Mechanisms of innate immunity in C. elegans epidermis
Clara Taffoni, Nathalie Pujol

To cite this version:
Clara Taffoni, Nathalie Pujol. Mechanisms of innate immunity in C. elegans epidermis. Tissue Barriers, Taylor & Francis, 2015, 3 (4), pp.e1078432. 10.1080/21688370.2015.1078432. hal-01768347

HAL Id: hal-01768347
https://hal-amu.archives-ouvertes.fr/hal-01768347
Submitted on 17 Apr 2018

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Distributed under a Creative Commons Attribution - NonCommercial 4.0 International License
Mechanisms of innate immunity in C. elegans epidermis

Clara Taffoni & Nathalie Pujol

To cite this article: Clara Taffoni & Nathalie Pujol (2015) Mechanisms of innate immunity in C. elegans epidermis, Tissue Barriers, 3:4, e1078432, DOI: 10.1080/21688370.2015.1078432

To link to this article: https://doi.org/10.1080/21688370.2015.1078432

© 2015 The Author(s). Published with license by Taylor & Francis Group, LLC
Clara Taffoni and Nathalie Pujol

Published online: 05 Oct 2015.

Article views: 1110

View related articles

View Crossmark data

Citing articles: 7 View citing articles
Mechanisms of innate immunity in C. elegans epidermis

Clara Taffoni and Nathalie Pujol*
Center d’Immunologie de Marseille-Luminy; Aix Marseille Université UM2; Inserm; Marseille, France

Keywords: collagen, cuticle, damage, Epidermal barrier, epidermis, GPCR, hemidesmosome, pathogen, Pathogen-barrier interactions, signaling, wounding

The roundworm C. elegans has been successfully used for more than 50 y as a genetically tractable invertebrate model in diverse biological fields such as neurobiology, development and interactions. C. elegans feeds on bacteria and can be naturally infected by a wide range of microorganisms, including viruses, bacteria and fungi. Most of these pathogens infect C. elegans through its gut, but some have developed ways to infect the epidermis. In this review, we will mainly focus on epidermal innate immunity, in particular the signaling pathways and effectors activated upon wounding and fungal infection that serve to protect the host. We will discuss the parallels that exist between epidermal innate immune responses in nematodes and mammals.

Introduction

A large body of knowledge has been accumulated for the tiny worm Caenorhabditis elegans since it was chosen as a model organism by Brenner 50 y ago.1 C. elegans, first described by Maupas at the start of the last century, 2 is a free-living nematode, 1 mm long, found in rotting fruit. In the lab it can easily be grown on agar petri dishes or in liquid culture. Being a self-fertilizing hermaphrodite, it can be expanded as a clonal population, grown on agar petri dishes or in liquid culture. Being a self-fertilizing hermaphrodite, it can be expanded as a clonal population, grown on agar petri dishes or in liquid culture. With a short life cycle of 3 days, and 300 progeny per adult, large homogenous populations can be generated easily. Its transparency facilitates live imaging, very useful for cell biological studies. C. elegans has a simple anatomy with less than 1000 cells organized in a small number of tissues and internal organs.3

In C. elegans, as in mammals, the epidermis acts as a barrier to protect the organism from environmental damage and pathogens, prevents leakage of internal molecules and blocks the entrance of foreign compounds. Unlike the multilayered stratified epithelium in mammals, the adult C. elegans epidermis (also termed “hypodermis” for historical reasons) is mainly composed of one cell layer, including a large syncytial cell, called hypodermal cell 7 (hyp7), which surrounds the worm and covers most of the body length, and a line of specialized lateral cells on each side of the worm, called the seam cells (Fig. 1).4

As in most invertebrates, C. elegans possesses an exoskeleton. In nematodes, this is a tough, but flexible external cuticle secreted by the epidermis. This collagenous extracellular matrix (ECM) maintains the integrity of the worm, defines the body shape of the animal and is required for locomotion, through its attachment to the underlying muscles. The cuticle is subdivided into basal, medial and cortical zones, overlaid by the epicuticle and the most external layer, the surface coat. While the cuticle is mainly composed of collagen and insoluble proteins, called cuticlins, the epicuticle and the surface coat are rich in lipids and structural glycoproteins such as mucins, respectively. Secreted non-structural proteins also likely make up an important part of the cuticle, such as enzymes involved in post-secretion modification and cross-linking of matrix proteins or structural proteins associated with the surface coat.5,6 In mammals, the most external layer of the skin, the stratum corneum (SC) is the final product of keratinocyte differentiation, resulting from denucleation and crosslinking of intracellular proteins. It is mainly composed of keratin, cholesterol, free fatty acids and ceramides.7 Although differing in composition, the C. elegans cuticle can be considered analogous to the SC as they both function as a permeability barrier.8

Cuticle Collagen: A Key Component in Barrier Integrity

Collagen is the main structural protein of the extracellular matrix in animals, and the most abundant protein in mammals. It is an essential component of the skin and plays a key role in organogenesis and body morphology. In C. elegans, 2 major collagen families are present, the cuticle and the basement membrane collagens. While only 3 genes encodes type IV and type XVIII basement membrane collagens, the cuticle collagens are encoded by more than 170 genes. They are most similar to

*Correspondence to: Nathalie Pujol; Email: pujol@ciml.univ-mrs.fr
Submitted: 04/29/2015; Revised: 07/17/2015; Accepted: 07/24/2015
This is an Open Access article distributed under the terms of the Creative Commons Attribution-Non-Commercial License (http://creativecommons.org/licenses/by-nc/3.0/), which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited. The moral rights of the named author(s) have been asserted.

© Clara Taffoni and Nathalie Pujol
www.tandfonline.com
the fibril-associated collagens with interrupted triple helices (FACIT) found in vertebrates. They are regulated differentially, most of them following the 4 molting cycles during larval development and are involved in the formation of distinct sub-cuticle structures (e.g., annular furrows and alae) and layers. Collagen biogenesis involves a complex maturation process, partially shared between nematodes and vertebrates, during which procollagen undergoes several co-translational and post-translational modifications. Collagen maturation ends with the secretion of collagen triple helices that are covalently cross-linked via tyrosine residues. This determines the characteristic stiffness and integrity of the worm cuticle. During this step, a peroxidase (MLT-7) cooperates with an NADPH dual oxidase (BLI-3), to promote reactive oxygen species (ROS)-mediated catalysis of di- and tri-tyrosine covalent bonds. Recently, a tetraspanin protein (TSP-15) was found to form a complex with BLI-3/DUOX and MLT-7/peroxidase and to be required for ROS generation. In the absence of any one of these 3 proteins, no di- and tri-tyrosine bonds are made and the cuticle is fragile and blistered. In vertebrates, the formation of covalent cross-links of proteins such as involucrin and loricrin in the SC is also essential for the function of mammalian skin as an external barrier. It occurs via the action of transglutaminases that lead to the oxidation of amino group of lysine or glutamate residues into aldehyde derivatives. Collagen helices are cross-linked by lysyl oxidase in vertebrates between lysine and hydroxyllysine residues. Lysyl oxidase-mediated cross-linkages are not found in the C. elegans cuticle, they only contribute to cross-linking of type IV collagen in the basement membrane.

In human skin, collagens are mainly found in the dermis. Many mutations in different collagen genes have been associated with various diseases. For example, patients with epidermolysis bullosa, carry mutations in COL7A1 or COL17A1 that lead to blisters in the skin and mucosal membranes, and mutations in COL3A1 or COL5A1 have been found in patients with Ehlers–Danlos syndrome, with various symptoms including a fragile skin. Similarly, in nematodes, mutations in genes encoding collagens, like DPY-9, ROL-6, BLI-1, or processing enzymes, including the prolyl 4-hydroxylase DPY-18 or the disulfide isomerase PDI-2, result in diverse body morphology defects described as ROLler (Rol: helical twisting of the animal’s body), DumPY (Dpy: shortening in the body length), or BLIster (Bli: blistering of cuticle material away from the surface of the animal).

The nematode cuticle is made of successive layers of different collagens. Interestingly, some mutants in specific collagen genes, such as dpy-2, dpy-7 or dpy-10, which are presumed to be the most external collagens of the cuticle, show high osmotic resistance, due to a higher internal level of glycerol. These same collagens are also required for proper circumferential furrow formation. Wheeler and Thomas proposed that these collagens on the furrows could act as a stretch sensor, monitoring the turgor pressure of the cuticle. OSM-7 and OSM-11, which are nematode-specific proteins structurally related to the Notch ligand Delta, possibly associated with the cuticle, could transduce the signal, leading to the increase in glycerol. The same mutants also show a constitutive activation of innate immunity (see below). As previous studies found these mutants to be involved in the suppression of diverse hypomorphic mutations,

![Figure 1. C. elegans anatomy. (A) Image of an early larval stage in DIC, scale bar is 20 μm. Schematic of an adult C. elegans body (B) and cuticle (C). The adult cuticle is approximately 0.5 μm in thickness and subdivided in (Bs) basal zone, (Md) medial zone, (Co) cortical zone, (Ep) epicuticle and (Sc) surface coat. Collagen is present in all the major layers, except for Ep and Sc which are rich in lipids and glycoproteins, respectively. The cuticle is synthesized by the epidermis, formed by the hypodermis (hyp), a simple epidermal syncitium and the lateral seam cells (adapted from Wormatlas, http://www.wormatlas.org/hermaphrodite/cuticle/mainframe.htm).](image-url)
probably through the induction of chaperone function,\textsuperscript{21-24} these results suggest that some collagens are part of a general stress sensing mechanism.

**Epidermal Wounding: Cell Autonomous Innate and Healing Response**

Just as mutation of genes required for the proper development of the worm’s cuticle can trigger an innate immune response, so too does directly wounding the worm. This can be done either by hand, with a fine glass needle, when the worm is pricked at the tail, or in a more controlled manner with a laser that enables precise wounding of the epidermis. In both cases, at the wound site there is an immediate deposition of as-yet uncharacterized autofluorescent material that persures for many hours. Later, the secretion of basal cuticle components leads to the repair of the damaged cuticle, leaving a scar at the wound site. Moreover, injury provokes a rapid induction of antimicrobial peptide (AMP) gene expression in the epidermis. For example a rise in the expression of nlp-29 is detectable within one hour after injury and is sustained for several hours (Fig. 2A).\textsuperscript{20} This induction of innate effectors upon wounding is not due to microbe invasion but is a preventive response of the tissues, as previously shown in other species.\textsuperscript{25} Indeed, mutants that are defective for the induction of defense gene expression upon wounding succumb to opportunistic infections.\textsuperscript{26}

In order to identify the first event which initiates the epidermal response to damage, Xu and Chisholm focused their attention on calcium signaling, as it was already known to be involved in embryonic wound healing and the response to membrane injury in single cells.\textsuperscript{27,28} Upon injury to the epidermal syncytium, an immediate Ca\textsuperscript{2+} wave, released from internal stores, can be detected at the wound site and provokes local formation of a dense actin ring which, in turn, will trigger complete wound closure in 24 hrs. This epidermal Ca\textsuperscript{2+} response involves the epidermal transient receptor potential channel, metastasin family (TRPM) GTL-2, the G\textalpha_q EGL-30 and its effector PLCβ EGL-8.\textsuperscript{29} Subsequently it was shown that specific Ca\textsuperscript{2+} uptake into the mitochondria close to the wound site leads to local mitochondrial ROS (mtROS) production and that mtROS release inhibits the Rho GTPase RHO-1, promoting actin polymerization and filopodial-like protrusion in a Cdc-42 and Arp2/3 dependent manner.\textsuperscript{25,30} This wound closure process is different from the
most common Rho GTPase-dependent purse-string mechanism that occurs, for instance, after wounding in the *Drosophila* embryo.31

While the pathways leading either to wound healing or AMP production (see below) are genetically and mechanistically distinct,29 the worm homolog of the tumor suppressor death-associated protein kinase, DAPK-1, functions as a negative regulator of both barrier repair and AMP expression. Loss of function of *dapk-1* leads to spontaneous cuticle hypertrophy with associated appearance of autofluorescent aggregates and to the constitutive upregulation of epidermal AMPs.26 Thus as for other immune responses, the epidermal response to damage has to be tightly regulated to prevent the deleterious consequences of chronic activation.

**Pathogen Infection**

While some pathogens will only passively enter the host through a wound, others have developed active ways to infect the worm through the cuticle. The nematophagous fungus *Leucinillium pudiliiata* is able to degrade the worm cuticle by secreting a serine protease, thus facilitating mycelia penetration and complete nematode degradation.32 Other fungi, such as *Arthrobotrys oligospora*, *Drechslerella stenobrocha* and *Dactylella ellipsoara*, are able to form traps to capture, kill and digest worms as a food source.33 One of the most studied pathogens able to infect the worms via its cuticle is *Drechmeria coniospora*. This endoparasitic fungus is frequently found associated with nematodes in nature.34,35 Its conidia (non-motile spores) can bind to the worm’s cuticle through specialized adhesive knobs and upon germination spread hyphae throughout the worm’s body, causing death after as little as 48 hours.36 Some bacteria also adhere to the cuticle, like *Leucobacter spp* Verde37 or *Microbacterium nematophilum* that colonizes the rectal cuticle and provokes swelling in the adjacent epithelia.38,39

Fungi and bacteria take advantage of the carbohydrate-rich surface coat and are believed to use lectins to adhere to the nematode cuticle. Recent studies in *C. elegans* have identified mutants with defects in their surface coat carbohydrates and an increased adhesion of *D. coniospora*. For instance, the *bus-2* mutant carries a mutation in the enzyme galactosyltransferase and is enriched in core-2 fucosyl O-glycans at the expense of core-1 glycans, suggesting that there is an increase in fucose exposure in this mutant.40,41 Moreover a fucose-specific lectin binds the vulva of wild-type worms, a preferential site for *D. coniospora* adhesion. Taken together, these results suggest that fucosyl glycans are a target for the binding of *D. coniospora* spores.42

It is interesting to note that if mutants like *bus-2* have an increased adhesion to *D. coniospora*, they conversely have a reduced adhesion to the bacterial pathogen *M. nematophilum*, hence their name *Bus* for bacterial unswollen.43 The same opposite effect has been revealed with 2 different strains of *Leucobacter spp* Verde. Thus, resistance to some natural pathogens can be associated with increased susceptibility to others, revealing the opposing selective forces that must shape the evolution of *C. elegans*.44

The surface coat composition changes during development through the different larval stages and it was also shown to change upon environmental signals,44,45 which could be presages, for instance, of the presence of pathogens. Thus, it is reasonable to think that this switch could enable evasion of pathogens or, in case of the parasitic nematodes, escape of the host immune response.46 This would complement the purely mechanical shedding of pathogens that can accompany molting.46

**AMP Induction in *C. elegans* Epidermis**

If pathogens have developed ways to attach to the cuticle, it is now clear that the worm is able to mount an innate immune response after infection. Using cDNA and genomic DNA microarrays or more recently RNaseq, transcriptomic profiling has identified many genes induced upon infection with *Monacrosporium haptostrymum*47 or *D. coniospora*.19,48-50 Among this latter class, some were predicted to encode AMPs, including 2 related families: a subset of neuropeptide-like proteins (NLPs) and the caenacins (CNCs).48,51 Members of these 2 families are present in 2 distinct genomic clusters: the *nlp-29* (including *nlp-27*, *nlp-28*, *nlp-29*, *nlp-30*, *nlp-31* and *nlp-34*) and the *cnc-2* (including *cnc-1*, *cnc-2*, *cnc-4*, *cnc-5*, and *cnc-11*) clusters, which contribute to resistance to *D. coniospora* infection.19,52,53 While these 2 classes of genes are upregulated by infection and act in the epidermis, they are regulated by different signaling pathways (see below, Fig. 2B). The main immune players were mostly identified through genetic screens, through the selection of mutants, termed Nipi, which fail to express antimicrobial peptide genes after *D. coniospora* infection.20

**A Cell Autonomous Response via the p38 MAPK Pathway**

After *D. coniospora* infection or wounding, the expression of *nlp* genes is induced by the activation of a G-protein-coupled receptor (GPCR) DCAR-1 in an epidermis-specific and cell-autonomous way (Fig. 2B). DCAR-1 is expressed in the apical part of the epidermal syncytium (hyp7) and is activated by HPLA (4-hydroxyphenyllactic acid). The level of this ligand is increased to resistance after infection, and also in mutants with defects in the cuticle like dpy-10. Thus, it seems that HPLA acts as a DAMP (damage-associated molecular pattern). Even though *dcar-1* orthologs are only found in nematodes, these findings point to an ancient role for GPCR in innate immunity.54 Interestingly HPLA, a common tyrosine metabolite, is detected across all species and its elevated concentration was detected in patients with phenylketonuria (PKU), tyrosinemia or with Zellweger’s syndrome.55-57

DCAR-1 acts upstream of the Gα protein GPA-12 and of the Gβ protein RACK-1 (orthologous to the mammalian Gα G12 and similar to the human Gβ-2-like 1, respectively), which in turn, act genetically upstream of the serine, threonine Protein Kinase C (PKC) TPA-1, whose activation is diacylglycerol (DAG) dependent. DAG is a second messenger produced from
phosphatidylinositol biphosphate (PIP2) by the catalytic action of the phospholipase C (PLC) EGL-8 and PLC-3 (a PLCγ homolog).54,58

In mammals and insect, a family of adaptor proteins containing a domain named TIR, for Toll Interleukin Receptor, appears to be crucial for Toll Like receptor (TLR) signaling.59 C. elegans has only 2 genes encoding proteins with a TIR domain, the TLR tol-1 and the SARM adaptor tir-1. Upon fungal infection, AMP induction is tol-1 independent but tir-1 dependent.48 tir-1 acts upstream of the nlp-1/nls-1/nlp-1/pmk-1 p38-MAPK pathway to induce epidermal AMP expression20 and it is required for full resistance to infection.48 Both the STAT family protein STA-2 and the GATA transcription factor ELT-3 are required for the transcription of nlp23,66 (Fig. 2B).

Parallel to this pathway, a conserved Tribbles-like kinase, NIPI-3, has been found to act upstream of TPA-1 and the p38/MAPK cassette specifically upon D. coniospora infection.20,58 The tribbles family of pseudokinases are key controllers of signal transduction via their interactions with diverse kinases, ubiquitin ligases and transcription factors. Early studies in the fly also associated Tribbles-like kinase with the control of cell cycle during development.61 It will be interesting to test if NIPI-3/TriB has also a general function in cell cycle regulation in C. elegans.

A Paracrine role of Non-Canonical DBL-1/TGFβ Pathway

cnc gene expression is induced both after D. coniospora infection and injury, but is controlled in a complex way depending on the stimulus (Fig. 2B). It was shown that the induction of the cnc-2 gene cluster is p38-MAPK dependent after wounding, while it is p38-MAPK independent upon infection. After infection, cnc genes are upregulated by a non-canonical DBL-1/TGFβ pathway.53

The canonical DBL-1/TGFβ pathway was previously shown to be involved in regulation of body size and male tail development in C. elegans. DBL-1 binding to the heterodimeric DAF-4/ SMA-6 receptor activates the intracellular Smad signal-transducer homolog SMA-2/SMA-3/SMA-4 complex which, in turn, will control gene expression through recruitment of the zinc-finger transcription factor, SMA-9, in the nucleus.62 If the entire SMA-2/SMA-3/SMA-4 complex is needed also for a response to P. aeruginosa,63 instead, only SMA-3 is required for a response to D. coniospora. The identity of the transcription factor involved in this process is still unclear (Fig. 2B).53 Strikingly, the DBL-1 ligand is mainly expressed in the nervous system, so it might act in a paracrine way to induce cnc gene expression in the epidermis,53 in addition to controlling defense gene expression in the intestine.64 It’s still unknown, however, how the signal is transmitted from the neurons to the epidermis. It’s possible that DBL-1-secreting neurons are directly in contact with the pseudo-coelom, suggesting that DAF-4/SMA-6 receptor could be activated from the basal side of the epidermis.46

Role of Solute Carrier Family (SLC) and Pseudokinases in Immunity

C. elegans genetic screens highlighted also the presence of unexpected players in these innate immune pathways, such as the sodium-neurotransmitter symporter SNF-12, a member of the solute carrier family (SLC6). Interestingly, SNF-12 controls the constitutive expression of cnc genes and nlp gene induction after infection and wounding. SNF-12 physically interacts with the transcription factor STA-2 and they have been found to colocalise in dyn-1-dependent-endocytic vesicles through which they regulate part of the innate immune response (Fig. 2B).60 SNF-12/SLC6 is predicted to be a transporter of a bioactive amine, but its endogenous substrate has yet to be identified.

Studies in vertebrates revealed that another class of SLC protein, transporting oligopeptides, might have a crucial function for the antibacterial or antiviral activity of macrophages, mature dendritic cells (mDCs)65 and plasmacytoid dendritic cells (pDCs).66 In mouse mDCs and macrophages, Slc15a mediates the internalization in late endosomes/lysosomes of fragments of peptidoglycan coming from bacteria cell wall, and stimulates NF-kB and other immune effector pathways.65 Moreover, it was shown that the peptide/histidine transporter Slc15a4 is present both in mouse pDCs66 and B cells lysesomes where, by maintaining the appropriate pH, it can promote optimal TLR7 and TLR9 activation.67 Interestingly, Slc15a4 is associated with systemic lupus erythematosus (SLE).68

Another player genetically interacting with STA-2 and SNF-12 in C. elegans is the pseudokinase NIPI-4 (Fig. 2B).69 Interestingly, even though pseudokinases lack the ability to phosphorylate substrates, they are still able to regulate cellular processes, forming heterodimers with other protein kinases and controlling their activity.70 For instance, in humans, the pseudokinase Ste20 Related ADaptor (STRAD)α, with its closed conformation typical of active protein kinases, can bind the tumor suppressor protein kinase LKB-1, induce a conformational change in LKB-1, which, in turn, enables cell proliferation, and regulates cell polarity as well as energy levels.70,71 NIPI-4 is expected to have a similar regulatory role. Although the identity of its target kinase is unknown, the p38 MAPK PMK-1 is a prime candidate.

Is MechanoSensation Involved in Innate Immunity Induction?

It is still not clear how p38 MAPK-dependent STA-2 activation and trafficking are regulated in C. elegans epidermis. A very recent study suggests that in the case of severe epidermal injury, upregulation of nlp-29 and cnc-2 could occur in a p38 independent, but STA-2 dependent way.72 STA-2 was found in the nucleus but also associated with hemidesmosomes (CeHDs),60,72 structures responsible for the attachment of the epidermal cells to the cuticle.73,74 Zhang et al. showed that STA-2 physically interacts with one apical component of CeHDs, MUP-4 a transmembrane multidomain protein. They hypothesized that the interaction of STA-2 with CeHDs inhibits its activity and that
severe damage or CeHDs disruption would free STA-2 to induce AMP production. They further propose that this mechanism is conserved in humans where β-defensin transcription correlates with disruption of hemidesmosomes protein complexes (HPC) and inactivation of STA-3 or STA-5B attenuates AMPs upregulation after HPC disassembly in human epidermal keratinocyte (HEKa) cells. The suggestion that hemidesmosomes will act as a damage-sensor raises very interesting questions, like how do the HDs perceive the damage? And what are the signals that will activate HDs? Some hints have come from studies of epidermal development.

In *C. elegans*, it was known that muscle contraction is required in the embryo during morphogenesis for epidermal elongation. From a recent study, it was shown that upon muscle contraction p21-activated kinase (PAK-1), a component of CeHDs, is activated and together with the adaptor GIT-1 (G-protein-coupled receptor kinase interactor) and its partner PIX-1 (PAK-interacting exchange factor) can phosphorylate epidermal intermediate filaments and promote CeHD biogenesis. In mutants defective for muscle contraction, GIT-1 progressively leaves HDs. Zhang et al. could show that applying external mechanical pressure on these mutants considerably retarded the diffusion of GIT-1. Thus, they demonstrated that CeHDs are mechanosensitive and direct targets of physical forces. In the future it would be of interest to address whether injury or spore adhesion could induce a mechanical signal that could be sensed by apical CeHDs.

Interestingly, there is evidence suggesting a role for collagens in HDs integrity. In humans, mature type-1 HDs contain the bullous pemphigoid (BP) antigens BP180/collagen XVII as well as integrin α6β4 and plectin. Mutations in the corresponding genes in patients with epidermolysis bullosa alter hemidesmosomal structure. Also during carcinogenesis, extracellular cleavage of collagen XVII by matrix-metalloprotease MMP-9 disrupts HDs. In *C. elegans*, worms with mutations in collagen genes, such as *dpy-9* and *rol-6*, present constitutive upregulation of AMPs. As the extracellular portion of MUP-4, the apical CeHD component, contains a von Willebrand factor A domain for collagen binding, it is possible that MUP-4 acts as a regulator of the epidermal immune response conceivably sensing, after injury or spore adhesion, collagen disruption.

References

1. Brenner S. *The genetics of Caenorhabditis elegans*. Genetics 1974; 77:71-94; PMID:4366476
2. Maupas E. Modes et formes de reproduction des nematodes. *Archives de Zoologie Experimentale et Generale* 1900(8):463-624
8. Xu S, Huiao TH, Chisholm AD. The wounded worm: Using *C. elegans* to understand the molecular basis of skin wound healing. *Worm* 2012; 1:134-8; PMID:24058838

Conclusions

*C. elegans* lacks both an adaptive immune system and specialized immune cells. This obviously limits its relevance for the understanding of human immunity. On the other hand, its relative simplicity and the extensive range of available experimental methods facilitate its use in biology. The major molecular mechanisms involved in the epidermal innate immune response in *C. elegans* have been identified. As in humans, they involve the upregulation of AMP expression and wound healing, governed by a combination of highly conserved and nematode-specific pathways. A common theme that has emerged is the importance of damage recognition as a trigger for the induction of defense genes. Future studies should reveal the mechanisms leading to the production and/or release of DAMPs and the identity of the elusive “fungus-receptor” putatively responsible for triggering pathogen-specific innate immune defenses. Together with a better understanding of how cross-tissue communication contributes to the regulation of host immunity, this will lead to a more comprehensive insight into how an organism can re-establish homeostasis after the integrity of epithelial barriers has been compromised by injury or infection.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Acknowledgments

We thank Jonathan Ewbank for critical reading.

Funding

Work in our lab is supported by institutional funding from INSERM, CNRS and AMU, and program grants from the Research National Agency ANR-12-BSV3–0001–01 (FUNEL) and ANR *Investissements d’Avenir* ANR-11-LABX-0054 (Labex INFORM) ANR-10-INSB-04–01 (France-BioImaging) and ANR-11-IDEX-0001–02 (A*MIDEX)).
30. Xu S, Chisholm AD. C. elegans epidermal wounding induces a mitochondrial ROS burst that promotes wound repair. Dev Cell 2014; 31:48-60; PMID:25313960; http://dx.doi.org/10.1016/j.devcel.2014.08.002
www.tandfonline.com Tissue Barriers e1078432-7