**Drosophila** blood as a model system for stress sensing mechanisms

**Jiwon Shim**
Department of Life Science, College of Natural Science, Hanyang University, Seoul 133-791, Korea

The *Drosophila* lymph gland is the hematopoietic organ in which stem-like progenitors proliferate and give rise to myeloid-type blood cells. Mechanisms involved in *Drosophila* hematopoiesis are well established and known to be conserved in the vertebrate system. Recent studies in *Drosophila* lymph gland have provided novel insights into how external and internal stresses integrate into blood progenitor maintenance mechanisms and the control of blood cell fate decisions. In this review, I will introduce a developmental overview of the *Drosophila* hematopoietic system, and recent understandings of how the system uses developmental signals not only for hematopoiesis but also as sensors for stress and environmental changes to elicit necessary blood responses. [BMB Reports 2015; 48(4): 223-228]

**Fig. 1.** *Drosophila* larval lymph gland. Blood cells that proliferate and differentiate during the larval stages reside in the primary lobe of the lymph gland. PSC (blue), MZ (green), CZ (red, orange, and purple - plasmatocytes, crystal cells, and lamellocytes, respectively). Yellow cells in between the MZ and the CZ are differentiating blood cells that exhibit characteristics of both progenitors and mature blood cells.

**DROSOPHILA LYMPH GLAND AS A HEMATOPOIETIC MODEL SYSTEM**

*Drosophila* hematopoiesis is divided largely into two waves and an additional wave: embryonic, lymph gland, and larval hematopoiesis (1). The initial wave of hematopoiesis takes place in the embryonic head mesoderm, where undifferentiated blood cells are formed, proliferate, and differentiate into mature blood cell types, including plasmatocytes and crystal cells (2). Upon differentiation, embryonic blood cells disperse through the embryo and migrate throughout along specific paths (3). Later in the larval stage, an independent set of blood cells are found in the lymph gland, which originate from the cardiacogenic mesoderm (4-6). The lymph gland is composed of three compartments. Stem-like progenitors in the lymph glands locate in the core, called the Medullary zone (MZ), and the MZ cells give rise to mature blood cells in the outermost region, called the Cortical zone (CZ) (Fig. 1) (6). Similar to other stem cell systems, maintenance of the stem-like progenitors is controlled mainly by a microenvironment niche, also known as the Posterior signaling center (PSC) (7, 8). Differentiation of the stem-like progenitors produces at least three types of blood cells that are reminiscent of vertebrate myeloid cell lineages (6). Plasmatocytes, corresponding to vertebrate macrophages, represent more than 95% of total blood cells and function in cellular immunity and phagocytosis (2, 4, 5). Crystal cells are named after their resident crystalline structures of unidentified proteins and are known to mediate wound healing and melanization, similar to platelet in vertebrates (9, 10). While crystal cells represent approximately 5% of total blood cells under normal conditions, there is a specific cell type, called lamellocytes, that only appears upon immune challenge and takes part in the encapsulation of relatively large particles (11, 12).

The embryonic phase of hematopoiesis corresponds to the ‘primitive’ hematopoiesis of vertebrates and the lymph gland phase of hematopoiesis is referred to as the ‘definitive’ hema-

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topoiesis. Recently, an additional phase, larval hematopoiesis, has been recognized and found to play a role in the segmental colonization and expansion of larval circulating blood cells with the support of the peripheral nervous system. Larval hematopoiesis expands the pool of embryonic blood cells, enabling embryonic blood cells to persist and function properly during larval stages (13). During metamorphosis, blood cells in the lymph gland are dispersed into the circulation, and are in great need as they play major roles in cell engulfment, together with embryonic-origin blood cells (14). Thus, unless otherwise challenged, blood cells produced by the lymph gland do not participate in the larval blood circulation and only start to be released at the onset of pupariation (14).

**IMMUNE CHALLENGES AND THE BLOOD SYSTEM**

In the wild, flies commonly confront dangers of microbial infection, physical injury, and animal infestation as they feed in a microorganism-enriched environment. As a defense strategy, cellular and molecular mechanisms involving a multi-layered innate immunity are well developed in *Drosophila*, and are phylogenetically conserved in higher organisms (15). *Drosophila* can mount both cellular and humoral responses when attacked by bacteria, fungi, parasitic animals, and viruses (16). The humoral response is largely mediated by the expression of antimicrobial peptide (AMP) families, produced by tissues such as the fat body - equivalent to mammalian liver and adipose tissue - and released into the hemolymph to attack infectious microbes directly (15, 17, 18). The *Drosophila* immune system recognizes various types of infectious microbes and specifically induces corresponding AMPs from specific cell types, under the control of the Toll or IMD pathway (19). In addition to the AMPs, oral infection induces reactive oxygen species (ROS) in the intestinal epithelia, suggesting that there are multiple layers of humoral immune responses upon microbial infection (20, 21).

The cellular immune response occurs simultaneously with the humoral response at the level of the blood cells. Plasmacytocytes are responsible for the removal of small particles in the hemolymph, such as microorganisms and apoptotic cell debris (22). Receptors on the surface of plasmacytocyte recognize target particles and modify the cytoskeleton to engulf, followed by a series of phagosome formation and disruption events. Several receptors involved in phagocytosis have been reported. A CD36 homolog, croquemort, is responsible for scavenging apoptotic cells in the embryo (23). In addition, members of the scavenger receptor family (dSR-CI), the EGF-domain protein Eater, and the Ig-domain protein Dscam mediate bindings to Gram-negative and Gram-positive bacteria (24-27). Encapsulation is a powerful immune reaction against parasitic invaders carried out by a specialized blood cell, the lamellocyte. Lamellocytes only appear in the presence of invaders whose size is bigger than the phagocytic capacity of plasmacytocytes (6). Active encapsulation has been analyzed using wasp instestation, in which wasps lay their eggs into the larval hemocoel (28). Once the wasp egg is detected by plasmacytocytes through an unidentified signal, massive differentiation of lamellocytes takes place, either by differentiation of existing circulating plasmacytocytes (29) or differentiation of progenitors in the lymph gland (30). JAKSTAT, JNK, Toll pathways, chromatin remodeling, and the PSC-driven EGF signal have been identified as signals responsible for the differentiation and proliferation of lamellocytes (11, 31-34). Lamellocytes form a multilayered capsule around a parasitic particle, followed by subsequent melanization and particle breakdown. Another immune reaction that involves both humoral and cellular components of innate immunity is melanization, carried out by crystal cells. Melanization is controlled by a series of serine protease cascades that ultimately produce active phenol oxidase, and this enzyme catalyzes tyrosine-derived phenols to form melanin. Several intermediate compounds generated during melanin synthesis are cytotoxic, so they participate in killing pathogens, indicating the humoral activity of melanization (35). The biggest reservoir of phenol oxidase is the crystal cells. Mature crystal cells accumulate unidentified crystalline proteins, including phenol oxidase, in the cytoplasm and are readily burst upon a specific signal, JNK, to release and activate the stored proteins (9). Thus, disruption of crystal cells is tightly associated with melanization at the level of the blood cells, which, in turn, leads to the humoral killing processes in a systematic manner.

**REACTIVE OXYGEN SPECIES AND THE BLOOD SYSTEM**

An interesting angle that has been identified with the *Drosophila* lymph gland is that wildtype Medullary zone cells contain significantly high ROS levels compared with their neighboring differentiated progeny (36). Scavenging ROS from the blood progenitors, by overexpression of antioxidants such as GTPx, retards their differentiation into mature blood cells. In contrast, increasing ROS above their normal levels by reducing Sod2 triggers precocious differentiation of mature blood cells. These studies thus provide the new insight that moderately high levels of ROS in blood progenitors serve as developmental signals and are essential for the maintenance of progenitors, while excessive ROS are harmful and deleterious to the cells. A recent study has found a detailed cellular mechanism underlying how high ROS expression in the progenitors is involved in both the maintenance and differentiation of blood cells at specific points in development (37). High ROS levels prime the blood progenitors in the early larval stages and are required for the expression of the adherent junction protein, E-cadherin, that is known to control stem cell integrity and facilitate signal transduction (38). Intriguingly, high levels of ROS are also detected in mammalian hematopoietic stem cells (HSCs) as well as in the common myeloid progenitors (CMPs), and maintaining moderate levels of ROS is key for their survival and normal cell cycle entry (39).
In addition to the role of ROS in cell fate decisions, ROS are also involved in the early proliferation of blood progenitors in the lymph gland (40). Increases in Target of rapamycin (TOR) signaling in the lymph gland cause its overgrowth, along with the elevated levels of ROS. Additionally, a decrease in ROS levels in the high TOR activity background causes abnormal early proliferation of the lymph gland to return to the normal proliferation profile, indicating a role of ROS in proliferation. Consistent with this, a genetic modifier screen of the leukemic gene,AML1-ETO identified FoxO and superoxide dismutase 2 (SOD2) as suppressors of the leukemic phenotype via reduction of ROS (41). Overexpression of AML1-ETO in the larval blood results in the aberrant proliferation of circulating blood cells that can cause the formation of melanotic tumors. This phenotype is due to an increase in the circulating blood progenitors that express high levels of ROS, similar to the progenitors found in the lymph gland. Overexpression of FoxO or SOD2 in the circulating blood progenitors is sufficient to reduce the number of ROS-positive progenitors, indicating a role for ROS in the proliferation of blood progenitors. The role of high ROS has been studied extensively, particularly with regard to effects in apoptosis and the stress response. However, studies using the Drosophila blood system highlight the physiological roles of internal ROS in the cellular and developmental signaling that are essential for cell fate decisions as well as blood system integrity. Given that ROS levels are influenced by an animal’s physiological status and by chemicals that alter redox status (37), it is possible that blood progenitors can be used as a redox sensor that monitors environmental toxicity.

**HYPOXIA AND THE BLOOD SYSTEM**

An adequate oxygen supply is one of the most critical determinants of an aerobic organism’s survival. A representative hypoxic response mechanism functions mainly via hypoxia inducible factor HIF1, which recognizes physiological oxygen levels and transduces environmental information into gene expression (42). At the organismal level in Drosophila, tracheal cells respond to hypoxia with a heightened sensitivity to declining oxygen levels and undergo tracheal branch remodeling for the better reception of oxygen (43). In addition to the trachea, Drosophila blood cells are highly susceptible to hypoxic condition and increase the number of mature crystal cells in the lymph gland (10). Although the physiological role of these crystal cells in the hypoxic response is still unclear, it is possible that the crystal cells react to hypoxia in the way that the hemoglobin does in vertebrates or is preparation for possible injuries that may be caused by prolonged hypoxia. Interestingly, unlike other somatic cells, crystal cells stably express Sima, the Drosophila ortholog of Hifα, even under normoxic conditions. Accordingly, either increase in Sima due to hypoxic stress or disruption of developmental Sima by modifying the Sima-Notch interaction further expands the number of crystal cells in the lymph gland, indicating that Sima in the crystal cells plays dual roles in both the stress response and developmental control of cell fate decision. It is noteworthy that vertebrate myeloid cells also maintain high Hifα in normoxic conditions to maintain their cellular energy pools and the ability to mount an inflammatory response (44), and this conservation indicates a possible parallel mechanism between the Drosophila blood system and that of vertebrates.

A study with adult hemocytes provides an interesting aspect that overexpression of Hsp70 (Heat shock protein 70) in blood cells contributes to a remarkable survival benefit in severe hypoxia and oxidative stress by inhibition of systemic ROS (45). In addition to the systemic role of blood cells in immunity, this study provides another example of a systemic function of hemocytes in the stress response, which is not yet fully understood.

**NUTRITION AND THE BLOOD SYSTEM**

Nutritional deprivation directly impinges on the maintenance of blood progenitors in the lymph gland and induces differentiation of mature blood cells (40, 46-48). Indeed, 24 h starvation of third instar larvae induces precocious differentiation of mature blood cells and a concomitant decrease in the progenitors. Moreover, chronic starvation, >48 h, results in blood cell differentiation accompanied by a decrease in the size of the PSC and the lymph gland. These phenomena are related to the Drosophila insulin-like growth factor 2 (Ilp2), which is a reminiscent of the mammalian insulin (49) and plays a key role in nutrition-mediated tissue growth. Systemic insulin secreted from the brain neuroendocrine cells is perceived by the insulin receptor (InR) either in the PSC or in the MZ cells, and is required for proper maintenance of the lymph gland integrity. Lack of InR in the PSC causes a reduction in the niche size, which indirectly affects progenitor cell fate through changes in the expression of Hedgehog (Hh) (8). Moreover, loss of InR in MZ cells gives rise to precocious differentiation of mature blood cells through a cell-autonomous PI3K/AKT and TOR pathway that, in turn, controls Wingless in the MZ cells (40, 46-48). These studies suggest that systemic nutrient and insulin signaling impact the local lymph gland-based signals to achieve blood homeostasis, and loss of the nutritional signal forces blood cells to differentiate. Interestingly, acute starvation generates hallmarks of inflammatory responses, including lamellocyte differentiation, crystal cell rupture, and infiltration of blood cells into the fat body, similar to the chronic inflammation phenotype observed in metabolic disorders (47). Moreover, it has been identified that there is a coordinated interaction between immune and metabolic signals to reallocate energy utilization from non-essential processes, such as growth, to more immediate needs, such as immunity (50). Thus, it is possible that starvation makes the blood progenitors withdraw their stemness and build up the blood cell repertoire to reinforce cellular immunity. From a developmental point of view, insulin-mediated blood regulation occurs naturally during the developmental timeline. Differenti-
ation of the mature blood cells takes place in accordance with animal's feeding behavior, and the lymph gland finally disintegrates upon pupariation when blood cells are in great needs during metamorphosis (14). Insulin- and nutrition-mediated stem cell control mechanisms are conserved in other stem cell types in Drosophila, indicating that acquiring proper amount of nutrition is universally indispensable for stem cell control, although the specific targets and roles of insulin in each stem cell vary.

**ADENOSINE AND THE BLOOD SYSTEM**

Homeostasis in the lymph gland is achieved by several factors including local maintenance signals from the PSC, cell-autonomous control of the MZ cells, systemic signals from the brain, and the equilibrium signal from differentiating cells in the CZ cells (51). It has been established that the PSC expresses Hh, which is the primary signal of MZ progenitors’ maintenance through the activation of Cubitus interruptus (Ci) (8). In addition to the Hh pathway, Pvf1 originating from the PSC is transported to the maturing blood cells in the cortical zone, where it binds to its receptor, Pvr, leading to production of Adgf-A via activation of STAT. Adgf-A is a secreted adenosine deaminase that functions to maintain low levels of adenosine in the progenitors. While adenosine is necessary for early progenitor proliferation, high levels of it lead to continuous progenitor proliferation and loss of their maintenance. Thus in this process, a balance between Adgf-A and adenosine plays a key role in achieving blood progenitor homeostasis and the presence of adenosine is likely to operate in a mechanism similar to the “quorum sensing” found in the prokaryotic colonization (52). Thus, a critical level of adenosine is decisive for the MZ population. Given that adenosine is a key precursor of various metabolic processes, adenosine-derived developmental signals may be associated with stress responses, particularly mitochondrial or metabolic stresses. Adenosine-associated developmental regulation is another example of a mechanism involved in both development and stress responses, making the system very efficient and sensitive to systemic environmental changes.

**OLFACTION AND THE BLOOD SYSTEM**

Niche-dependent mechanisms of hematopoietic stem/progenitor cell maintenance have been studied extensively in both vertebrates and invertebrates (53). Mechanisms independent of niche-derived signals that operate in a more systemic manner but affect blood progenitors have recently started to emerge (54). In addition to nutrition-directed blood regulation, olfactory sensation impacts the blood progenitors in the lymph gland via secretion of physiological levels of γ-aminobutyric acid (GABA) into the blood stream (55). GABA is expressed in a subset of neuroendocrine cells, and the release of GABA is highly dependent on olfactory stimulation. Thus, olfactory dysfunction decreases GABA secretion from the neuroendocrine cells and reduces levels of GABA in the bloodstream. Blood progenitors in the lymph gland express the metabotropic GABA_{B} receptor that allows these cells to sense GABA in the circulation, leading to an increase in their cytosolic calcium concentration, essential for blood progenitor maintenance. Although downstream targets of high calcium in the progenitors are unclear, it is possible that developmental signals in the MZ cells are able to use the calcium and cross-talk to each other. GABA is an evolutionarily conserved molecule. In plants, GABA serves as a stress signal as well as a metabolite (56), while GABA in vertebrates has been studied extensively with regard to its neurotransmitter function (57). Interestingly, there are related studies that GABA can be measured in the bloodstream of mammals, including humans (58), and the GABA_{B} receptor is expressed in primary human HSCs (59), indicating a possible parallel mechanism conserved in higher organisms. It is interesting to note that olfaction can be associated with the developmental timing and stress response. Because olfactory receptors are dissociated upon pupariation, premature loss of olfactory receptors during larval stages is reminiscent of pupariation when extensive mature blood cells are to differentiate and disintegrate from the lymph gland.

**PERSPECTIVE**

The overall physiological status of an organism is critical for determining stem and progenitor cell fate. Given that purpose of harboring stem/progenitor cells is to provide a regenerative capacity, systemic controls of stem/progenitor cells are critical to ensure their cell fate determination as stem cells are tightly associated with rapid growth, immunity/injury, and other environmental challenges to meet the needs of an organism (60). Simultaneously, there is a developmental timeline that larval stem/progenitors should follow to complete normal development. Thus, instead of using two different mechanisms for development and stress response, animals use identical mechanisms that allow easy conversion, depending on their external circumstances.

In the lymph gland, mature blood cells appear in the mid-to-late second instar larval life, followed by rapid differentiation in late third instar (6). At the onset of pupariation, the lymph gland disintegrates and bursts open to release blood cells (14). Even with no infection, mature blood cells are essential during metamorphosis to scavenge and phagocyte larval cells and bacteria released from the reforming gut. We can view the blood progenitor control mechanisms introduced in this review as signals that exist physiologically during larval development and, as the larvae approach pupariation, these processes are attenuated. As third instar larvae start wandering, the larval insulin level drops naturally (61). At this time, larvae generate a cuticle, causing reduced oxygen diffusion, and dissociate larval olfactory receptors to decrease sensory perception. Finally, remodeling of the gut releases chemical/bacterial substances and metabolic intermediates, which are
likely to induce immune responses, cause ROS induction, and change adenosine levels. Thus, all the conditions that facilitate blood differentiation are also developmental consequences of the initiation of pupariation. Interestingly, different types of stresses are also sensed by these pathways and cause precocious progenitor differentiation, thus functioning as stress sensors in the system.

It seems likely that increased differentiation of mature blood cells in various stress responses may facilitate the initiation of more rapid immune reactions by acquiring additional challenges. However, whether this blood differentiation response is directly related to the systems’ protective mechanism is not yet clear. It will be interesting to understand this mechanism, of which process contains primitive features of boosted immunity by the increased blood repertoire, similar to the adaptive immune system in vertebrates. Given that larvae are challenged by various external changes, it is possible that Drosophila has additional and novel mechanisms that can affect both development and stress responses.

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