Innate Immunity and Mobilization of Hematopoietic Stem Cells

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Published online: 10 July 2017 © The Author(s) 2017. This article is an open access publication

Abstract

Purpose of Review Several mechanisms have been postulated to orchestrate mobilization of hematopoietic stem/progenitor cells (HSPCs), and still more work is needed to better understand this process and to gain better mechanistic insight.

Recent Findings Evidence accumulated that mobilization of HSPCs is a part of innate immunity response to tissue organ injury, stress, and infection. This evolutionary ancient process is orchestrated by granulocytes and monocytes that trigger activation of complement cascade and the coagulation cascade.

Summary We will present data from our laboratory that initiation of complement cascade activation and subsequently activation of the coagulation cascade during mobilization process are dependent on mannan-binding lectin (MBL). The mannan-binding pathway activates MBL-associated serine proteases (MASP-1 and MASP-2) that cleave the third complement component C3 and prothrombin. Cleavage of C3 leads to formation of classical C5 convertase and cleavage of prothrombin generates thrombin, which has \( \text{C5-like convertase} \) activity. Finally, both C5 convertase and thrombin cleave the fifth complement component C5, and activate distal part of the complement cascade that is crucial for egress of HSCPs from bone marrow niches into peripheral blood.

Keywords Complement cascade · Hematopoietic stem cells · Stem cell mobilization · HO-1 · iNOS

Introduction

The alpha-chemokine stromal derived factor-1 (SDF-1)–C-X-C chemokine receptor type 4 (CXCR4) receptor and vascular adhesion molecule-1 (VCAM-1)–very late antigen-4 (VLA-4) integrin receptor axes play important roles in retention of hematopoietic stem/progenitor cells (HSPCs) in bone marrow (BM) niches [1, 2]. However, in steady-state conditions low numbers of HSPCs always circulate in peripheral blood (PB) and lymph, and undergo a circadian rhythm in their circulation, with the peak occurring early in the morning and the nadir at night [3, 4]. In addition to these circadian rhythms, the number of circulating HSPCs increases in PB in response to (i) systemic or local inflammation, (ii) strenuous exercise, (iii) hypoxia, and (iv) tissue/organ injuries [5, 6, 7, 8, 9].

Moreover, in clinical settings, administration of some agents may induce forced egress of HSPCs into PB and increase their number in PB up to 100-fold in a process known as “pharmacological mobilization.” Such drugs include cytokine granulocyte colony stimulating factor (G-CSF) and the small molecular CXCR4 antagonist AMD3100, also known as Plerixafor. The growth-related oncogene protein beta (Gro-β) [10, 11, 12, 13, 14], some cytostatics (e.g., cyclophosphamide), and non-steroidal anti-inflammatory drugs could be employed as mobilizing agents as well [15–18]. Pharmacological mobilization is exploited as a means to obtain HSPCs for hematopoietic reconstitution in the clinical setting for mobilized peripheral blood (mPB) grafts and our previous research demonstrated that the complement cascade (ComC), as part of innate immunity and the evolutionary...
ancient response to tissue/organ injury and stress, is an important modulator of stem cell trafficking [12••, 13].

Understanding the mechanisms that govern mobilization of HSPCs is crucial for optimizing hematopoietic stem cell transplants using mPB grafts. Unfortunately, in autologous transplant settings, ~10% of normal patients and ~25% of patients after chemotherapy do not respond efficiently to currently recommended mobilization protocols and are deemed poor mobilizers [19••, 20, 21]. Therefore, it is important to develop more efficient mobilization protocols in order to harvest the required number of CD34+ HSPCs for transplantation.

In this review, we present data on a role of elements of innate immunity including (i) granulocytes, (ii) monocytes, (iii) naturally occurring antibodies (NAbs), and (iv) activation of ComC and coagulation cascade (CoaC) in mobilization of HSPCs.

Chemoattractants for HSPCs and the Role of Accessory Cells Involved in Mobilization Process

Under steady-state conditions, HSPCs are actively retained in BM niches, and retention mechanisms counteract a continuous sphingosine-1-phosphate (S1P) chemoattractant gradient originating in blood plasma that can promote HSPCs egress into the blood, as recently demonstrated [22••, 23, 24]. The S1P concentration in PB is higher than in tissues as this active phosphosphingolipid is transported by erythrocytes, albumins, and high-density lipoproteins (HDL) [25].

Overall, there have been very few chemoattractants identified for HSPCs besides S1P, the abovementioned alpha-chemokine SDF-1 [26•], ceramide-1-phosphate (C1P) [27], and some extracellular nucleotides such as ATP [28, 29]. The responsiveness of HSPCs to SDF-1 is modulated by several molecules related to innate immunity [12••, 30•, 31•] and inflammation [32, 33]. Migration of HSPCs is facilitated by certain cationic antimicrobial peptides [34]; innate immunity mediators, including ComC-derived C3 cleavage fragments C3a and a desArgC3a; and cathelicidin LL-37, which enhance or prime the responsiveness of HSPCs to an SDF-1 gradient [32, 34, 35•, 36, 37, 38]. These findings demonstrate that the responsiveness of HSPCs to SDF-1 is modulated by several molecules related to innate immunity [12••, 30•, 31•] and inflammation [32, 33]. As reported by other investigators, the responsiveness of HSPCs to an SDF-1 gradient may also be enhanced by the presence of prostaglandin E2 (PGE2) [11, 39] and independent from SDF-1, gradient HSPCs may also follow in the homing process of calcium-sensing receptors and RIG-I-like receptors) [50, 55, 56, 57, 58••, 59].

PRRs are classified according to their ligand specificity, function, localization, and/or evolutionary relationships and may be divided into cell membrane PRRs (toll-like receptors, TLRs) and intracytoplasmic PRRs (NOD-like receptors and RIG-I-like receptors) [50, 55, 56, 57, 58••, 59]. Our most recent data demonstrated that TLRs play a negative role in mobilization, as they upregulate anti-inflammatory activity in HSPCs that as mentioned above inhibits cell migration and mobilization process [60].

We became interested in other PRRs that are secreted and activate the innate immune system or circulate free in PB to identify and bind to two classes of molecules: (i) pathogen-associated molecular patterns (PAMPs), which are expressed by microbial pathogens and what is relevant to this review, and (ii) damage-associated molecular patterns (DAMPs), which are associated with cell components and are released during cell activation, cell damage, or cell death [50, 56, 57, 58••].

Elements of Innate Immunity and Their Relevance to Stem Cell Mobilization

The innate (non-specific or inborn) immune system is an important part of host defense from infection by other organisms. The most important components of innate immunity are the ComC, NAbs, and Gr-1+ leucocytes. In addition, recent evidence indicates that ComC plays also an important pleiotropic role in embryonic development and tissue regeneration, which at least partly involve stem cell migration [49]. The ComC is composed of several plasma proteins synthesized in the liver and is an important damage-sensing mechanism activated by the (i) classical, (ii) mannann-binding lectin (MBL), and (iii) alternative pathways (Fig. 1) [50–52]. NAbs are produced without any previous infection, vaccination, or foreign-antigen exposure. In addition to B cell derived NAbs, monocytes and neutrophils are also involved in mobilization of HSPCs [53, 54].

An important role in innate immunity is also played by pattern-recognition receptors (PRRs) [55, 56, 57, 58••]. These proteins are expressed on the surface of cells of the innate immune system or circulate free in PB to identify and bind to two classes of molecules: (i) pathogen-associated molecular patterns (PAMPs), which are expressed by microbial pathogens and what is relevant to this review, and (ii) damage-associated molecular patterns (DAMPs), which are associated with cell components and are released during cell activation, cell damage, or cell death [50, 56, 57, 58••].
mannose-binding lectin (MBL), is a major PRR of the innate immune system that binds to a wide range of pathogens [51, 52, 61]. MBL also binds phospholipids and several DAMPs released from activated cells, such as high-mobility group box 1 (HMGB1), extracellular ATP, DNA, and hyaluronan fragments [55, 56, 59, 62]. Once bound to ligands, MBL recruits MBL-associated serine proteases (MASP-1 and MASP-2) [51, 62, 63] and initiates activation of the ComC. After cleavage of C3, the next step is generation of classical C5 convertase that cleaves C5 into C5a and iC5b. In parallel, MASPs also activate prothrombin that gives rise to thrombin which has C5B convertase-like activity. C5 cleavage fragments [64•] are crucial for egress of HSPCs from BM into PB by augmenting degranulation of granulocytes (release of proteolytic and lipolytic enzymes) and chemoattracting granulocytes and monocytes into PB. In addition, C5b-C9 (MAC) may release S1P from erythrocytes and platelets and thus enhances the S1P level in PB, which directs egress of HSPCs. Of note, the alternative pathway is continuously active at a very low level under steady-state conditions. Contribution of this pathway to mobilization requires further studies.

A Novel View of ComC-Related Pro-mobilizing Mechanisms—Turning Away from the Classical to Implicate the Mannan-Binding Pathway for ComC Cascade Activation

We are aware that, since mobilization of HSPCs is an evolutionarily ancient and important biological process, there may be redundant mechanisms that modulate this process. Nevertheless, elements of innate immunity, such as ComC activation, granulocytes, monocytes, and NAbs, play major roles.

When we initially discovered a requirement for ComC activation in HSPC mobilization, we assumed that the classical activation pathway of ComC would play a pivotal role [12••]. However, to our surprise, mice with mutations in components that initiate the classical pathway (e.g., C1q−/− mice) (Fig. 1) do not show impairment in mobilization of HSPCs [68]. Thus, the main question emerged, which ComC activation pathway triggers mobilization of HSPCs after administration of G-CSF or AMD3100? We recently demonstrated that it is likely to be the aforementioned MBL pathway that comes into play Fig. 1. Specifically, mice deficient in MBL and MASP-1 are poor mobilizers [66••]. Moreover, by contrast, mobilization is enhanced in mice HO-1- and iNOS-deficient animals [42••, 43••] indicating that both these enzymes are negative counterbalancing regulators of this process.

To provide a step-by-step mechanistic picture (Fig. 2), we propose that activation of Gr-1+ granulocytes and monocytes by mobilizing agents leads to the release of proteolytic enzymes [69••] and lipolytic enzymes [48] that disrupt the SDF-1–CXCR4 and VCAM-1–VLA-4 retention axes that keep HSPCs in their BM niches and to the secretion of reactive oxygen species (ROS) that expose auto-antigens known...
Neoepitopes in the BM microenvironment, which bind NAbs, mainly of the IgM class [37, 62]. It is known that NAbs that are produced as mentioned above without any previous infection, vaccination, other foreign antigen exposure, or passive immunization may preferentially recognize oxidation-specific epitopes exposed by ROS during inflammation [62, 66]. The binding of NAbs to neoepitopes creates a damage-associated molecular pattern (DAMP) complex that is recognized by MBL, being a major soluble pattern-recognition receptor (PRR) of the innate immune system that triggers activation of the ComC (Fig. 2). In addition, Gr-1+ cells also release other soluble DAMPs that are recognized by MBL including high-mobility group box 1 (HMGB1), extracellular ATP, DNA, and hyaluronan fragments [55, 56, 59, 62]. Our preliminary data show that all these molecules are potentiating mobilization in MBL-MASP-dependent manner.

In this context, it is important to more thoroughly explore the role of the ComC in modulating the function of Gr-1+ cells (granulocytes and monocytes) in BM, as they are the first cells that egress from BM into PB and pave the way for HSPCs to cross the BM–PB endothelial barrier [12++, 54, 70, 71**]. This occurs in response to increase in C5a level in PB due to ComC activation. C5a is a most potent activator and chemoattractant for Gr-1+ cells [12**]. In addition, it has been postulated that monocytes may secrete factors that inhibit SDF-1 expression in BM niches [70, 72].

Based on these considerations, we propose the following involvement of innate immunity in the mobilization of HSPCs and distinguish three main phases of this process: (i) initiation, (ii) amplification, and (iii) execution phase (Fig. 2). These three phases will be shortly discussed below.

**Initiation Phase** Mobilizing agents G-CSF or AMD3100 (which besides its blocking properties is also a partial agonist of CXCR4) [73] activate Gr-1+ neutrophils and monocytes and enhance secretion of reactive oxygen species (ROS) by these cells. In the BM microenvironment, ROS expose neoepitopes. Neoepitope–IgM complexes and DAMPs are recognized by MBL, which activates the ComC and CoaC in a MASP-dependent manner. Step 2 (amplification). Convertases (classical C5 and C5-like) generated in this step cleave C5 to release cleavage fragments crucial in egress of HSPCs from BM. This step is also modulated by a C3 auto-amplification loop, with a possible contribution from the alternative ComC activation pathway. Step 3 (execution). In this step, C5 cleavage fragments promote release of HSPCs from BM, and this process is negatively regulated by HO-1 and iNOS—shown as negative sign

![Fig. 2](image) Proposed three-step model for triggering mobilization of HSPCs. All the phases of mobilization process are depicted. Step 1 (initiation phase). Activation of Gr-1+ neutrophils and monocytes by mobilizing agent (e.g., G-CSF or AMD3100) induces secretion of ROS and DAMPs by these cells. In the BM microenvironment, ROS expose neoepitopes. Neoepitope–IgM complexes and DAMPs are recognized by MBL, which activates the ComC and CoaC in a MASP-dependent manner. Step 2 (amplification). Convertases (classical C5 and C5-like) generated in this step cleave C5 to release cleavage fragments crucial in egress of HSPCs from BM. This step is also modulated by a C3 auto-amplification loop, with a possible contribution from the alternative ComC activation pathway. Step 3 (execution). In this step, C5 cleavage fragments promote release of HSPCs from BM, and this process is negatively regulated by HO-1 and iNOS—shown as negative sign.
C5α and iC5b that are crucial in egress of HSPCs from BM. This step is also modulated by a C3 auto-amplification loop, with a possible contribution from the alternative ComC activation pathway. To support involvement of this pathway, MBL−/− and MASP-1−/− mice still mobilize HSPCs, however in much less effective manner. Overall, the alternative pathway is activated by foreign or the organism’s own damaged cells and is facilitated by the continuous spontaneous hydrolysis of C3. The alternative pathway of ComC activation also has an important function by providing an amplification loop enhancing C3 activation, independent of which ComC pathway was initially activated. This effect is mainly due to properdin, the only positive regulator in the complement system, which stabilizes C3 convertase. To confirm involvement of alternative ComC activation pathway, it would be important to perform mobilization studies in B factor- and properdin- and factor H-deficient mice, as these factors regulate the spontaneous C3 amplification loop. Impaired mobilization in these animals would support involvement of alternative pathway of ComC activation.

**Execution Step** In this step, C5 cleavage fragments promote release of HSPCs from BM, and this process is negatively regulated by HO-1 and iNOS. To support this later notion, activation of ComC-mediated mobilization must be strictly controlled, and our most recent research revealed that both HO-1 and iNOS are negative regulators of cell trafficking that attenuate mobilization process [41, 42••, 43••]. HO-1 is an inducible stress–response enzyme that not only catalyzes the degradation of heme (e.g., released from erythrocytes) but also has an important function in various states associated with cellular stress [74–76]. Inherited HO-1 deficiency in humans and in mice results in vulnerability to stressful injury and inflammation, which can be explained by hyperactivity of the ComC [77].

We demonstrated for the first time that HO-1 plays an important and heretofore unrecognized role in retention of HSPCs in BM niches by (i) negatively modulating activation of the mobilization-promoting ComC and (ii) attenuating the chemotaxis of HSPCs in response to SDF-1 and SIP gradients [41, 42••]. Furthermore, our results showing a positive mobilizing effect by a non-toxic, small-molecule inhibitor of HO-1 (SnPP) suggest that blockade of HO-1 would be a promising strategy to facilitate mobilization of HSPCs in so-called poor mobilizers [42••]. Thus, this observation is highly relevant for developing more efficient mobilization strategies for HSPCs. Moreover, iNOS is also an inducible stress-response enzyme and produces nitric oxide (NO), which is a gaseous free radical molecule involved in several biological processes related to inflammation, tissue damage, and infections, and iNOS activity is enhanced during ComC activation as well [78]. Similarly, we found that inhibition of iNOS in HSPCs by the small-molecule inhibitor NIL also enhances mobilization of HSPCs. We propose again that these simple and inexpensive strategies to inhibit iNOS could be employed in poor mobilizers similarly as HO-1 inhibition to enhance process of HSPC mobilization [43••].

**Does an Inherited Defective Activation of the Mbl in Patients Explain Poor Mobilization Status?**

Since MBL ComC activation pathway plays a crucial role in triggering mobilization of HSPCs, one can ask if MBL deficiency may explain the poor mobilization status of some patients. Human MBL (MBL2) deficiency, the most common form of complement deficiency, is seen in 5–10% of the population, and correlates with the percentage of poor mobilizers. In humans, MBL is produced in the liver, and structural mutations in exon 1 of the human MBL2 gene at codon 52 (Arg → Cys, allele D), codon 54 (Gly → Asp, allele B), and codon 57 (Gly → Glu, allele C) independently reduce the level of functional serum MBL by disrupting the protein structure. Furthermore, several nucleotide substitutions in the promoter region of the human MBL2 gene at positions −550 (H/L polymorphism); −221 (X/Y polymorphism); and −427, −349, −336, del (−324 to −329), −70, and +4 (P/Q polymorphisms) affect MBL2 serum levels [79–81]. It remains to be shown whether MBL2 would be a good biomarker to predict poor mobilization; this concept is worth pursuing as could help to predict outcome of mobilization in HSPCs donors.

**Conclusions**

Optimization of stem cell mobilization is an important goal to obtain optimal graft that could significantly improve clinical outcomes following transplantation. Our data indicate (i) the crucial role of MBL–MASP in triggering ComC-directed trafficking of HSPCs, (ii) presence of functional crosstalk between the ComC and the CoA, and (iii) identify HO-1 and iNOS as negative regulators of ComC-mediated mobilization processes. Our observations are also highly relevant for other processes in which an increase in stem cell trafficking is observed in response to stress related to infection, tissue/organ injury, or strenuous exercise. Thus, these observations by modulating innate immunity responses will enable development of innovative treatment approaches, not only in hematopoietic settings but also when stem cells are employed to treat tissue/organ injuries, as seen in regenerative medicine.

**Acknowledgments** This work was supported by NIH grants 3 R01 DK074720 and R01HL112788, the Stella and Henry Endowment, and the Harmonia NCN grant UMO-2014/14/M/NZ3/00475 to MZR.
Compliance with Ethical Standards

Conflict of Interest  Mateusz Adamiak and Mariusz Z. Ratajczak declare that they have no conflict of interest.

Human and Animal Rights and Informed Consent  This article does not contain any studies with human or animal subjects performed by any of the authors.

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