become activated. Activation of all three cascades was measured by generation of C5a, decrease in prothrombin time and activated partial thromboplastin time as well as an increase in the concentrations of plasmin/antiplasmin and thrombin/antithrombin. More importantly, the CoaC and FibC, by generating thrombin and plasmin, respectively, provide C5 convertase activity, explaining why mobilization of HSPCs in C3-deficient mice, which do not generate ComC-generated C5a convertase, is not impaired. Our observations shed more light on how the CoaC and FibC modulate stem cell mobilization and may lead to the development of more efficient mobilization strategies in poor mobilizers. Furthermore, as it is known that all these cascades are activated in all the situations in which HSPCs are mobilized from BM into PB (for example, infections, tissue/organ damage or strenuous exercise) and show a circadian rhythm of activation, they must be involved in both stress-induced and circadian changes in HSPC trafficking in PB.

INTRODUCTION

Hematopoietic stem/progenitor cells (HSPCs) express the chemokine receptor CXCR4 and the very late antigen-4 receptor (VLA-4). They also known as CD106) that are expressed by cells in the BM microenvironment (for example, osteoblasts and fibroblasts). This is known as α4β1 integrin) on their surface and are retained in bone marrow (BM) niches by interaction of these receptors with their respective ligands, α4-chemokine stromal-derived growth factor-1 (SDF-1) and vascular adhesion molecule-1 (VCAM-1, also known as CD106) that are expressed by cells in the BM microenvironment (for example, osteoblasts and fibroblasts). This helps them to circulate under steady-state conditions at detectable levels in the peripheral blood (PB), and their number increases in response to (1) systemic or local inflammation, (2) strenuous exercise, (3) stress, (4) tissue/organ injury and (5) pharmacological agents. Activation of the complement cascade (ComC), and mice deficient in a downstream component of ComC, complement protein 5 (C5), are very poor mobilizers. It has been explained by demonstration that the C5 cleavage fragment C5a, which is activated in BM sinusoids, is crucial for egress of granulocytes and monocytes from the BM and that these cells pave the way for HSPCs through the PB–BM barrier during mobilization. At the same time, C5a activates granulocytes and monocytes to release several proteolytic enzymes in the BM microenvironment that attenuate SDF-1–CXCR4 and VLA4–VCAM-1 retention signals in BM niches.

It is also known that activation of the ComC, similar to coagulation cascade (CoaC) and fibrinolytic cascade (FibC), is based on sequential activation of proteolytic proenzymes from the top to the bottom of the cascade. Therefore, the lack of upstream C3 should theoretically affect generation of ComC-generated C5 convertase, a proteolytic enzyme activating C5. Surprisingly, C3−/− mice are easy mobilizers that suggests that other proteolytic enzymes in blood plasma substitute for ComC-generated C5 convertase. To explain how C5 can be activated during the mobilization process even when C3 is missing, we hypothesized that other proteases that are products of the activated CoaC and FibC compensate for the lack of proteolytic activity of ComC-derived C5 convertase. In support of this hypothesis, it has been demonstrated that both CoaC and FibC have vigorous crosstalk with ComC during some innate immunity-mediated responses. It has also been demonstrated that thrombin (Dr T Lapidot, personal communication) or plasmin administration may enhance mobilization of HSPCs, suggesting a role for these enzymes in the mobilization process. To support this further, G-CSF-induced mobilization of HSPCs was facilitated in plasminogen activator inhibitor-1- and plasmin inhibitor-α2 antiplasmin-deficient mice. Moreover, both membrane-anchored plasminogen activator, urokinase receptor and cleaved form of soluble urokinase receptor have been implicated in HSPCs mobilization.

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In our experiments, mobilization was evaluated in C3-deficient mice (C3−/−) and normal wild-type (WT) littermates mobilized by granulocyte colony-stimulating factor (G-CSF) or the CXCR4 receptor antagonist AMD3100 in the presence or absence of refludan (a direct inhibitor of thrombin) and tranexamic acid (an inhibitor of plasminogen activation). In parallel, we measured the activation of all three cascades by detecting the level of CSa in PB and measuring prothrombin time (PT) and activated partial thromboplastin time (APTPT) as well as the concentrations of thrombin/antithrombin and plasmin/antiplasmin complexes.

The data presented in this work demonstrate for the first time the existence of vigorous crosstalk between all three evolutionarily ancient proteolytic enzyme cascades, the ComC, CoaC, and FibC, in the process of mobilizing HSPCs. We observed that G-CSF-induced mobilization of HSPCs was significantly reduced in normally easy-mobilizing C3−/− mice when the mice were treated with refludan (a CoaC inhibitor) or tranexamic acid (an FibC inhibitor) and that this reduction correlated with significant inhibition of C5 activation/cleavage. Significantly, we also observed that inhibitors of the CoaC and FibC had a negative effect on mobilization of HSPCs in normal WT animals. Our observations of crosstalk between the ComC, CoaC, and FibC better explain the effect of thrombin and plasmin on stem cell mobilization and may lead to the development of more efficient mobilization strategies in poor mobilizers and mobilized patients who are treated with drugs that interfere with CoaC/FibC.

MATERIALS AND METHODS

Animals
Mobilization experiments were performed on 6–8-week-old C57Bl/6 C3−/− and WT C57Bl/6 C3+/+ age- and sex-matched littermates (breeding colony purchased from The Jackson Laboratory, Bar Harbor, ME, USA). C57Bl/6 and C3−/− mice were identified by screening for C3 protein levels (using radial immunodiffusion), and before the experiment, PCR was used to confirm the genotype of the C3−/− mice. Animal studies were approved by the Animal Care and Use Committee of the University of Louisville (Louisville, KY, USA).

Mobilization studies
Mice were mobilized by G-CSF (6 days, 250 μg/kg, subcutaneously) or AMD3100 (5 mg/kg, subcutaneous) in the absence or presence of refludan, a direct inhibitor of thrombin (5 mg/kg/day, divided in two doses). Bacteri, Whippy, NJ, USA) or tranexamic acid, an inhibitor of plasminogen activation (20 mg/mice/day; Sigma-Aldrich, MO, USA). Control mice were injected with vehicle, refludan or tranexamic acid only. At 6 h after the last G-CSF injection or 1 h after AMD3100 injection, PB was obtained from the vena cava (with a 25-gauge needle and 1 ml syringe containing 250 U heparin).

Peripheral blood parameter counts
To obtain leukocyte counts, blood samples were collected from the retro-orbital plexus of the mice into microvette EDTA-coated tubes (Sarstedt Inc., Newton, NC, USA) and run within 3 h of collection on a HemaVet 950 hematology analyzer (Drew Scientific Inc., Oxford, CT, USA; http://www.drew-scientific.com).

Fluorescence-activated cell sorting analysis of SKL cells
The following monoclonal antibodies were employed to stain Sca-1 (clone RA3–6B2), anti-mouse TCRαβ–PE (clone H57–597), anti-mouse CD45R/B220–PE (clone E13–161.7), streptavidin–phycoerythrin (PE)–Cy5-conjugated anti-mouse c-Kit (clone 2B8) and lineage markers anti-mouse CD45R/B220–PE (clone 53-6.7), anti-mouse CD3ε–PE (clone 145-2C11), anti-mouse CD8α–PE (clone 53-6.7), anti-mouse CD4–PE (clone 145-2C11), anti-mouse CD11b–PE (clone M1/70), anti-mouse Ter119–PE (clone TER-119) and anti-mouse Gr-1–PE (clone RB6–8C5), as described. All monoclonal antibodies were added at saturating concentrations, and the cells were then incubated for 30 min on ice, washed twice, resuspended in RPMI-1640 + 2% fetal bovine serum, and analyzed with an LSR II flow cytometer (BD, San Diego, CA, USA).

RESULTS

Enumeration of the number of colony-forming unit-granulocyte/macrophage (CFU-GM) mobilized into PB
After PB red blood cell lysis (BD Pharm Lyse Buffer, San Jose, CA, USA), nucleated cells were washed, counted and 1 × 10^6 cells were resuspended in 10% culture medium with 90% human methylcellulose base media supplemented with 25 ng/ml recombinant murine GM-CSF and 10 ng/ml recombinant murine IL-3 (PeproTech, Rocky Hill, NJ, USA). After 1 week of culture, the numbers of CFU-GM colonies were scored using an inverted microscope (Olympus, Center Valley, PA, USA).

Evaluation of HSPC mobilization
For evaluation of circulating CFU-GM and Sca-1+/c-Kit+/Lin− (SKL) cells, the following formulas were used: (number of white blood cells (WBCs) × number of CFU-GM colonies)/number of WBCs plated = number of CFU-GM per μl of PB; and (number of WBCs × number of SKL cells)/number of gated WBCs = number of SKL cells per μl of PB.

Plasma concentration of C5b-C9 (MAC complex)
The concentration of C5b-C9 was measured by employing the commercially available, highly sensitive enzyme-linked immunosorbent assay (ELISA) kit K-ASSAY (Kamiya Biomedical Company, Seattle, WA, USA), according to the manufacturer’s protocol. For analysis, PB from C3−/− mice was collected on day 6 of G-CSF-induced mobilization by retro-orbital plexus bleeding into cold microvette EDTA-coated tubes (Sarstedt Inc.). Subsequently, blood was centrifuged at 2000 g for 20 min in 4°C to obtain plasma.

Activation of coagulation cascade
PT and APTT were evaluated by employing coagulometry. For measurement of CoaC components, PB from C57Bl/6 mice was collected on day 6 after G-CSF- or 1 h AMD3100-induced mobilization into tubes with 3.2% sodium citrate (ratio: 1:9). Subsequently, blood was centrifuged for 10 min at 700 g at 4°C to obtain plasma. PT and APTT were measured within 4 h of blood collection. For PT evaluation, THROMBOPLASTIN PT-S (Cormay, Lomianki, Poland) was warmed to 37°C, and 100 μl was added to 0.05 ml of serum. The time of appearance of a clot was measured using a coagulometer. For APTT evaluation, to 0.05 ml of serum, 0.05 ml of APTT-P-REAGENT (Cormay) was added, and after 5 min of incubation (37°C), calcium chloride (warmed to 37°C) was added. The time of appearance of a clot was evaluated using a coagulometer. Thrombin/antithrombin complexes were measured by employing ELISA assay, according to the manufacturer’s protocols (USCN Life Science, Wuhan, China).

Activation of fibrinolysis cascade
Plasmin/antiplasmin complexes were measured by employing ELISA assay, according to the manufacturer’s protocols (USCN Life Science). Plasma was collected in the same way, as reported above.

Statistical analysis
Arithmetic means and s.d. were calculated using Excel. Statistical significance was defined as P < 0.05. Data were analyzed using Student’s t-test for unpaired samples (Excel), and ANOVA was performed using STATISTICA 10.0 (StatSoft, Krakow, Poland).

G-CSF and AMD3100 mobilization is impaired in C3−/− mice in the presence of refludan and/or tranexamic acid
We have already reported that inhibition of the CoaC by refludan impairs G-CSF-induced mobilization of HSPCs in C3−/− mice. In this report, we show that, in addition to inhibiting the CoaC, inhibition of the FibC has a similar effect. Moreover, inhibition of both these proteolytic cascades at the same time even more significantly affects the numbers of mobilized WBCs, SKL and CFU-GM circulating in PB (Figure 1a). We report here, also for the first time, that refludan and tranexamic acid have similar negative effects on AMD3100-induced mobilization of HSPCs in C3−/− mice (Figure 1b). These data support the hypothesis that both thrombin and plasmin in fact possess C5 convertase-like activity.

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Nevertheless, as is shown in Figure 1, mobilization was not completely inhibited after simultaneous inhibition of both CoaC and FibC that implies that some residual activity of both thrombin and plasmin may be present, or some other plasma proteases partially compensate for thrombin and plasmin activity. It also shows that, when employed alone, neither refludan nor tranexamic acid affects the pool of circulating HSPCs.

Administration of refludan and/or tranexamic acid also impairs stem cell mobilization in normal WT littermates.

In this study, we also present data showing that inhibition of both CoaC and FibC has a negative effect on mobilization of HSPCs after either G-CSF- (Figure 2a) or AMD3100-induced (Figure 2b) mobilization. This effect was partially visible in G-CSF-induced mobilization when refludan and tranexamic acid were employed together. However, in AMD3100-induced mobilization, the addition of tranexamic acid to refludan did not significantly increase mobilization. This effect was particularly visible in G-CSF-induced mobilization. This implies that some residual activity of both thrombin and plasmin may be present, or some other plasma proteases partially compensate for thrombin and plasmin activity.

As described in the past, we observed that in response to G-CSF administration, C3$$^{-/-}$$ mice mobilize greater numbers of HSPCs into PB (Figure 1a) than WT littermates (Figure 2a) that suggests that C3 cleavage fragments promote CXCR4-dependent retention of HSPCs in BM. Interestingly, when we employed the CXCR4 antagonist AMD3100 in our current study, this difference in mobilization was not apparent (Figure 1b vs Figure 2b). Again, when employed alone, neither refludan nor tranexamic acid affected the pool of circulating HSPCs.

All three evolutionarily ancient proteolytic cascades are activated during G-CSF- and AMD3100-induced mobilization. In support of our in vivo mobilization data, in the presence or absence of CoaC and FibC inhibitors, we directly measured activation of both cascades in the PB of mice mobilized by G-CSF or AMD3100.

Figure 3 demonstrates that administration of G-CSF (Figure 3A) or AMD3100 (Figure 3B) significantly decreases both PT and APTT and AMD3100 significantly increases thrombin/antithrombin that indicates activation of the CoaC. G-CSF administration also increased the concentration of thrombin/antithrombin complexes in PB, although this increase was not significant. At the same time, as shown in Figure 3A, AMD3100 significantly activated the FibC, as seen by an increase in plasmin/antiplasmin concentration. Finally, activation of the ComC during G-CSF- and AMD-3100-induced mobilization, as previously reported, was inhibited after administration of refludan and/or tranexamic acid (Figure 3C).

DISCUSSION

The mechanisms that regulate mobilization of HSPCs are still not well understood, and a relatively large number of patients,
particularly those pretreated by chemotherapy, are poor mobilizers in response to G-CSF.\textsuperscript{7,18-20}

It has been reported that administration of G-CSF or AMD3100 induces a highly proteolytic microenvironment in BM because of several proteolytic enzymes released by activated granulocytes and monocytes.\textsuperscript{21,22} Thus, it is widely accepted that the first step in the release of HSPCs from their niches in BM into PB is related to attenuation of the interaction between the major homing/retention receptors (CXCR4 and VLA-4) expressed on the surface of HSPCs with their corresponding ligands (SDF-1 and VCAM-1) in BM niches.\textsuperscript{1-5} This could be achieved by the action of the above-mentioned proteolytic enzymes or blockade with specific receptor-blocking molecules (for example, AMD3100). However, the role of BM-expressed proteolytic enzymes, such as neutrophil elastase, cathepsin-G or metalloproteinases,\textsuperscript{15} has been challenged by findings that knockout of several of these enzymes has only a negligible effect on the mobilization process,\textsuperscript{17} and this also suggests the existence of compensatory mechanisms involving other proteolytic enzymes. We hypothesize that proteases generated in the activation of CoaC and FibC may play a role here. In fact, it has been suggested that CoaC and FibC play a role in the release of HSPCs from BM,\textsuperscript{13-15} however, the molecular mechanisms behind this phenomenon are still not fully understood.

It is well known that three evolutionarily ancient proteolytic enzyme cascades, ComC, CoaC and FibC, are activated in PB during several events in which HSPCs are released from the BM into PB (for example, inflammation or strenuous exercise). All of these cascades also show circadian rhythms in activation because of a drop in blood pH during deep sleep at night and are most likely responsible for circadian changes in HSPC trafficking in PB.\textsuperscript{23,24} Therefore, in addition to circadian changes in the tonus of the vegetative nervous system, circadian activation of the ComC, CoaC and FibC is most likely also involved in the circadian release of HSPCs from BM into PB.\textsuperscript{6} What is also important, the three cascades show a vigorous crosstalk because of mutual cross-activation by proteolytic enzymes released, for example, during inflammation.\textsuperscript{10}

We previously reported on the importance of the ComC in G-CSF- and AMD3100-induced mobilization.\textsuperscript{4} We have also shown that mice defective in C3, which is a proximal component of the ComC, are easy mobilizers.\textsuperscript{12} We have explained this phenomenon by a lack of C3 cleavage and C3a formation that increases because of a drop in blood pH during deep sleep at night and are most likely responsible for circadian changes in HSPC trafficking in PB.\textsuperscript{23,24} Therefore, in addition to circadian changes in the tonus of the vegetative nervous system, circadian activation of the ComC, CoaC and FibC is most likely also involved in the circadian release of HSPCs from BM into PB.\textsuperscript{6} What is also important, the three cascades show a vigorous crosstalk because of mutual cross-activation by proteolytic enzymes released, for example, during inflammation.\textsuperscript{10}

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potent chemoattractant for granulocytes and monocytes, the first cells to egress from the BM and thus permeabilize the BM–PB barrier, facilitating subsequent egress of HSPCs that express proteolytic enzymes at an ~100-fold lower level.\(^{4,9}\) In addition, another C5 cleavage fragment, C5b, is involved in formation of \( \text{C5b-C9 (membrane attack complex)} \) that may increase the PB level of sphingosine-1-phosphate,\(^{12} \) a crucial direct chemoattractant in PB directing egress of HSPCs. Thus, the failure of C5-deficient mice to generate both C5a and membrane attack complex explains why these animals are poor mobilizers.\(^{9}\)

Taking into consideration that C3 is necessary for generation of C5 convertase, an enzyme that cleaves C5 during activation of CoaC, one would expect that C3-deficient mice, similar to C5-deficient animals, would be poor mobilizers. However, as mentioned above, this is not the case. To address why C3\(^{-/-}\) mice, despite with a defect in generation of C5 convertase, are easy mobilizers, we considered the involvement of alternative proteolytic pathways that provide C5 convertase-like activity and thus cleave/activate C5 in C3\(^{-/-}\) mice. Therefore we focused on the CoaC and FibC, knowing that both thrombin and plasminogen may possess C5 convertase-like activity, as demonstrated in some experimental settings.\(^{10}\) In our previous report we showed that inhibition of CoaC by defibraned negatively affects G-CSF-induced mobilization in C3\(^{-/-}\) animals, suggesting the involvement of the CoaC in this process.\(^{16}\) In further support of this interpretation, as mentioned above thombin and plasmin have been shown to mobilize HSPCs when administered \( \text{in vivo}. \) To our knowledge, we have demonstrated for the first time that administration of both G-CSF and AMD3100 activate CoaC, ComC and FibC at the same time and shed more light on the role of CoaC and FibC in the mobilization process. Specifically, based on the observation that C5 cleavage is crucial for egress of HSPCs from BM,\(^{9}\) we show here that both thrombin, as a product of the activated CoaC, and plasmin, as a product of the activated FibC, provide C5 convertase-like activity to facilitate C5 cleavage as a crucial step in HSPC mobilization (Figure 4).

Thus, these results confirm that C5 can be activated/cleaved in a thrombin- and plasmin-dependent manner\(^{10}\) in the absence of classical C3-dependent, CoaC-generated C5 convertase—and thus both thrombin and plasmin may provide C5-convertase-like activity during mobilization of HSPCs. We employed C3\(^{-/-}\) animals that are easy mobilizers as a model of this effect, although our data also support thrombin and plasmin providing C5-convertase-like activity in G-CSF- and AMD3100-mobilized WT animals. However, as inhibition of both proteases did not completely inhibit egress of HSPCs from BM, one has to consider the involvement of other components of both CoaC and FibC in providing C5-convertase-like activity. In support of this conclusion, C5 can also be cleaved by several other proteases released from granulocytes and monocytes unrelated to the CoaC and ComC, such as cathepsin, matrix metalloproteinase (MMP)-2 and MMP-9.\(^{17}\)

Figure 3. Evidence for activation of CoaC, FibC and ComC during mobilization of HSPCs. (A and B) Activation of the CoaC in WT mice mobilized by G-CSF or AMD3100, as measured by decrease of prothrombin time (PT) (a), APTT (b; \( *P < 0.05, **P < 0.01 \)) and by the increase in serum level of thrombin/antithrombin (TAT) complexes (c; \( *P < 0.005 \)). (d) Activation of the FibC in WT mice mobilized by G-CSF or AMD3100 as measured by the increase in serum level of and plasmin/antiplasmin (PAP) complexes. \( *P < 0.005 \). These are combined results from two independent experiments \( (n = 10 \text{ mice/group}) \).

(C and D) Activation of ComC as measured by increase in concentration of C5a cleavage fragments in PB (control values in nonmobilized mice were assumed to be 1.0). These are combined results from two independent experiments \( (n = 6 \text{ mice/group}) \). \( *P < 0.005 \)
In conclusion, the data presented in this work demonstrate, for the first time, the existence of crosstalk between three evolutionarily ancient proteolytic cascades, ComC, CoaC and FibC, in the mobilization of HSPCs by providing C5 convertase activity. Further study is needed to see whether proteolytic enzymes from those cascades also penetrate the blood–BM barrier and affect CXCR4–CXCL12 interaction in the mobilization of the patients who are treated with drugs that potentially interfere with CoaC/FibC.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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