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Review

Innate immune system and tissue regeneration in planarians: An area ripe for exploration

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ABSTRACT

The immune system has been implicated as an important modulator of tissue regeneration. However, the mechanisms driving injury-induced immune response and tissue repair remain poorly understood. For over 200 years, planarians have been a classical model for studies on tissue regeneration, but the planarian immune system and its potential role in repair is largely unknown. We found through comparative genomic analysis and data mining that planarians contain many potential homologs of the innate immune system that are activated during injury and repair of adult tissues. These findings support the notion that the relationship between adult tissue repair and the immune system is an ancient feature of basal Bilateria. Further analysis of the planarian immune system during regeneration could potentially add to our understanding of how the innate immune system and inflammatory responses interplay with regenerative signals to induce scar-less tissue repair in the context of the adult organism.

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1. Introduction

The process by which form and function is reestablished after tissue injury has puzzled the scientific community for hundreds of years [1–3]. Tissue injury triggers localized and systemic signals that orchestrate mechanical and cellular responses aimed at reducing the surface area of the wound, repair damage, and coordinating functional integration between new and pre-existing structures [4,5]. The immune system and the inflammatory responses associated with injury have been implicated as critical modulators of wound repair and regeneration [6–10]. For instance, effective repair of wounded skin in mammals, and regeneration of complex structures in amphibians (e.g. tail and limbs) rely on the correct synchronization of the immune response upon injury [6–11]. Recent studies suggest that the difference in regenerative capacity of some species is inversely correlated with the complexity of the immune system (Fig. 1) [7,9]. However, the mechanisms integrating immune responses to injury and the process of tissue repair remain poorly understood.

Upon injury the host-mediated immune response not only defends against infection, it facilitates the removal of cellular debris near the site of the wound. These functions may also modulate the cellular response that initiates repair and reestablishes tissue function. It is conceivable that both the injury-induced immune response and the process of tissue repair evolved together to promote and preserve multicellularity. The possibility of shared origin is supported by the fact that regeneration is an attribute widely distributed across multicellular organisms, and immunity is an ancient feature that emerged from the common ancestor of Cnidaria and Bilateria [12–18]. Coexistence of both regeneration and the immune system is observed in the simplest animals such as the diploblastic cnidian Hydra, which regenerate entire body parts in presence of a primitive complement system and which have many genes associated with the mammalian immune response [16,17,19,20]. Evaluating the evolution and intermingling of the immune response and process of regeneration following injury may provide a new depth to our understanding of tissue repair mechanisms across the animal kingdom.

Planarians are non-parasitic flatworms that provide compelling evolutionary reasons to analyze injury-induced immune responses and the process of regeneration in metazoans [21,22]. Planarians display bilateral symmetry and derivatives of all three germ layers (ectoderm, mesoderm and endoderm). Adult planarians can regenerate any part of their body and constantly renew aging and damaged tissues (Fig. 2), which are features missing...
in other commonly used invertebrate models (e.g. Drosophila and Caenorhabditis elegans). Regeneration and adult tissue turnover in planarians proceed through activation of somatic stem cells known as neoblasts [23–25]. Planarians have provided a classical model to study regeneration for over 200 years, but their immune system remains largely unexplored [1,26–28]. To our knowledge, no systematic analysis of the planarian immune system has been published to date. We present a brief survey of the most salient components of the innate immune system through genomic analysis between the freshwater planarian Schmidtea mediterranea and other animal species. We analyzed two sources: the S. mediterranea genome database, SmedGD [29] and transcriptomic work of Sandman et al. [30] to evaluate genes associated with the immune system and their expression during regeneration. Our findings suggest evolutionary conservation of the trilobed immune system. We identified components of the S. mediterranea immune system that may protect against infections, distinguish between commensal and pathogenic microorganisms, and that actively participate during wound repair, regeneration and maintenance of adult tissues.

2. The planarian model system and tissue regeneration

Planarians are known for their extraordinary capacity to regenerate adult tissues. These animals can regenerate any part of their body including their entire digestive system, brain and neural connections, muscles, and connective tissues. Planarians are easy and inexpensive to maintain under laboratory conditions and are amenable to molecular, genetic, behavioral and computational analysis [1,26,31–34]. During the past 20 years, planarian research has attracted attention due to the accessibility of this model organism and the opportunities to progress our understanding in long standing biomedical problems associated with regeneration, cancer and degenerative diseases. Several tools available to aid in the study of planarians, include the S. mediterranea genome, proteomic data, results from high-throughput RNAi-screens and transcriptomic analysis [1,25,26,29,32,34–38]. Altogether with the availability of standard genetic and biochemical techniques, planarians present an attractive model to study wound repair and regeneration and the possible interplay of the immune system during these processes.

Detailed information about the biology of planarians and recent advances on research pertaining to its regenerative capacities has been reviewed elsewhere [1,23–25,28]. In this section, we aim at providing a brief overview of the process of regeneration in S. mediterranea, the most commonly used planarian species. Planarian regeneration is quick, taking about a week to reestablish form and function. Immediately following amputation, contraction around the injury reduces the damaged surface. In the first 24 hours, the regenerative process can be observed with H+/K+-ATPase-mediated tissue depolarization around the wounded area [39–42]. The antagonistic process of cell death and proliferation a process of cell death and proliferation around the injured area is critical for the progression of regeneration. The H+/K+-ATPase-mediated ion flux and consequent tissue depolarization is required for anterior tissue specification and may provide signals that guide neoblast proliferation, migration and remodeling [5,39,40]. Decisions regarding polarity are defined within the first few hours post-amputation and differentiated tissues may provide molecular cues and guidance during regeneration [1,25,28,43]. Neoblasts, the only dividing cells in adult planarians survive as the exclusive source of new cells during the formation of the blastema (a regenerative outgrowth around the injured area).

Neoblast division gives rise to cellular progeny that migrate and differentiate within the blastema. Remodeling and integration between the new and pre-existing tissue is an active process during regeneration but little is known about the signals mediating these
events. The planarian model provides the opportunity to analyze local and systemic signaling during tissue repair and regeneration, providing a unique prospective for understanding injury-immune responses in the adult body.

3. The planarian innate immune system

The immune system in adult animals is essential to maintaining body homeostasis as it distinguishes between commensal and pathogenic microorganisms, protects against infections, and serves as a signaling system during tissue repair. Most studies associating the immune system and flatworms focus on parasitic species that invade and activate host immune responses [44–46]. There is a wealth of literature associated with this aspect of host-interaction that has important practical and clinical applications and is discussed elsewhere [44–46]. We instead concentrate on the immune response against pathogens that invade non-parasitic planarians and the injury-immune activation associated with adult tissue regeneration.

The immune response is comprised of two parts: the innate and adaptive responses. Innate immune responses make up the first responders of the immune system and impart broad (often termed non-specific) protection from classes of pathogens. This response is immediate, occurring within only a few hours of infection. The innate immune system provides the first line of defense giving the adaptive (or acquired) immune response the days to weeks needed to fully develop. The adaptive immune system then has the task of clearing the microbial infection using pathogen-specific immune responses [47,48]. There is no systematic proof of an advanced adaptive immune response in invertebrate animals [49]. On the other hand, innate immune responses are markedly similar between many invertebrates and vertebrates [50–52].

The innate immune system is the first barrier of defense against invasive microbes. It is comprised of many players, from the physical barrier and secretion of mucus, anti-microbial peptides, fatty acids and lysozymes to phagocytes that capture and digest the invading pathogens [53]. Microbes express classes of surface proteins, termed pathogen-associated molecular patterns, which are

**Table 1**

<table>
<thead>
<tr>
<th>Innate immune component</th>
<th>Domain found</th>
<th>Maker ID from Planarian database</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toll-like receptor (TLR) or MyD88 or CD14</td>
<td>Leucine rich repeat</td>
<td>mk4.000112.15, mk4.000148.12, mk4.002858.00, mk4.007365.00, mk4.019024.00, mk4.026212.00</td>
</tr>
<tr>
<td></td>
<td>TIR domain</td>
<td>mk4.000285.01, mk4.000346.04, mk4.027932.00</td>
</tr>
<tr>
<td>SARM 1</td>
<td>SAM domain + TIR domain</td>
<td>mk4.008544.03</td>
</tr>
<tr>
<td>Selectins</td>
<td>C-type lectin like domain</td>
<td>mk4.000042.08, mk4.000389.11, mk4.020584.01</td>
</tr>
<tr>
<td>Complement</td>
<td>CUB domain</td>
<td>mk4.001431.05, mk4.001438.03, mk4.000798.07, mk4.003730.02, mk4.007851.01</td>
</tr>
<tr>
<td></td>
<td>Alpha2-macroglobulin</td>
<td>mk4.004852.05, mk4.014934.00, mk4.043975.00, mk4.008692.00, mk4.008429.01, mk4.043975.00, mk4.008251.00</td>
</tr>
<tr>
<td>Complement factors or Integrins</td>
<td>Von Willebrand factor type A domain</td>
<td>mk4.007402.00</td>
</tr>
<tr>
<td>Cell adhesion molecules (ICAM, VCAM)</td>
<td>Immunoglobulin I-set domain</td>
<td>mk4.006497.00</td>
</tr>
<tr>
<td>Urokinase-type plasminogen activator</td>
<td>Kringle Domain</td>
<td>mk4.000652.03</td>
</tr>
<tr>
<td>IRAK like kinase 1</td>
<td>Possible IRAK similar to C. elegans and Protein Kinase domain</td>
<td>mk4.004443.00, mk4.004443.01, mk4.027874.00</td>
</tr>
<tr>
<td>TRAF-6</td>
<td>TIR domain of TRAF and TRAF MATH domain</td>
<td>mk4.000002.01–mk4.000002.47</td>
</tr>
<tr>
<td>TRAF related</td>
<td>MATH domains</td>
<td>mk4.000007.04–mk4.000007.34</td>
</tr>
</tbody>
</table>

Source: SmedGD [29].

recognized by host pathogen recognition receptors (PRR) expressed on host cells. In vertebrates, recognition of pathogens through these PRRs results in inflammatory signals, differentiation of immune cells and migration of leukocytes into infected or damaged tissue [53].

The planarian immune system remains largely unexplored, but it is known that under laboratory conditions planarians suffer from infection just as other eukaryotic species, with bacterial infection a common threat in routine maintenance [54]. Treatment with antibiotics usually resolves the infection but it is unknown what types of bacteria commonly infect *S. mediterranea* under laboratory conditions. Nonetheless, a mechanism of pathogen recognition must exist in *S. mediterranea* to prevent or control these infections. Histological and functional work identified active phagocytic cells around areas of wounding; suggesting the possibility that a primitive innate immune system participates during tissue repair, cancer, and infection [55–57]. Table 1 includes a collection of candidate genes from the *S. mediterranea* genome that are potential components of the innate immune system [29]. Table 2 provides a sample of candidate genes of the innate immune system and their expression during head regeneration in *S. mediterranea* [30]. Details about these planarian candidate innate immune genes are provided below.

3.1. Mucus and anti-microbial peptides

Surfaces that secrete mucus are highly susceptible to pathogens and mucosal surfaces are known to contain large amounts of bacteria that could be pathogenic or symbiotic [58,59]. Mucus containing antimicrobial activity can be generated as a front line of defense. Microbial adhesion to mucus limits colonization of the pathogens, providing a defense mechanism to infection [60]. Planarians are normally covered by a thin layer of mucus that may act as a protective barrier and allows the presence of commensal microorganisms. Stressful situations, such as wounding, in *S. mediterranea* induces secretion of mucus that may have a double role in blocking pathogen entry and aiding in the healing process. Recent evidence suggests that planarian derived mucus
proteins display similarities with mucus secretion from humans [35]. However, it is unclear whether wound repair and regeneration in planarians require components of the mucus to repair tissue.

Anti-microbial peptides are small peptides with clusters of positively charged and hydrophobic amino acids [61]. These small peptides are evolutionarily conserved and are known to have immunomodulatory functions [52]. For example, anti-microbial peptides are known to be secreted upon injury in insects [62]. Altincicek et al. screened for septic wound inducing genes by introducing bacterial lipopolysaccharide into the wound site of planaria, and identified the induction of several potential anti-microbial peptides [63].

These anti-microbial homologs are likely involved in regulation of commensal bacterial and pathogen killing in planaria. Further, a sequence suggested to be an antimicrobial peptide resistance and lipid A acylation protein, PagP, was found within the planarian genome (mk4.054786.01.01). How these anti-microbial peptides are activated during injury remains to be ascertained.

### 3.2. Pathogen recognition receptors

Initiation of innate immune responses relies on early signals following recognition of pathogen- or damage-associated molecular patterns (PAMPs and DAMPs) [64]. These early events involve recognition of molecules such as lipopolysaccharide, flagellin or dsRNA, found on or in invading pathogens or released by damaged and dying cells [64]. Two members of the PRR family, NOD-like receptors and toll-like receptors (TLRs) are conserved from early invertebrates to mammals and are maintained with a small number of protein structures [65].

TLRs are comprised of a cytoplasmic tail, a transmembrane and an extracellular domain that contains leucine rich repeat (LRR) motifs [66]. Based on crystallographic studies LRRs are shaped in a 3D horseshoe format that contain the leucine rich repeats. This domain is important for interaction with other molecules, pathogen detection and provides structure [66,67]. In mammals interactions through these LRRs trigger a conformation change to the TLR that results in a signaling cascade that activates proinflammatory cytokines and type 1 interferons [68]. LRRs are highly conserved across species, and their primary function is in the formation of protein–protein interactions [67]. Vertebrate and invertebrates have various numbers of TLRs each with specific functions, mostly targeted toward pathogen recognition [69–71].

<table>
<thead>
<tr>
<th>Innate immune component</th>
<th>Domain found</th>
<th>Maker ID</th>
<th>Time post-amputation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toll Like Receptor (TLR) or MyD88</td>
<td>Leucine rich repeat</td>
<td>mk4.007365.00</td>
<td>↑ ↑ ↑ ↑</td>
</tr>
<tr>
<td></td>
<td>Leucine rich repeat</td>
<td>mk4.002858.00</td>
<td>↑ ↑ ↑ ↑</td>
</tr>
<tr>
<td></td>
<td>TIR domain</td>
<td>mk4.000285.01</td>
<td>↑ ↓ ↓ ↓</td>
</tr>
<tr>
<td>Selectins</td>
<td>C-type lectin like domain</td>
<td>mk4.000389.11</td>
<td>↑ ↓ ↑ ↓</td>
</tr>
<tr>
<td></td>
<td>C-type lectin like domain</td>
<td>mk4.020584.01</td>
<td>↑ ↓ ↓ ↓</td>
</tr>
<tr>
<td>Complement</td>
<td>CUB domain</td>
<td>mk4.000798.07</td>
<td>↓ ↓ ↓ ↓</td>
</tr>
<tr>
<td></td>
<td>CUB domain</td>
<td>mk4.003730.02</td>
<td>↓ ↑ ↓ ↓</td>
</tr>
<tr>
<td></td>
<td>CUB domain</td>
<td>mk4.001431.05</td>
<td>↓ ↑ ↓ ↓</td>
</tr>
<tr>
<td></td>
<td>alpha2-macroglobulin</td>
<td>mk4.005639.01</td>
<td>↓ ↑ ↓ ↓</td>
</tr>
<tr>
<td></td>
<td>alpha2-macroglobulin</td>
<td>mk4.008692.00</td>
<td>↓ ↑ ↓ ↓</td>
</tr>
<tr>
<td></td>
<td>C1q domain</td>
<td>mk4.008251.00</td>
<td>↑ ↓ ↓ ↓</td>
</tr>
<tr>
<td>Complement factors or Integrins</td>
<td>Von Willebrand factor type A domain</td>
<td>mk4.007402.00</td>
<td>↓ ↑ ↑ ↓</td>
</tr>
<tr>
<td>IRAK like kinase 1</td>
<td>Possible IRAK similar to C. elegans and Protein Kinase domain</td>
<td>mk4.027874.00</td>
<td>↑ ↓ ↓ ↓</td>
</tr>
<tr>
<td>TRAF-6</td>
<td>zn finger domain of TRAF</td>
<td>mk4.000002.06</td>
<td>↑ ↑ ↓ ↓</td>
</tr>
<tr>
<td></td>
<td>TRAF MATH domain</td>
<td>mk4.000002.07</td>
<td>↑ ↑ ↓ ↓</td>
</tr>
</tbody>
</table>

Our analysis of the planarian genome identified several gene segments that contain LRRs (mk4.007365.00, mk4.002858.00, mk4.000112.15, mk4.026212.00, mk4.000148.12). Whether these sequence similarities have TLR-like activity or are unrelated repeats remains to be investigated. CD14, also involved with innate immune function contain multiple LRR repeats, and whether the LRR repeats found in planarians belongs to TLR or CD14 functionality is unknown. Recent transcriptomic analysis performed on a head regeneration time course found upregulation of LRR homolog mk4.007365.00 and mk4.002858.00 from one to 72 h post-amputation [30].

A planarian genome wide search for TIR, the cytoplasmic domain of TLR, identified four TIR-like domain sequences (mk4.008544.03, mk4.000285.01, mk4.000346.04, mk4.027932.00). One gene segment (mk4.008544.03) contains two SAM domains suggesting that this might be a SARM1 protein. SARM1 proteins negatively regulate innate immune function by blocking signaling downstream of the PRRs [72]. None of the TIR domain containing gene segments included any LRR sequences suggesting that these might not be TLRs, or that the pattern recognition motifs in the planaria differ from traditional LRR repeats. One TIR-like sequence in planaria (mk4.000285.01) is also upregulated for 6 h post-amputation [30]. Upregulation of these TIR and LRR motifs may correspond with detection of, or response to, infection during injury, and warrants further attention.

Mammalian TLR family members share several signaling molecules including the adaptor protein MyD88, TRAF6 (TNF receptor-associated kinase 1), the protein kinase IRAK (IL-1R-associated kinase) and TAK1 (TGF-beta-activated kinase) binding protein. MyD88 protein also contains a TIR domain and a death domain. Though we found no significant homology for death domains near the four planarian TIR domains, we cannot eliminate the existence of a MyD88-like adaptor protein in planaria. TRAF proteins contain three domains: a MATH domain, RING-type zinc finger domains and TRAF-type zinc finger domains. Analysis of the planarian genome identified MATH homologs domains (mk4.000007.00 and mk4.000002). The gene segments between mk4.000002.00.01 and mk4.000002.33.01 contains both MATH and RING-type zinc finger domains and is upregulated throughout regeneration [30]. Region mk4.000007.00 also remain upregulated 1–72 h post-amputation. The gene segments between mk4.000002.00.01 and mk4.000002.33.01 contains both MATH and RING-type zinc finger domains and is also upregulated throughout regeneration. We identified one possible TAK1 homolog...
Based on these findings, it seems likely that planaria have some form of pathogen-recognition and signaling to distinguish between self-tissue and pathogenic microbes.

3.3. Complement cascade

The complement system is a highly conserved component of innate immunity. In mammals, complement functions to recognize and clear invading pathogens by inducing inflammatory responses, activating phagocytic cells and directly lysing microbes [73]. In mammals the complement system is made up of a cascading series of protein interactions that begin via three independent pathogen recognition pathways [73]. These three pathways converge on one protein C3 and continue through an identical cascade to form a membrane attack complex and amplify inflammatory signals. One arm of this system uses antibodies-coated pathogens in the recognition of pathogens and initiation of the complement cascade [73]. Although in mammals complement links the innate and adaptive immune responses, the complement system appears to be very ancient, with complement-like proteins found in most invertebrates and vertebrates [17,74,75]. We identified several complement factors and domain homologies within the S. mediterranea genome. The CUB domain (for complement C1r/C1s, Uegf, Bmp1) is found in the C1 component of the complement cascade. We have identified several homologs for the CUB domain (mk4.001431.05, mk4.001438.03, mk4.000798.07, mk4.007851.01, mk4.003730.02) and found mk4.000798.07 to be upregulated 6h post-amputation [30]. mk4.001431.05 and mk4.003730.02 which are found adjacent to a Sushi domain (mk4.003591.01), another domain found in the C1 complement subcomponent, remained upregulated very early during regeneration. A C1q-like domain (mk4.008251.00), the C terminal domain of the C1 enzyme that triggers classical complement activation [76] was also upregulated post-amputation up to 6h [30]. Several C3 complement homologs for the alpha-2-macroglobulin domain were also identified in planaria. As with other early invertebrates, planarians appear to have a conserved complement system that may act to control invading pathogens, but this requires confirmation and further study.

3.4. Phagocytic cells (reticuloctyes)

Macrophages are known to play an important role in the resolution of acute inflammation by clearing apoptotic and necrotic tissue due to injury and immune responses [77]. Macrophages also ingest and breakdown invading pathogens and in this way act in host defense. In mammals macrophages differentiate down two distinct pathways dependent upon the local signals encountered. M1 classical macrophage activation results in macrophages that produce inflammatory cytokines and act in host defense to control microbial infections. Alternatively activated macrophages aid in wound repair by phagocytosing damaged tissue and secreting growth factors [78]. Macrophage function is essential during tissue repair in most vertebrate animals including mice [79], zebrafish [80], and salamanders [8]. However, wound healing in PU.1 mice (no neutrophils or macrophages) occurs without scarring, establishing alternative regulatory modes for phagocytic cells during regeneration.

In planarians, a phagocytic, mesenchymal cell, termed the reticular cell has been previously described [55–57]. Within 10h of injury in the presence of bacteria, the reticular cell migrates into the wound, phagocytoses and encapsulates bacteria. Reticular cells thus have the capability to recognize foreign particles as distinct from self, migrate and phagocytose pathogens. Phagocytic engulfment of cellular debris has also been observed during regeneration after fissioning and after amputation [55–57]. We speculate the reticular cells in planarians may represent a primitive form of macrophage.

Although it is unconfirmed whether planarian phagocytic cells express PRRs, in other organisms, macrophages are one of the major cells types that express PRRs and respond to pathogen-associated molecular patterns [53,81]. Macrophages produce many proteins that induce inflammation, migration of other cells into the site of infection or injury, killing of pathogens and down-modulation of these responses following clearance of the infection or damage [78]. Perforin is a lytic enzyme produced and secreted by mammalian macrophages that lyses invading pathogens and infected cells. A membrane attack complex component/perforin/C9 homolog was identified during sepsis-injury in planaria [63], suggesting a role for this enzyme in host defense of planaria. Alterations in the debris removal process or inflammatory responses might alter the final outcome of the repair, thus indicating a potential role for phagocytic cells in both host defense and wound repair and regeneration.

A recently identified group of molecules secreted by macrophages, maresins (macrophage mediators in resolving inflammation) enhance macrophage phagocytosis and limit local polymorphonuclear neutrophil infiltration [82]. When exposed to human maresin MaR1, the rate of planarian anterior tissue regeneration increased [82]. Further, planaria biosynthesized MaR1 upon injury, and the formation of MaR1 was blocked by a lipoxigenase (LOX) inhibitor, eliminating the enhancement in tissue regeneration. Together these data indicate that key planarian signaling components can respond to human MaR1, and thus conservation exists between human and planarian maresins and their signal cascades. These data further support a link between immune cell activities and regenerative mechanisms.

3.5. Cell adhesion molecules

Cell adhesion molecules are typically transmembrane proteins expressed on the cell surface. These proteins mediate cell–cell interactions including, signal transduction, cellular communication, and migration. In mammals, cell adhesion molecules have been divided into four families based on protein structure: Immunoglobulin (Ig) superfamily, cadherins, selectins and integrins [84,85].

The Ig superfamily is calcium-independent and composed of variable numbers of Ig-like repeats. Analysis of the planarian genome revealed several Ig-like domains (mk4.006497.00) that are commonly found in adhesion molecules such as the ICAMs. The cadherin family, a calcium-dependent cell adhesion molecule, usually contains 3–5 internal repeats of cadherin and form homodimers. Cadherins play a crucial role in mediating innate and adaptive immune functions by maintaining epithelial barrier function and regulating leukocyte migration [86,87]. Most integrins function as receptors for proteins within the extracellular matrix and can bind to a single ligand or to multiple ligands. In mammals, selectins are adhesion molecules with a trans-membrane glycoprotein that are expressed on leukocytes and activated epithelial cells [88]. The vertebrate selectin family consists of L-selectin, E-selectin and P-selectin and these are mostly involved with leukocyte homing and migratory responses to inflammation or pathogens. They mediate leukocyte rolling and adhesion during mammalian wound repair and leukocyte extravasation [89]. Selectins belong to the C-type lectin superfamily of proteins, are calcium-dependent and are widely found in metazoans [90]. Although C-type lectin domains
have been linked to a large range of functions, pathogen recognition remains a primary function amongst most of this family [90].

Fusaoka et al. recently identified several vertebrate neural cell adhesion molecule orthologs in planarian Dugesia japonica that when knocked-down using RNAi altered neural network morphology [91]. Our analysis of the planarian genome revealed homologs for several Ig-like domains typically found in adhesion molecules, integrin-related sequences, C-type lectin domains and Cys-FGFR domains. Some of these domains were upregulated post-amputation and remained mostly elevated at 30 min to 72 h [30]. C-type lectin domains are associated with immune responses, cell adhesion and cell death [90]. Studies have shown that invertebrate homologs have an abundance of C-type lectin-like domains [92]. C-type lectin receptors play a crucial role in pathogen recognition and could mediate functions associated with cell adhesion. We found several homologs of C-type lectin domains (mk4.010284.01, mk4.000424.08, mk4.000389.11) that were upregulated during regeneration in planaria [30]. Analysis of the planarian genome also identified homologs for 8, 11 and 15 cadherins and for E-Cadherin. Several of these homologs were upregulated during regeneration. However, the specific function these proteins during planaria wounding and regeneration or in host defense has not been evaluated.

3.6. Nitric oxide

Nitric oxide is utilized in many animal species as a cell signaling and cytotoxic molecule as it diffuses readily through fluids and across cell membranes [93]. It is synthesized from l-arginine by the enzyme nitric oxide synthase in response to calcium-dependent and independent signals. Apart from pathogen defense nitric oxide stimulates clearing of cellular debris and the activation, growth and death of immune cells [94,95]. A homolog for nitric oxide synthase has been discovered in several flatworm species [96]. Adh3 an evolutionarily conserved gene that encodes the enzyme alcohol dehydrogenase, is another protein involved in nitric oxide metabolism that has been identified in planaria [97]. The existence of a nitric oxide synthase homolog and an alcohol dehydrogenase suggests another host defense mechanism by which planarians might control pathogens. Nitric oxide also has a documented role in mammalian wound repair. In rats fed with an arginine-free diet (arginine suppresses nitric oxide production) wound healing was impaired [98], and in patients nitric oxide levels are low in diabetic sores [95]. Nitric oxide may represent another evolutionarily conserved example with functions in both host defense and wound repair.

3.7. Double strand RNA (dsRNA)

RNAi technology is widely used in planarian research. dsRNA and miRNA have been implicated biologically as important transcription regulators that mediate many signaling pathways during development, repair, regeneration and maintenance of homeostasis [99]. Existence of dsRNA creates red flags within eukaryotic cells, indicating viral infection [100,101]. Eukaryotic cells have evolved with specific rapid and successful mechanisms to remove dsRNA as foreign material. Scientists have manipulated this knowledge to successfully downregulate genes and their functions from cells. Within an eukaryotic organism this RNAi silencing pathway can be used not only for the removal of the viral bi-products, but for changes in gene expression levels and subsequent signaling pathways that might play a crucial role in immune functions. Analysis of the planarian genome revealed homologs for Dicer and AGO2 (mk4.000678.05 and mk4.001736.00), however these genes were not upregulated during regeneration in the planaria [30] implying this mode of immune regulation might not be important in wound repair and regeneration.

3.8. Metalloendopeptidases

Metalloendopeptidases are highly conserved across evolution and are ubiquitously expressed [102,103]. Matrix metalloproteinases (MMPs) in mammals play a critical role in extracellular matrix remodeling and thus wound repair. There are 24 mammalian MMPs, some that may also have a role in immune responses, activating cytokines and chemokines to induce inflammation and immunity [104,105]. Injury induces the expression of MMPs, and these perform multiple, distinct functions in wound repair, cell migration and bacterial inhibition and killing [106]. MMP-like proteins have been identified in a diverse array of invertebrates [107,108]. Planarians have four MMP-like genes (Smed-mmp1, Smed-mmp2, Smed-mt-mmpA and Smed-mt-mmpB) with roles in proliferation, apoptosis and cell migration [109]. Two of these, Smed-mmp-1 and Dj-mmp-1, are essential for homeostatic remodeling of extracellular matrix, but not wound repair [109]. Although gene expression was unchanged following wounding, knock-down of Smed-mt-mmpA resulted in a delay in blastema regeneration [109]. Smed-mmp1 was also found to be expressed during septic-wounding in planaria, suggesting a role for this MMP in innate immune responses in addition to structural homeostasis [63]. A detailed analysis of MMP-like gene activity in planaria should be evaluated in response to injury and infection as these proteins have potential to be important in both biological activities.

4. Concluding remarks

The presence of molecules associated with innate immune response in planarians along with their activation during regeneration implies that the injury-mediated immune response is an ancient feature of basal Bilateria. Specific regulatory mechanisms of the immune system during planarian regeneration remain to be addressed. In addition to the potential vertebrate immune homologs that we identified, it is unknown if other non-vertebrate immune mechanisms exist in planarians for fighting pathogens. It is possible that further analysis of the injury-induced immune response may reveal invertebrate-specific responses as part of their ancient immune defense mechanisms. Thus, the planarian model provides great opportunities to evaluate evolutionary mechanisms and studies of molecular players during tissue regeneration and host defense that are difficult to interrogate in other experimental models. Planarian studies mostly focus on the adult organism, which facilitate analysis of local and systemic signals in the presence of a fully developed immune system. RNAi, transcriptomic and proteomic approaches could be used to categorize components of the immune system modulating particular phases of wound healing, blastema formation and tissue remodeling (Fig. 3). Critical aspects such as the role of commensal bacteria during tissue repair and the specific role of primitive phagocytic cells (reticulo-cyes) could be investigated in the context of the whole organism. The absence of an adaptive immune system in planarians presents some limitations within the model for comparison to human mechanisms. However, comparative studies between planarians and salamanders evaluating scar-free repair and regeneration [9] may point toward common factors within the innate immune system with potential clinical applications.

Evaluation of healing in autoimmune patients suggests that a dysregulated inflammatory response delays or impairs the wound repair process [110]. It thus appears that a proper balance of immune response must be maintained during wound repair. The data seem to suggest that too strong of an immune response (or
Injury to planarian tissue

Mechanical Contraction
Cellular Responses

Immune Inducing Signals

Neoblasts
Reticular Cells

Overlap??

Fig. 3. Injury induces immune and repair signaling in planarians. Following injury or amputation, molecular signals induce proliferation of neoblasts and formation of the blastema. Reticular cells migrate into the injury site and can be found with phagocytosed dead tissue. Septis-injury induces expression of potential host defense molecules [63]. As immune-inducing and regeneration signaling events likely occur simultaneously, it is possible that some of these signals impact both biological pathways. In addition, immune components alter the rate of regeneration and potentially impact the structural outcome of the tissue reconstruction.

perhaps a more developed immune system) may contribute to scarring during wound repair and reduced regenerative capacity, while a lower level of inflammation (or more primitive immune signaling) may be critical in ensuring that the appropriate cell types migrate into the damaged environment and contribute to the clean-up of localized debris and dead cellular tissue (Fig. 3) [6,7,5,11]. This information could be readily addressed by analyzing planarian regeneration and their enigmatic immune system.

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