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Abstract The study of molluscan immune systems, and in particular those of bivalve molluscs (clams, oysters, scallops, mussels, etc.) has experienced great growth in recent decades, mainly due to the needs of a rapidly growing aquaculture industry to manage the impacts of disease and the application of -omic tools to this diverse group of invertebrate organisms. Several unique aspects of molluscan immune systems highlighted in this chapter include the importance of feeding behavior and mucosal immunity, the discovery of unique levels of diversity in immune genes, and experimental indication of transgenerational immune priming. The development of comparative functional studies using natural and selectively bred disease-resistant strains, together with the potential but yet to be fully developed application of gene-editing technologies, should provide exciting insights into the functional relevance of immune gene family expansion and molecular diversification in bivalves. Other areas of bivalve immunity that deserve further study include elucidation of the process of hematopoiesis, the molecular characterization of hemocyte subpopulations, and the genetic and molecular mechanisms underlying immune priming. While the most important aspects of the immune system of the largest group of molluscs, gastropods (e.g., snails and slugs), are discussed in detail in Chap. 12, we also briefly outline the most distinctive features of the immune system of another fascinating group of marine molluscs, cephalopods, which include invertebrate animals with extraordinary morphological and behavioral complexity.

Keywords (separated by “ - ”) Mollusca - Bivalves - Oyster - Mussel - Scallop - Clam - Cephalopods - Aquaculture - Lectins - Opsonization - Pathogen recognition - Immune signaling - Antimicrobial peptides - Apoptosis - Complement system - Immune priming - Phagocytosis - Prophenoloxidase - Neuroendocrine immunomodulation

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Q4 This chapter has been lightly edited for clarity, correctness and house style. Please check it carefully to make sure the intended meaning has been preserved. If the intended meaning has been inadvertently altered by the editing changes, please make any corrections needed.

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[AU2]	Marco Gerdol, Marta Gomez-Chiarri, Maria G. Castillo, Antonio Figueras, Graziano Fiorito, Rebeca Moreira, Beatriz Novoa, Alberto Pallavicini, Giovanna Ponte, Katina Rumbedakis, Paola Venier, and Gerardo R. Vasta	4
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An Introduction to Bivalve Molluscs

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Evolution and Life Cycle

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[AU5]	The phylum Mollusca includes eight taxonomic classes comprising more than	10
[AU6]	85,000 living species, and 60,000 additional species documented by fossil	11
	records (Fig. 1). This ranks molluscs as the second most abundant phylum of	12
[AU7]	animals after arthropods and before chordates (Ponder and Lindberg 2008).	13
	Molluscs are successful invertebrates characterized by a broad morphological	14

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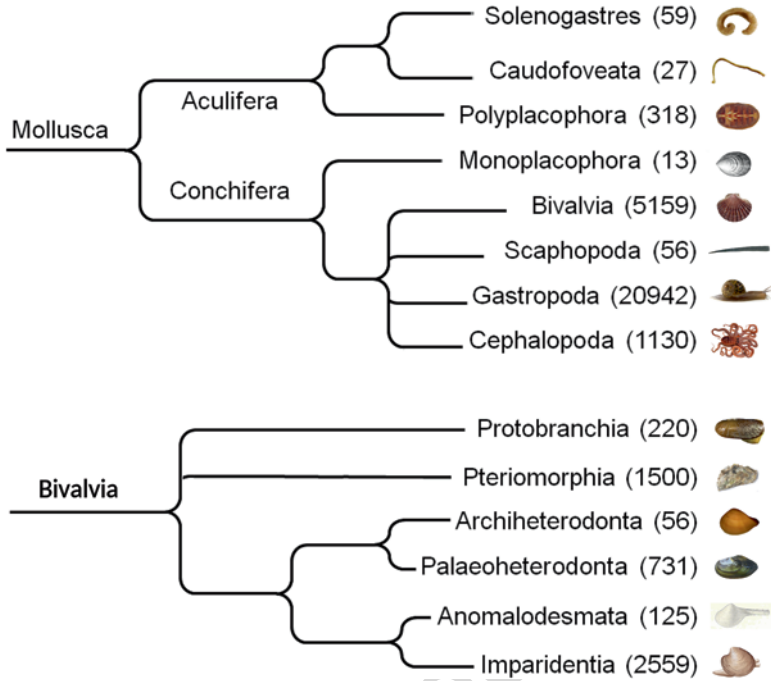


Fig. 1 Simplified tree of life of molluscs (above) and bivalves (below), based on Bieler et al. (2014) and the Tree of Life web project (<http://tolweb.org/Mollusca/2488>). The number of species currently registered in the NCBI Taxonomy database for each taxon (data retrieved in December 2017) is displayed between brackets

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and physiological diversity. They are extraordinarily well adapted to adverse environmental conditions and, starting from the early radiation that occurred in the Late Cambrian era, they have colonized almost all ecological niches: from terrestrial habitats over 3000 meters above sea level to deepsea hydrothermal vents, coping with extreme levels of heavy metals, pH, temperature, CO₂, methane, and sulfide (Plazzi and Passamonti 2010)

Bivalvia represent the second largest class within the phylum Mollusca, with over 5000 recognized species, mostly adapted to marine environments. Although the phylogenetic relationship among the different groups of bivalves and, more generally, of all molluscs have been the subject of debate for decades (Kocot et al. 2011; Smith et al. 2011; Sigwart and Lindberg 2015), recent studies tried to reorganize the bivalve tree of life into six major lineages, as shown in Fig. 1 (Bieler et al. 2014). Briefly, the authors recognized the primitive and relatively small group of Protobranchia, the large groups of Pteriomorphia (comprising oysters, mussels, and scallops, among others), Palaeoheterodonta (mostly freshwater clams and mussels), Imparidentia (the largest and most diverse group of bivalves, comprising over 2500 clam species), and two additional small groups with peculiar morphological features, i.e., Archiheterodonta and Anomalodesmata.

Bivalves can be protandric hermaphrodites (oysters in the genera *Magallana* and *Crassostrea*), simultaneous hermaphrodites (scallops in the genus *Pecten*), and rhythmical consecutive hermaphrodites (oysters in the genus *Ostrea*). As exemplified in Fig. 2, the general life history of the majority of molluscan bivalve species starts during the main spawning season when adult animals with mature gonads release oocytes and spermatozoa in the water column and external fertilization occurs (Pechenik 2010). Bivalve larvae are planktonic (free-living) and remain in the water column for days to weeks, depending on the species and the environmental conditions. During larval development, the molluscan embryo becomes a

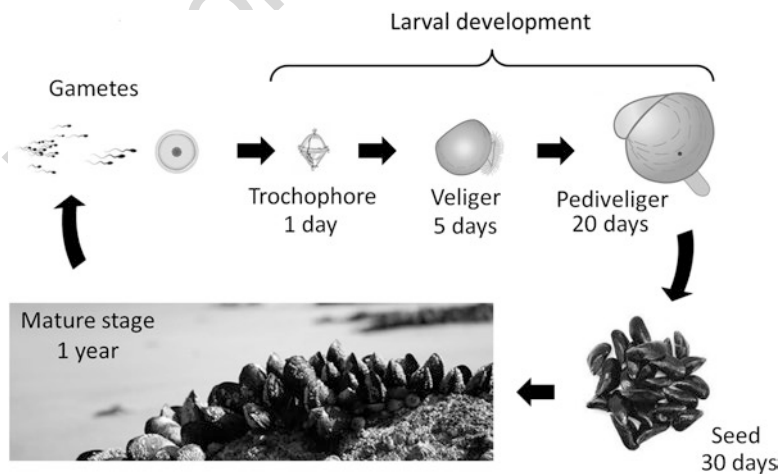


Fig. 2 Life cycle of a bivalve, as exemplified for the Mediterranean mussel, *Mytilus galloprovincialis*

42 planktonic (free swimming) trochophore larva. The late trochophore is the phylotypic
 43 stage, defined as the ontogenetic stage, characterized by maximum similarity among
 44 the species within a phylum (Xu et al. 2016a). After a few days, the primordium of
 45 the shell appears and the bands of cilia used by larvae to feed and swim develop into
 46 the velum, a characteristic organ of the veliger stage. Then, larvae develop a foot,
 47 characteristic of the pediveliger stage, and undergo metamorphosis. Once meta-
 48 morphosis is complete, their body plan and physiological aspects resemble those of
 49 the adult form and the larvae will settle out of the water column where, depending
 50 on the species, they might attach to a substrate, lie on a substrate and swim, or bury
 51 themselves in sediments (Balseiro et al. 2013). When adults become mature, gameto-
 52 genesis occurs, with modalities that depend on the species, geographic region, water
 53 depth, and season (Shumway and Parsons 2006).

54 **Anatomy and Physiology of Bivalves**

55 Although the adult anatomy of molluscs can greatly differ from one taxon to another,
 56 they share a general basic plan derived from a hypothetical shared ancestor (Fig. 3).
 57 This includes a soft oval body with bilateral symmetry, a muscular foot, a mantle—
 58 which secretes the shell (absent or internalized in some groups) or the spicules—
 59 and a feeding organ formed by chitinous sharp structures, called radula (absent in
 60 bivalves).

61 Overall, this shared body plan results in a great morphological diversity of
 62 bivalve groups adapted to different ecological niches, as shown in Fig. 4 (Ruppert
 63 et al. 2004). Bivalve shells consist of two, sometimes symmetric, hinged valves.

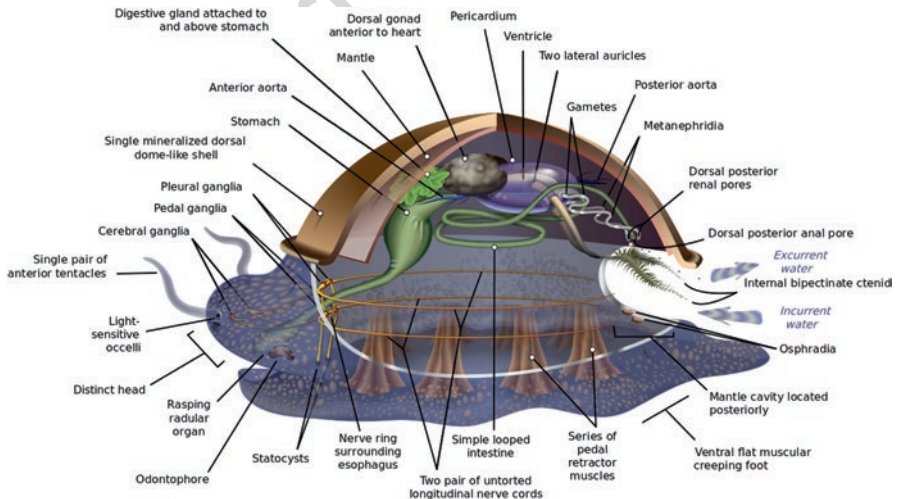


Fig. 3 Anatomy of the hypothetical common ancestor of all molluscs. (Author: KD Schroeder—Archimollusc-en.svg from Wikimedia Commons—License: CC-BY-SA 3.0)

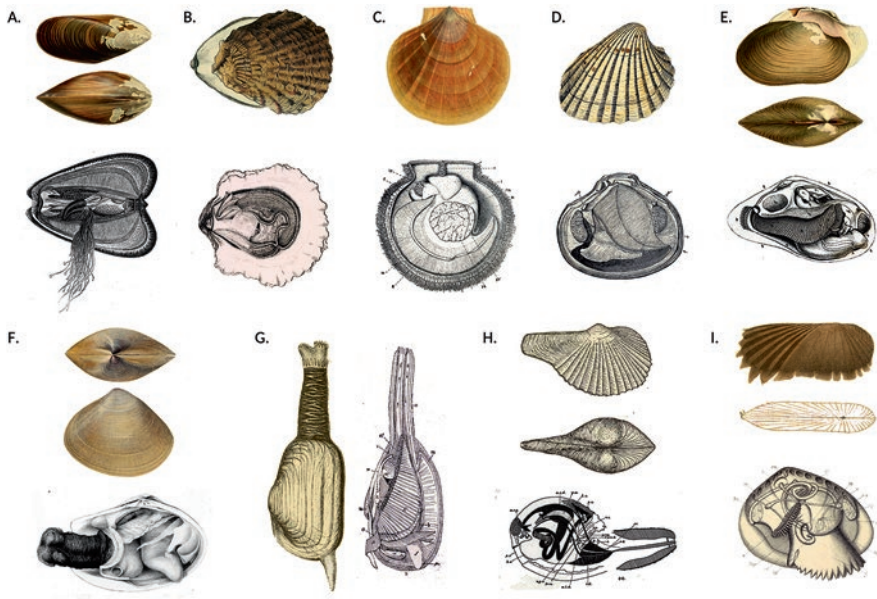


Fig. 4 Examples of diversity in the basic anatomy of different bivalve lineages. (a) Anatomy of Mytiloida (Pteriomorpha): *Mytilus uguiculatus* (external) and *Mytilus galloprovincialis* (internal). (b) Anatomy of Ostreoida (Pteriomorpha): *Ostrea edulis*. (c) Anatomy of Pectinoida (Pteriomorpha): *Placopecten magellanicus*. (d) Anatomy of Archiheterodonta: *Cardites floridanus* (external) and *Astarte borealis* (internal). (e) Anatomy of Palaeoheterodonta: *Anodonta cygnaea*. (f) Anatomy of Mactroidea (Imparidentia): *Mactra antiquate* (external) and *Tresus capax* (internal). (g) Anatomy of Myida (Imparidentia): *Mya arenaria*. (h) Anatomy of Anomalodesmata: *Cardiomya reticulata* (external) and *Laternula elliptica* (internal). (i) Anatomy of Protobranchia: *Solemya velum* (external) and *Ennucula delphinodonta* (internal). To better show anatomic internal details, in most cases one of the valves and the mantle have been removed. (The anatomic tables have been taken from multiple sources, kindly provided by the Biodiversity Heritage Library)

The shell is produced by secretory cells in the epithelium of the mantle or pallium, with contributions from the hemocytes (blood cells) (Mount et al. 2004). Bivalve shells are formed mainly of conchiolin, which is composed of protein-hardened calcium carbonate (aragonite or calcite) and has three layers: the outer layer (periostracum), a middle layer, and the inner layer, which is often nacreous and in some cases has exceptional economic value. The mantle encloses a chamber surrounding the bivalve body called the mantle or pallial cavity, which is in direct contact with the environment when the shell is open. Organs that have direct contact with the pallial cavity include the gills (or ctenidia), the osphradia (chemical sensors), and the openings of the nephridia, gonads, and digestive system. The space between the mantle and the shell constitutes the extrapallial cavity (Ruppert et al. 2004).

The movement of shell valves is controlled by one, two, or (rarely) three adductor muscles that control shell closure and keep it tightly shut when needed, and by an elastic ligament that acts as a spring, allowing the shell to open when muscles are relaxed. Some bivalves also possess a pair of siphons (inhalant and exhalant) used

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79 in the exchange of water. These systems ensure the flow of water into the pallial
80 cavity for feeding and respiration.

81 The gills divide the mantle cavity into distinct chambers and their cells possess
82 cilia, which produce a laminar flow of water that facilitates feeding and enhances
83 respiratory gas diffusion and exchange. Gills also exhibit osmoregulatory, ion trans-
84 port, homeostasis, and sensorial functions (Moreira et al. 2015). Gas exchange
85 occurs mainly in the center of the gill filament, where the hemocytes circulate
86 through hemolymph vessels. Most bivalves absorb oxygen directly from water
87 through their tissues and oxygen-carrying molecules such as hemocyanin have been
88 identified in only a few genera. As coelomates, bivalves have another characteristic
89 cavity, the coelom, a small pericardial cavity enclosing the heart. Hemolymph is
90 pumped throughout the body by the heart, which receives oxygenated blood from
91 the gills and pumps it into the main blood vessel, a short artery that opens directly
92 into the hemocoel. Bivalve molluscs have an open circulatory system, with the
93 hemolymph reaching all of the organs by passive diffusion aided by the pumping
94 effect of the heart, which also has excretory functions. A pair of nephridia con-
95 nected to the coelom extracts any reusable materials from the coelomic cavity,
96 dumps additional unwanted products into it, and then excretes all of the materials
97 into the mantle cavity. In bivalves, gonads are located within the connective tissue
98 at the edge of the mantle, with spawning occurring directly in the mantle cavity
99 (Ruppert et al. 2004).

100 Depending on the species, bivalves feed on suspended particles in the water col-
101 umn, using an inhalant opening or siphon and ctenidia (e.g., *Magallana* and
102 *Crassostrea* spp. oysters); on deposits or particles on top of sediments, using an
103 inhalant siphon and ctenidia (e.g., *Macoma* spp. clams); or on deposits in the sedi-
104 ments, using proboscides (e.g., *Yoldia* spp. clams). Many bivalves are able to pump
105 large volumes of water while feeding. In bivalve species that use the ctenidia to
106 feed, food particles (mainly phytoplankton) are selectively trapped in a thick layer
107 of mucus covering the gills, transported with the aid of the cilia, sorted, and directed
108 to the outer labial palps, where particles are further sorted on the basis of size and
109 other physical and chemical characteristics. Some particles are then transferred to
110 the mouth by the inner palps, while other particles are rejected in pseudofeces
111 released into the pallial space. Mucus and cilia facilitate particle movement toward
112 the stomach, where there is further sorting and selection of particles (Ward and
113 Shumway 2004), leading to the prostyle, a mass of food and mucus. The prostyle is
114 extracellularly digested by the action of the enzymes produced by the digestive
115 gland. In most bivalve species, phagocytic cells have been evidenced in the tubules
116 of the digestive diverticula, where they contribute to intracellular digestion of the
117 selected particles reaching this organ. The remaining particles are excreted via the
118 nephridia or via the gut and finally reach the mantle cavity through the anus (Ruppert
119 et al. 2004).

120 Although mostly a sedentary group in their adult life stages, some bivalve spe-
121 cies are able to move. Most bivalves rely on the foot, a muscular organ with senso-
122 rial abilities achieved through balance receptors, the statocysts (Williamson 1993).
123 Larval pediveligers use the foot to sense and locate appropriate substrate for

settlement. In burrowing species such as clams, the foot is used by adults to burrow into the sediments. In mussels, the foot is linked to the production of byssus, an extremely resistant extracellular protein used to attach to the substrate (Carrington et al. 2015). Some species of bivalves (e.g., scallops) are also able to swim by rapidly opening and closing the two valves of the shell (Ruppert et al. 2004).

The nervous system of bivalve molluscs has a simple structure, organized in paired ganglia connected by nerve commissures within them and nerve cords along them in a “rope ladder structure.” The visceral cords innervate the internal organs and the pedal cords innervate the foot. The ganglia are divided in two groups: (1) cerebral, pleural (absent in bivalves), and visceral above the esophagus; and (2) the pedal ganglia below. These two differentiated parts are connected by the collar nerve, which surrounds the esophagus (Ruppert et al. 2004).

Ecological and Economical Roles

Bivalve molluscs cover multiple important roles, from both ecological and socio-economic points of view. Ecologically, bivalves have a key role in the environmental energy flux, in the maintenance of water quality by filter feeding and, for reef-building species such as oysters, in providing substrates and habitats for other species (Zu Ermgassen et al. 2012). Several bivalve species, and mussels in particular, have been used worldwide as sentinels for environmental pollution because of their sedentary and cosmopolitan nature in coastal waters, ease of sampling, ability as filter feeders to concentrate pollutants, and commercial use as an important food staple (Campos et al. 2012; Farrington et al. 2016; Burgos-Aceves and Faggio 2017). Bivalves can also concentrate pathogens and marine toxins, reaching harmful levels for consumers (Visciano et al. 2016). Moreover, as exemplified in Fig. 5, bivalves constitute a major sector of world fishery and aquaculture production, with more than 16 million metric tons with a value of almost US\$18 million produced in 2015, representing 15% of total aquaculture production (FAO 2016).

The main purpose of the molluscan aquaculture industry is to produce food, although this industry also has other applications such as ecosystem restoration, extraction of pharmaceutical and industrial products, and ornamentation (aquaria, nacre, pearls). The most important cultured species of molluscs are bivalves such as oysters, mussels, clams, cockles, and scallops, hence the focus of this chapter on these species. The culture process generally starts with the “conditioning” of broodstock in hatcheries by feeding them nutrient-rich cultured microalgae. Spawning is initiated by manipulation of environmental conditions (i.e., temperature, food availability) or, in some cases, gametes are surgically harvested. Fertilization is achieved by mixing of sperm and eggs. Larvae are kept in the hatchery while being fed cultured microalgae until they undergo metamorphosis and settle, and the small juveniles (also called spat) are moved out of the hatchery to a nursery and/or grow-out facility in open water to take advantage of the natural food supply. Grow-out culture technology varies depending on the species and location but can include the use of rope culture (mussels), cages/bags (oysters), and planting in natural beds (clams).

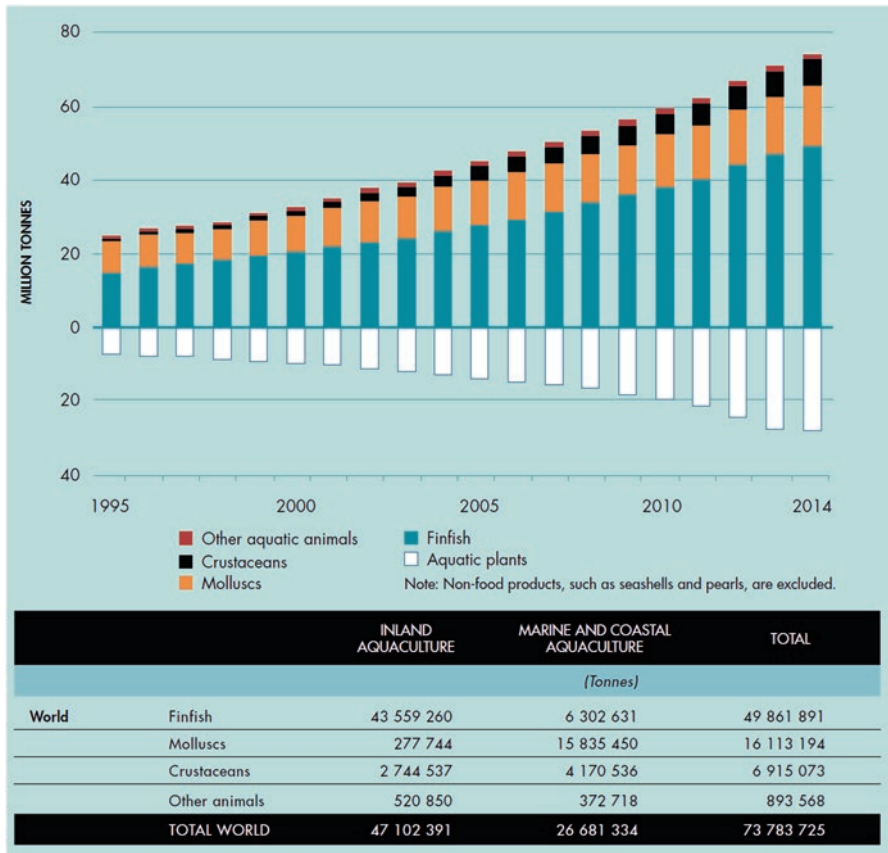


Fig. 5 World aquaculture production from 1995 until the present day. (FAO 2016)

166 Feeding relies on natural phytoplankton production at the site, and most of the labor
 167 involves predator and biofouling control.

168 Major Infectious Diseases Affecting Bivalve Molluscs

169 The commercial importance of many bivalve molluscs and efforts to manage dis-
 170 eases that severely impact the aquaculture industry have driven much of the research
 171 in the immunology of these species. Bivalve aquaculture has been severely impacted
 172 in recent years by infectious diseases and toxins from harmful algal blooms causing
 173 morbidity and mortality, as well as closures of the industry due to the accumulation
 174 of toxins and pathogens affecting the health of human consumers (GLOBEFISH
 175 2017). The relevance of these diseases is highlighted by the fact that the World
 176 Organization for Animal Health (most commonly known as the OIE) lists six dis-
 177 eases affecting bivalve molluscs among those with major relevance for animal

protection (OIE 2017). While pathologies caused by viruses, bacteria, and parasites have been documented in nearly all major molluscan classes, in this chapter we will present an overview of the pathological agents that have so far been relevant causes of concern for marine aquaculture activities and most commonly used as models in the study of bivalve immunity, leaving a discussion of infectious agents targeting other molluscs to the section “An Overview of Infectious Agents with Which Molluscs Must Contend” in Chap. 12.

AU8

Many diseases affecting bivalves result from an accidental side effect derived from the transfer of aquaculture species, leading to naïve hosts (indigenous or introduced) being exposed to new pathogens. Disease dynamics are heavily influenced by environmental factors, mainly temperature and salinity (Carella et al. 2015; Lafferty and Hofmann 2016; Stentiford et al. 2017), which are remarkably influenced by human activities, as thoroughly discussed in the section “Challenges for Molluscs in the Anthropocene Epoch” in Chap. 12. The study of bivalve immunology has benefited from many decades of research on host–pathogen interactions, the identification of species displaying natural resistance to diseases, the development of disease-resistant strains through selective breeding, and the recent application of -omic tools to bivalve research (Allam and Raftos 2015; Gómez-Chiarri et al. 2015). Most of the research has been focused on pathogens that can be cultured (Fernández Robledo et al. 2014).

Major Viral Diseases of Marine Bivalves

Although the characterization of viral diseases in bivalves has been hampered by the lack of cell lines from marine molluscs, recent advances in sequencing and the development of challenge models and disease-resistant strains have resulted in a better understanding of viral pathogenesis and immunity in several commercially important marine molluscs (Arzul et al. 2017). The best-characterized viral disease of bivalves is caused by oyster herpesvirus 1 (OsHV-1) and its variants (OsHV-1 Var and several microvariants, μ Var). Massive mortalities of bivalve larvae and/or juveniles due to OsHV-1 infection have seriously impacted the oyster industry in Europe, but also in Mexico, the USA, Australia, New Zealand, China, Japan, and Korea. These infections are recurrent in Pacific oysters (*Magallana gigas*), but other species of oysters, clams, mussels, and scallops are affected as well (Arzul et al. 2017). As shown for other diseases, some strains and species of bivalves appear to be resistant to or tolerant of the disease, such as the Sydney rock oyster, the eastern oyster, and mussels (Masood et al. 2016). Susceptibility to the disease also varies with age, size, and genetics within a species, and several selectively bred lines of Pacific oysters with increased resistance have been developed (Dégremont et al. 2015). In contrast to herpesviruses infecting vertebrates, both inter- and intraspecies horizontal transmission of OsHV-1 have been shown, with more tolerant individuals or species acting as disease carriers and reservoirs (Arzul et al. 2017).

220 Morphological and genomic characterization has led to the classification of this
221 large enveloped virus as a member of the *Malacoherpesviridae* (Mushegian et al.
222 2018). The function of most of the 124 ORFs found in the OsHV-1 viral genome is
223 unknown, mostly because of lack of homology with sequences with known function
224 (He et al. 2015; Arzul et al. 2017). Infection of oysters with OsHV-1 causes reduced
225 feeding and swimming in larvae. High levels of viral replication are observed
226 mainly in connective tissues, leading to changes in tissue and cellular architecture,
227 including dilation of the digestive tubules, nuclear chromatin margination and pyc-
228 nosis, and damage to the cytoskeleton and organelles. The disease is also character-
229 ized by massive infiltration of hemocytes. High levels of mortality occur within 48 h
230 postinfection in susceptible animals (He et al. 2015; Young et al. 2017).

231 Exposure of oysters to the virus through experimental challenges indicates that
232 the viral particles infect the host through the digestive gland and/or other mucosal
233 surfaces, probably exploiting hemocytes to reach target tissues (Segarra et al. 2016;
234 Morga et al. 2017). The virus is able to rapidly (within 1 h) infect and initiate repli-
235 cation in hemocytes. The formation of viral particles has not been observed in
236 hemocytes, however, suggesting that these cells impede completion of the viral
237 cycle, as observed in vertebrate macrophages infected with other herpesviruses
238 (Morga et al. 2017). Viral infection leads to activation of the integrin pathway in the
239 host cells, followed by activation of the actin pathway, indicating that the virus
240 exploits these pathways to enter the cell and eventually deliver the viral genome into
241 the nucleosome. Proteomic and metabolomic studies in challenged oysters show
242 that OsHV-1 causes substantial alterations in central carbon metabolism and gly-
243 colysis (Warburg effect) in the host, as well as alterations in lipid metabolism and a
244 characteristic fatty acid signature indicative of lipolysis. These metabolic alterations
245 increase the availability of substrates for virion synthesis and assembly. They can
246 also lead to increased inflammation and pathology through the activation of immune-
247 responsive gene 1 protein/*cis*-aconitic acid decarboxylase (IRG1/CAD), a protein
248 linking cellular metabolism with immunity, activation of the respiratory burst,
249 increased permeabilization of the mitochondrial membrane, and reduced ATP pro-
250 duction (Corporeau et al. 2014; Young et al. 2017).

251 Major Bacterial Diseases of Marine Bivalves

252 With a few exceptions (detailed below), mass mortalities caused by bacterial patho-
253 gens in bivalves are observed in larvae and, less often, in juveniles in hatcheries and
254 nurseries (Travers et al. 2015). Experimental challenges with bacterial pathogens,
255 however, are commonly used to study immune responses in bivalves because of the
256 ability to perform culturing and ease of isolation and characterization (Gómez-
257 Chiarri et al. 2015). A wide variety of *Vibrio* spp., including several belonging to the
258 *V. splendidus*, *V. harveyi*, and *V. tubiashii/coralliilyticus* clades, have been isolated
259 from outbreaks in bivalve hatcheries. In general, early signs of infection of bivalve
260 larvae by pathogenic vibrios include decreased feeding and damage to the velum,
261 followed by widespread necrosis of tissues and rapid mortality (Travers et al. 2015).

Strains of *V. aestuarianus*, *V. splendidus*, *V. crassostreae*, and others are often detected during summer mortality events in juvenile and adult Pacific oysters, also associated with infection with OsHV-1. Mass mortalities are, in general, seen during the spawning season and other conditions of stress (De Decker et al. 2011). The genomes of many of these pathogenic vibrios have been sequenced, facilitating the identification of mechanisms of virulence (Travers et al. 2015; Gómez-Chiarri et al. 2015). Examples of virulence factors involved in vibriosis include a variety of metalloproteases, hydrolases, cytotoxins, siderophores, the type III secretion system, and an OmpU from *V. tasmaniensis* LGP32, which is involved in internalization of the bacteria into *M. gigas* hemocytes (Travers et al. 2015; Le Roux et al. 2016).

Two bacterial pathogens of bivalves—*Aliiroseovarius crassostreae* and *Vibrio tapetis*—are notable for their ability to colonize the periostracal lamina of the inner side of bivalve shells. These pathogens cause Roseovarius Oyster Disease (ROD, also called Juvenile Oyster Disease) in the eastern oyster *Crassostrea virginica* and Brown Ring Disease in *Ruditapes* spp. clams, respectively. Susceptible bivalves respond to the presence of the pathogen in the inner side of the shell and the pallial cavity by producing conchiolin mixed with melanin and other quinones with antimicrobial action, resulting in pathognomonic brown deposits that surround the edge of the mantle (Travers et al. 2015). Little is known about mechanisms of virulence in ROD, but it is likely that formations of polar fimbriae and biofilm on the shell of oysters by *A. crassostreae* are involved in the disease (Boardman et al. 2008). Virulence factors identified in the genome of *A. crassostreae* include a hemolysin/cytotoxin and a putative type IVA secretion system (T4ASS) (Kessner et al. 2016). The metabolic demand of the chronic infections derived from an unsuccessful immune response in susceptible animals may contribute to mortality (Paillard et al. 2014; McDowell et al. 2014).

A few selected bacterial pathogens have been associated with sporadic episodes of mortality in adult bivalves, most notably *Nocardia crassostreae* and several intracellular Rickettsia-like organisms (RLOs). Little is known, however, about mechanisms of virulence and host immunity in these diseases (Travers et al. 2015; Zannella et al. 2017).

Major Parasitic Diseases of Marine Bivalves 294

Haplosporidian Parasites 295

Protistan parasites constitute the largest cause of adult bivalve morbidity and mortality. Among the most devastating groups of protozoan parasites of bivalve molluscs are several parasites belonging to the phylum Haplosporidia (Arzul and Carnegie 2015). In particular, the haplosporidians *Bonamia ostreae*, *B. exitiosa*, and *Haplosporidium nelsoni* have been well known for decades for causing significant economic and ecological losses, mainly in Europe and the USA. The growth of the bivalve aquaculture industry has led to the recent identification of many other haplosporidian parasites affecting a variety of bivalves. Most of the outbreaks caused

304 by the best-known representatives of this phylum, *B. ostreae* and *H. nelsoni*, have
305 been observed in adult oysters. While species from the genus *Bonamia* are only
306 known to affect oysters, have a direct mode of transmission, and are mostly intracel-
307 lular, other haplosporidian taxa have representatives affecting a wide variety of
308 bivalve hosts, are transmitted through intermediate hosts, and are typically extracel-
309 lular. Many aspects of the life cycle of these parasites are unknown, as they cannot
310 be maintained in culture. However, it is presumed that infective stages of *H. nelsoni*
311 enter the host through the epithelial lining of the gill, developing into multinucle-
312 ated plasmodia, which are seen in all tissues in heavily infected oysters. Depending
313 on the haplosporidian species, sporulation occurs in the epithelium of the digestive
314 diverticula or in connective tissues of the host, leading to the development of sporo-
315 cysts, which are thought to eventually burst upon death of the host, releasing spores
316 into the environment. Sporulation of *H. nelsoni* has rarely been observed in *C. virgi-*
317 *nica*, indicating that this oyster may be an atypical host. Oysters that have survived
318 outbreaks of *H. nelsoni* and *B. ostreae* show increased resistance to these diseases,
319 a fact that has been exploited in the development of selectively bred disease-resistant
320 strains (Arzul and Carnegie 2015; Morga et al. 2017).

321 **Cercozoan Parasites**

322 Several *Marteilia* spp. (Cercozoa, Paramyxida) have been responsible for flat and
323 Sydney rock oyster epizootics in Europe and Australia. These parasites affect a
324 diversity of molluscan hosts, including oysters, clams, and mussels, and disease
325 pathogenesis varies depending on the *Marteilia* spp. and the host. Clinical signs of
326 the disease may include nodules (a gross manifestation of an encapsulation response)
327 and, in many of the species, necrotic damage to the digestive gland. As other
328 Paramyxean parasites, *Marteilia* spp. show a characteristic cell-within-cell develop-
329 ment by budding. Therefore, most aspects of their complex life cycle, pathogenesis,
330 mechanisms of virulence, and modes of transmission remain a mystery, since efforts
331 to culture these parasites or transmit the disease using cohabitation challenges have
332 been unsuccessful (Carrasco et al. 2015).

333 **Perkinsozoan Parasites**

334 Perkinsosis is caused by a variety of species belonging to the genus *Perkinsus* (phy-
335 lum Perkinsozoa, superphylum Alveolata). The first *Perkinsus* spp. to be character-
336 ized, *Perkinsus marinus*, was identified in the 1940s as the cause of mass mortalities
337 of eastern oysters in the Gulf of Mexico. As is the case for haplosporidian parasites,
338 many other species have been described with the growth of the bivalve aquaculture
339 industry, including *P. olseni*, *P. chesapeaki*, *P. mediterraneus*, *P. beihaiensis*, *P. hon-*
340 *shuensis*, and *P. qugwadi*. While the geographic range of *P. marinus* seems to be
341 limited mainly to that of *C. virginica* in North America, other *Perkinsus* spp., such
342 as *P. olseni*, have a wider geographic and host range. Therefore, *Perkinsus* spp.
343 affect oysters, clams, scallops, cockles, and mussel species in Australia, New
344 Zealand, Asia, America, and Europe (Reece et al. 2017). These parasites have a
345 direct life cycle with four described life stages: trophozoites, hypnospores (or pre-
346 zoosporangia), zoosporangia, and biflagellated spores (Soudant et al. 2013). The

disease is transmitted horizontally, infecting the host through the epithelia of the digestive tract and mantle after the parasites are brought into the pallial cavity and ingested through feeding. Although *Perkinsus* spp. can cause relatively rapid mortality with few clinical signs in the most susceptible individuals within a population, it is most frequently manifested as a chronic disease in adult bivalves. Signs of disease are characterized by severe hemocytic infiltration of tissues, a decrease in gametogenesis and the condition index and, in some individuals, death by occlusion of vascular sinuses, tissue necrosis, and/or emaciation. In some host species, such as *Ruditapes* spp. clams infected by *P. olseni*, the chronic response is characterized by granuloma-like formations, which can be visibly detected as nodules at the base of gills. Parasites are transmitted to other hosts after being released to the water through diapedesis, in feces, or at the death of the host (Soudant et al. 2013; Ruano et al. 2015). Clonal cultures of most *Perkinsus* spp. are available, allowing for the characterization of putative virulence factors through genetic, genomic, and proteomic studies (Gómez-Chiarri et al. 2015; Hasanuzzaman et al. 2016; Fernández-Boo et al. 2016). Some interesting examples of mechanisms of virulence potentially contributing to the ability of *P. marinus* to survive within the hemocytes of the eastern oyster (Alavi et al. 2009) include antioxidant enzymes, such as superoxide dismutases (Schott and Vasta 2003; Schott et al. 2003; Asojo et al. 2006; Fernández-Robledo et al. 2008) and ascorbate-dependent peroxidases (Schott et al. 2003), and a natural resistance-associated macrophage protein (NRAMP) (Lin et al. 2011). Exposure of *P. marinus* to oyster tissue homogenates or pallial fluid in vitro modulates the production of serine proteases and the expression of genes coding for anti-apoptotic proteins, heat shock proteins, and proteinase inhibitors (Soudant et al. 2013; Pales Espinosa et al. 2014). Another interesting feature of *Perkinsus* spp. may be the presence of a relic plastid with no photosynthetic capabilities (Fernández Robledo et al. 2011) and the ability to secrete several fatty acids, including arachidonic acid (Soudant et al. 2013). Differences in resistance to or tolerance of infection by *Perkinsus* spp. have been documented within and between bivalve species, and selectively bred lines with moderate resistance to or tolerance of *P. marinus* are available (Proestou et al. 2016).

Quahog Parasite Unknown

The protist Quahog Parasite Unknown (Labyrinthulomycetes, Stramenopiles), better known as QPX, causes an opportunistic disease in the quahog *Mercenaria mercenaria* in the northeast and mid-Atlantic regions of the USA (Burge et al. 2013). The disease caused by QPX is characterized by the presence of areas of massive focal inflammation, visibly manifested as nodules commonly observed at the edge of the mantle or the base of the siphon. Differences in susceptibility to QPX infection have been observed between clam populations from different geographic locations (with clams originating south of Virginia being more susceptible than northern clams) and lines of clams derived from survivors of disease outbreaks. Resistance is probably due to a combination of factors, including adaptation to local conditions, as well as selection for molecules involved in more effective immune responses against the parasite (Wang et al. 2016b). QPX is a saprophyte that secretes a thick

391 mucus layer while in tissues of the clam that appears to protect the parasite from
392 the immune response of the host. Putative virulence factors include a variety of
393 hydrolytic enzymes and proteases, antioxidants, polysaccharide production, and
394 factors involved in recognition, such as lectins. The expression of many of these
395 putative virulence factors—in particular, genes that may be involved in the forma-
396 tion of the protective mucus layer—are significantly regulated by temperature
397 (Rubin et al. 2017).

398 **Metazoan Parasites**

399 Some metazoan parasites have been documented in marine bivalves, including the
400 copepod *Mytilicola intestinalis* (a parasite of mussels) and the trematode
401 *Schistosoma mansoni* (a parasite of humans that also infects snails). Trematode
402 infections are common in molluscs, which act as intermediate hosts. This complex
403 host–parasite interplay is modulated by pattern recognition and effector molecules,
404 as thoroughly reviewed by other authors (Zhang and Loker 2004; Adema et al.
405 2010; Pila et al. 2017) and discussed in detail in the section “Disease-Transmitting
406 Snails” in Chap. 12.

407 **A General Overview of Bivalve Immunity**

408 **Feeding: An Aspect Not to Be Overlooked**

409 Invertebrates, including molluscs, lack the acquired response in a narrow sense
410 (Criscitello and de Figueiredo 2013), but they possess a potent and efficient cellular
411 and humoral innate immune system, physical barriers such as the shell and the
412 mucus, and behavioral avoidance. This innate response involves, as its major play-
413 ers, circulating hemocytes and a broad range of diverse molecular effectors. A gen-
414 eral overview of immune defenses in bivalves is depicted in Fig. 6. One of the first
415 lines of defense of bivalves against pathogens derives from their ability to sense the
416 environment and sort particles during feeding (Ben-Horin et al. 2015). As described
417 in the section “Anatomy and Physiology of Bivalves”, bivalves are filter feeders,
418 and the surfaces of the mantle and the gills are exposed to large volumes of water
419 containing microbes and plankton. Bivalves are able to distinguish non-nutritious or
420 potentially harmful particles on the basis of size, physical, and chemical cues, and
421 reject (expel) these particles using mucociliary mechanisms. Bivalves are also able
422 to shut down feeding and keep the valves tightly closed under unfavorable environ-
423 mental conditions (e.g., low oxygen or blooms of an undesirable phytoplankton
424 species). Although the specific roles of sensing and behavioral responses in disease
425 resistance and immunity have not been well studied, some recent evidence indicates
426 that these may be an interesting avenue for further study. For example, it is thought
427 that oysters accumulate relatively less domoic acid (a toxin produced by the harmful
428 algae *Pseudo-nitzschia* spp.) than mussels, in part because oysters ingest fewer algal
429 cells (Mafra et al. 2010). There is also evidence that feeding behavior is responsible
430 for increased resistance to the parasite *P. marinus* observed in some selectively bred

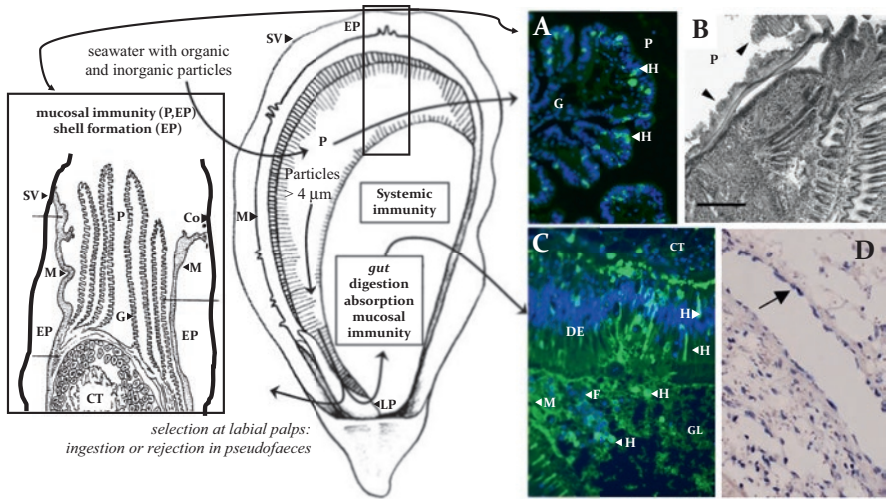


Fig. 6 Overview of immune responses in a representative bivalve (an oyster). *Center*: View of oyster tissues on top of one of the valves, illustrating the flow of water and particles during feeding (Troost 2010). *Left*: Lateral view of the ventral side of an oyster, showing the pallial (P) and extra-pallial (EP) cavities. *Right*: Micrographs illustrating examples of cellular responses in different immune compartments. **(a–c)** Examples of mucosal immune responses. **(d)** Example of a systemic immune response (see below for more details). When the two shell valves (SV) characteristic of bivalves open to allow for feeding, water is pumped through the gills (G) and particles are selected to be either rejected or brought into the gut (central panel). Cells in the mucosal epithelium of the gills and mantle (M) secrete mucus and other effectors. The mantle is also responsible for sealing the edge of the shell valve from the environment (left panel) and producing conchiolin (Co, in the drawing on the right and the arrowhead in **(b)**). Hemocytes (H) can migrate into the pallial and extrapallial cavities **(a and b)**, the gut **(c)**, and the blood sinuses **(d)** to recognize, capture, and digest particles and pathogens. **(a)** Immunofluorescence image of a section of oyster gill (G) tissue, showing hemocytes labeled in green (H). Shown in blue are cell nuclei stained with Hoestch. **(b)** H&E-stained sections of a challenged oyster showing degeneration and erosion of the mantle associated with hemocytic infiltration (arrows) and the presence of conchiolin (arrowheads) (scale bar = 100 μm) (Gomez-Leon et al. 2008). **(c)** Immunofluorescence image of a section of oyster gut showing the digestive epithelium (DE), with hemocytes labeled in green (H). The presence of mucus (M) and algal food (F) can be observed in the gut lumen (GL). Shown in blue are cell nuclei stained with Hoestch. **(d)** Big-defensin labeling in hemocytes (arrow) at the edge of a blood vessel in Pacific oysters challenged with *V. anguillarum* (Rosa et al. 2011)

AU10

AU11

families of eastern oysters, with oysters from resistant families removing (filtering) 431
 fewer algal cells from the water when mixed with *P. marinus* than susceptible oysters 432
AU9 (Ben-Horin and Proestou personal communication). 433

Mucosal Immunity: An Important Yet Understudied Topic 434

Mucosal immunity constitutes the next barrier to infection on those tissue surfaces 435
 in contact with the external environment, while maintaining tolerance of nonharm- 436
 ful commensal microbes and innocuous substances. Mucosal immunity represents 437

438 an important, but understudied, first line of immune defense, extending the defensive
439 role of mucus beyond that of a simple physical barrier (Allam and Pales Espinosa
440 2016) in all molluscs, as detailed in the section “[Molluscan Immunity Begins at the](#)
441 [Mucosal Surface, an Immunologically Active Site That Remains Understudied](#)” in
442 Chap. 12. This aspect seems to be of primary importance in bivalves, as their life is
443 tightly linked to aquatic environments. Indeed, bivalves can overcome an experi-
444 mental pathogen challenge by bath exposure but cannot overcome experimental
445 challenge with smaller amounts of the same pathogen if exposed by injection.
446 Pathogens able to bypass these initial barriers to infection (either by surviving inside
447 phagocytic cells or by directly migrating through epithelial junctions) then trigger a
448 systemic immune response. In general, for both mucosal and systemic immunity,
449 the recognition of nonself (in the form of microbe-associated molecular patterns
450 (MAMPs)) by lectins and other pattern recognition receptors (PRRs) and opsonins
451 in hemolymph (see section “[Recognition, Agglutination, and Opsonization](#)”), and
452 by sentinel cells (most probably hemocytes), present in the tissues, triggers signal-
453 ing transduction cascades and the release of cytokines (see section “[Signaling and](#)
454 [Regulatory Pathways](#)”), leading to humoral immune responses (see section
455 “[Humoral Immune Effectors](#)”) and cellular immune responses (see section “[Cellular](#)
456 [Immune Responses](#)”) that vary according to the nature and location of the immune
457 stimuli. A fine regulation of the immune response is achieved through the neuroen-
458 docrine immunomodulation (NEI) regulatory network (see section “[Connections](#)
459 [with the Neuroendocrine System](#)”), a cross talk between the nervous, endocrine,
460 and immune systems that maintains homeostasis and tunes innate immune response
461 in all animals.

462 In particular, mucosal immune responses include (a) the production of humoral
463 defense factors secreted into the mucus covering the epithelium of tissues in either
464 the pallial or the extrapallial space; (b) chemotaxis and the transepithelial migration
465 of hemocytes into the pallial and extrapallial spaces, followed by phagocytosis and
466 intracellular killing; (c) phagocytosis and intracellular digestion by cells in the
467 digestive epithelium; and, if needed, (d) an encapsulation response in the extrapal-
468 lial cavity characterized by the secretion of conchiolin and antimicrobial products
469 and activation of the prophenoloxidase cascade (see section “[The Phenoloxidase](#)
470 [Cascade](#)”) (Allam and Raftos 2015; Allam and Pales Espinosa 2016; Zannella et al.
471 2017). Systemic immune defenses include (a) recognition, opsonization, phagocy-
472 tosis, and intracellular killing by circulating hemocytes and other, yet to be identi-
473 fied, phagocytic cells within tissues; (b) killing in plasma through secretion of
474 humoral effectors and activation of an ancient complement system and the pheno-
475 oxidase system; and, if needed, (c) an encapsulation response that leads to
476 granuloma-like formations, grossly visible as nodules in extreme cases.

477 **Hemocytes: Key Cellular Players in Bivalve Immune Response**

478 Hemocytes are a key component of the bivalve immune system. These cells are
479 present in all cavities of bivalves, circulating in the hemolymph (which bathes all

tissues) and migrating into the pallial and extrapallial spaces. Different types of hemocytes have been described in molluscs on the basis of morphological characteristics (see section “[A Short Journey in the ‘Immune System’ of Cephalopods](#)” for a brief comparative overview between bivalve and cephalopod hemocytes and the section “[Hemocytes Play a Central Role in Molluscan Immune Responses: Some Basics Regarding Their Morphology and Origins](#)” in Chap. 12 for a broader discussion), and their roles in both physiological processes (e.g., digestion and shell formation) and immune functions (e.g., phagocytosis, synthesis of immune effectors, and modulation of immune responses) are well known (Cheng 1984; Ordás et al. 2000; Goedken and De Guise 2004; Costa et al. 2009b; Wang et al. 2017c; Ivanina et al. 2017).

The lack of specific cell markers, however, has so far prevented detailed characterization of the functionality and mechanism of action of specific cell populations; thus, recent efforts dedicated to the development of these markers are particularly exciting (Donaghy et al. 2009; Sekine et al. 2016; Allam and Pales Espinosa 2016). Moreover, the location of the hematopoietic organ and the process of hematopoiesis and maturation into distinct hemocyte populations are still controversial topics (Pila et al. 2016; Dyachuk 2016). While the hematopoietic organ in gastropods is the amoebocyte-producing organ (Jeong et al. 1983) and that in cephalopods is the white gland (Cowden and Curtis 1973), a variety of tissues in different species and developmental stages have been proposed as hematopoietic organs in bivalves. These include an irregularly folded structure in the gills (Jemaà et al. 2014) and unspecified locations within the mantle and gills (Song et al. 2016) of adult oysters, the mantle edge of mussel larvae (Balseiro et al. 2013), the connective tissues and gill epithelium of recently settled larvae from the flat oyster *Ostrea edulis* (Xue and Renault 2001), and a ring structure around the dorsal side of the embryo in oyster trochophore larvae (Song et al. 2016).

Expansion and Molecular Diversification: The Bivalve Immune System Is Not as “Simple” as We Thought

Exploration of molluscan genomes has revealed massive expansion and functional divergence of gene families involved in immune recognition and opsonization (detailed in section “[Recognition, Agglutination, and Opsonization](#)”), adhesion (syndecan, protocadherin), acute phase responses (hsp70), signal transduction (see section “[Signaling and Regulatory Pathways](#)”), cytokine production (see section “[Production of Cytokines](#)”), apoptosis (see section “[Apoptosis and Autophagy](#)”), or oxidation and antioxidation (cytochrome p450, superoxide dismutase) (Zhang et al. 2012a; Simakov et al. 2013; Albertin et al. 2015; Murgarella et al. 2016; Sun et al. 2017; da Silva et al. 2017; Mun et al. 2017; Du et al. 2017). Many of these immune gene family expansions are lineage (bivalve) specific (Zhang et al. 2015; McDowell et al. 2016). The mechanisms (i.e., gene duplications, rearrangements, polymorphism, etc.) and functional relevance of these gene expansions and divergence are still being studied, but there are indications that gene diversity may be responsible

522 for a certain level of species specificity in bivalve immune responses (see Chap. 12,
523 section “[Expansion and Diversification of Innate Immune Gene Families](#)” for a
524 comparative overview of a few specific cases).

525 **Evidence of “Immunological Memory” in Bivalves**

526 The plasticity of bivalve immune responses is also evidenced by indications that
527 the immune system can be primed, leading to short-term memory. For example,
528 scallops and oysters showed enhanced pathogen-specific phagocytosis upon a sec-
529 ondary challenge and upregulation of expression of genes involved in phagocytosis
530 and hematopoiesis (Zhang et al. 2014d; Wang et al. 2015b; Green et al. 2015;
531 Pinaud et al. 2016; Wang et al. 2017a). Recent experiments have further indicated
532 that experimentally infected juvenile oysters can mount a long-lasting antiviral
533 immune memory, persisting for at least 5 months, which protects them from sub-
534 sequent viral infections (Lafont et al. 2017). Furthermore, transgenerational
535 immune priming has been demonstrated in bivalves (Green et al. 2016). The spec-
536 ific mechanisms involved in these two types of priming are still unclear, but the
537 switch from cellular to humoral response and epigenetic regulation are believed to
538 play crucial roles. An in-depth discussion of the relevance of this poorly under-
539 stood phenomenon in molluscs is provided in the section “[Immune Priming](#)” in
540 Chap. 12. The role of maternal transfer has been also studied as a part of the innate
541 immune response in molluscan larvae, making transgenerational immune priming
542 possible. Bivalve oocytes possess significant antibacterial, lysozyme, and agglutinat-
543 ing activities against pathogens, and several immune factors have been identified in
544 embryos (Wang et al. 2015b).

545 **How Do Environmental Factors Affect the Bivalve Immune** 546 **Response?**

547 Bivalves are poikilotherm species living in highly diverse and variable environ-
548 nments. Consequently, immune responses are heavily affected by environmental con-
549 ditions, such as temperature, salinity, dissolved oxygen, pH, and pollution.
550 Therefore, an extensive body of knowledge has been built about the potential effect
551 of environmental stress and pollution on immune parameters in these organisms and
552 other molluscan groups—in particular, in connection with human activities, as dis-
553 cussed in detail in the section “[Challenges for Molluscs in the Anthropocene Epoch](#)”
554 in Chap. 12. For example, exposure of bivalves to environmental toxins of natural
555 origin, like those derived from harmful algal blooms or toxic cyanobacteria, has
556 been shown to affect the phagocytic responses of bivalves, generally leading to
557 immunosuppression (Hégaret et al. 2011; Soudant et al. 2013; Queiroga et al. 2017).
558 Exposure of oyster hemocytes to pollutants such as TBT in vitro and in vivo reduces
559 their production of ROS and phagocytic activity (Soudant et al. 2013), and exposure
560 of bivalve hemocytes in vitro to nanomaterials leads, in general, to decreased

phagocytic activity, increased antioxidant levels, and increased apoptosis, indicating immunotoxicity (Rocha et al. 2015). The effects of environmental stressors on bivalve immunity, however, depend on the evolutionary history of the bivalve species and the history of exposure to different environmental conditions between populations within a species.

Recognition, Agglutination, and Opsonization

The Role of Lectins in Immune Recognition

A critical step of innate immune responses against an infectious challenge is the immediate recognition of the “nonself” carbohydrate moieties on the surface of potential pathogens and parasites, such as viral envelope glycoproteins, bacterial lipopolysaccharides and exopolysaccharides, and various surface glycans on eukaryotic parasites (Boehm 2012). These surface structures encode vast information that is “decoded” by the hosts’ carbohydrate-binding proteins (lectins) (Vasta and Ahmed 2008) which, upon binding to the recognized ligand, can immobilize the infectious agents and activate downstream signaling pathways, leading to their uptake and intracellular killing by phagocytic cells. Furthermore, lectin-mediated activation of the complement system can also promote phagocytosis and killing of potential pathogens (Fujita et al. 2004; Vasta et al. 2007) (see section “Evidence of an Ancient Complement System in Bivalves?”). Thus, lectins are critical components of innate immune mechanisms as both recognition and effector factors—functions that are facilitated by the oligomerization of lectin peptide subunits, leading to increased avidity for the multivalent glycan ligands typically found on the microbial surface (Taylor and Drickamer 2003; Vasta et al. 2007). On the basis of the identification of unique amino acid sequence motifs and the structural fold of the carbohydrate recognition domain (CRD), and the requirement of divalent cations or a reducing environment for ligand binding, lectins have been classified into several major families. These include C-type lectins (CTLs), FTLs, RTLs, HTLs, PTLs, XTLs, I-type lectins, pentraxins, galectins (formerly S-type lectins), ficolins, and others (Vasta et al. 2007). Members of several lectin families such as CTLs, RTLs, FTLs, peptidoglycan-binding proteins, ficolins, pentraxins, and galectins have been implicated in immune surveillance and homeostasis (Vasta and Ahmed 2008) (Fig. 7).

Unlike immunoglobulins (Igs) and Ig superfamily members such as DSCAM (Yue et al. 2016) and FREPs (Zhang et al. 2004), which generate recognition diversity by genetic mechanisms, lectins are typically described as “hard wired” in the germline (Vasta et al. 2007). Therefore, given the great diversity of potential infectious agents present in the aquatic or terrestrial environments that molluscs inhabit, how their innate immune systems are able to cope with these infectious challenges is an outstanding question that remains to be fully addressed (Harvell et al. 1999). However, the complexity of the lectin repertoires in organisms that lack the typical Ig-mediated adaptive immunity, such as molluscs, strongly suggests that a wide variety of molecular topologies can be effectively recognized in surface carbohydrate

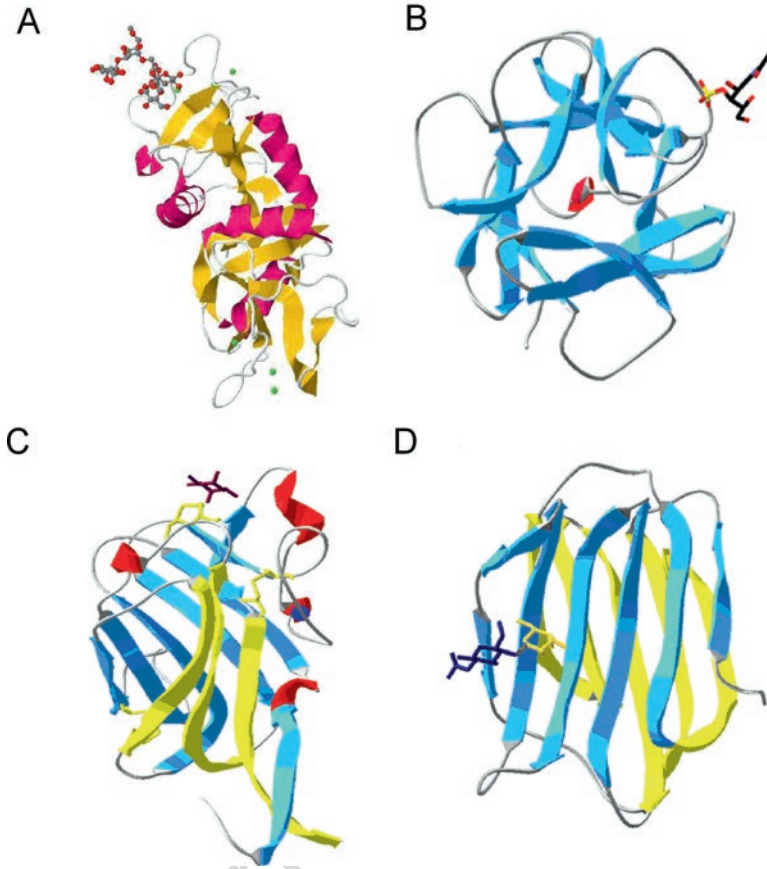


Fig. 7 Typical structural fold of four of the most important lectin families with functions in immune recognition in bivalve molluscs. (a) C-type lectin with bound carbohydrate ligand (PDB accession ID: 2MSB). (b) R-type lectin with bound 4-sulfated GalNAc (PDB accession ID: 1DQ0). (c) F-type lectin with bound fucose (PDB accession ID: 1K12). (d) Galectin with bound LacNAc (PDB accession ID: 1KJL)

602 moieties common to diverse microbial pathogens, leading to activation of effector
603 mechanisms that can kill and eliminate them for successful innate immune protec-
604 tion (Vasta et al. 2007, 2012a; Vasta and Ahmed 2008). A discussion of the best-
605 characterized lectin families identified in molluscs follows below.

606 C-Type Lectins

607 Together with the S-type lectins (currently known as galectins; see below) C-type
608 lectins (CTLs) were the first two families to be rigorously defined by the presence
609 of unique sequence motifs in their CRDs (Drickamer 1988). CTLs are characterized
610 by the CTL-like domain (CTLD) of the unique structural fold and the requirement
611 of Ca^{2+} for ligand binding. The CTLD can be structurally diversified and associated

with a variety of lectin and nonlectin domains constituting “mosaic” or “chimeric” proteins endowed with multiple functional properties (Zelensky and Gready 2005; Pees et al. 2016). In mammals, this highly heterogeneous lectin family is currently subdivided into 17 groups based on their domain organization (Zelensky and Gready 2005; Vasta and Ahmed 2008; Pees et al. 2016). CTLs participate not only in the initial step of pathogen recognition via the CRD but also in various antimicrobial effector functions, including pathogen recognition, opsonization, and activation of the complement cascade (Vasta et al. 2007). In invertebrate taxa, CTLs are also key factors in carbohydrate-mediated recognition of the infectious challenge, but also in effector roles such as immobilization, phagocytosis, clearance, and encapsulation of the infectious agent. Furthermore, they have also been implicated in nodule formation, in the activation of the prophenoloxidase/melanization cascade, and in other functions, including direct antimicrobial activity and regulation of antimicrobial peptide (AMP) expression (Vasta et al. 2007; Vasta and Ahmed 2008; Wang et al. 2014b; Pees et al. 2016; Zhao et al. 2016b). Numerous studies have been conducted in various mollusc species, aimed at investigating the potential role of CTLs in immune defense, and their roles in recognition, agglutination/immobilization, and opsonization of bacterial pathogens have been firmly established (Zheng et al. 2008; Zhu et al. 2008; Jing et al. 2011; Huang et al. 2013a; Zhang et al. 2014b; Mu et al. 2014; Martins et al. 2014; Chovar-Vera et al. 2015; Huang et al. 2015b; Yang et al. 2015). In general, the CTL repertoire in any single species appears to be highly diversified and complex, and the temporospatial expression and localization of CTLs includes hemocytes, plasma, and pallial mucus, as well as organs and tissues relevant to immune responses such as the mantle, gills and gut. Additionally, infectious challenge experiments have revealed that in most cases their expression is modulated by exposure to potential pathogens (Zhu et al. 2008; Mu et al. 2014; Martins et al. 2014; Chovar-Vera et al. 2015). The report that molluscs can express components of the complement system (see section “Evidence of an Ancient Complement System in Bivalves?”) (Li et al. 2015a; Wang et al. 2017b) has suggested that CTLs may function not only as pathogen agglutinins and opsonins but also in activating the complement cascade with further antimicrobial activity.

R-Type Lectins

The R-type lectins (RTLs) are lectins characterized by a CRD of unique structure, consisting of three lobes arranged around a threefold axis CRD (β -trefoil), in which each lobe may contain a carbohydrate-binding site (Cummings and Schnaar 2017). This structure is found in RTLs from higher plants as well as in hydrolases from prokaryotes, mammalian glycosyltransferases, and macrophage mannose receptors (Cummings and Schnaar 2017). RTLs with binding preference for α -D-galactose/GalNAc moieties and a very similar amino acid sequence have been isolated from the mussels *Crenomytilus grayanus* (CGL) (Jakób et al. 2015; Chernikov et al. 2017a, b), *Mytilus galloprovincialis* (Mytilectin-1) (Hasan et al. 2016; Terada et al. 2016), *Mytilus trossulus* (MTL) (Chikalovets et al. 2016), and *Mytilus californianus* (García-Maldonado et al. 2017). The RTL known as MytiLec-1 displays the typical β -trefoil structure (Terada et al. 2016), whereas two additional isoforms (MytiLec-2

656 and -3) identified in the same mussel species contain an additional pore-forming
657 aerolysin-like domain (Hasan et al. 2016; Terada et al. 2016). The structure of CGL
658 was resolved recently and shows a similar β -trefoil structure (Jakóbc et al. 2015).
659 RTLs from mussels can recognize and agglutinate both Gram-positive and Gram-
660 negative bacteria in a carbohydrate-dependent manner, display bacteriostatic activity,
661 and also show antifungal activity by binding to and inhibiting hyphal growth (Jakóbc
662 et al. 2015; Hasan et al. 2016; Terada et al. 2016; Chernikov et al. 2017a, b). It is
663 noteworthy that mytillectins and CGL also show immunomodulatory activity for
664 mammalian macrophages, and proapoptotic/antitumoral activity by binding to glo-
665 botriose [Gb3; Gal α (1,4)Gal β (1,4)Glc α 1] on the cell surface glycolipids such as glo-
666 botriosyl ceramide (Chernikov et al. 2017a, b)—properties that have revealed their
667 promise as effective diagnostic and therapeutic agents and have already led to the
668 computational design of an artificial lectin named Mitsuba-1 (Terada et al. 2017).

669 F-Type Lectins

670 F-type lectins (FTLs) are the most recent lectin family to be identified (Odom and
671 Vasta 2006), and they are characterized by a fucose recognition domain (F-type
672 lectin domain; FTLD) that displays a novel β -barrel jellyroll fold (“F-type” fold),
673 and unique carbohydrate- and calcium-binding sequence motifs (Bianchet et al.
674 2002). FTLs may exhibit single, double, or greater multiples of the FTLD and are
675 widely distributed in nature (Bianchet et al. 2002; Odom and Vasta 2006; Bianchet
676 et al. 2010). Like the CTLs, FTLs may display FTLDs combined with other struc-
677 turally and functionally distinct domains, yielding lectin subunits of pleiotropic
678 properties even within a single species (Bianchet et al. 2002; Odom and Vasta 2006;
679 Bianchet et al. 2010; Vasta et al. 2012a). Although the F-type fold is distinctive for
680 FTLs, it is not unique to these lectins, as other proteins with various functions also
681 display the FTLD fold (Bianchet et al. 2002). Interestingly, although a phylogenetic
682 analysis of FTLD sequences from viruses to mammals has revealed consistency
683 with the taxonomy of extant species, the surprisingly discontinuous distribution of
684 FTLDs within each taxonomic category suggests not only an extensive structural/
685 functional diversification of FTLs along evolutionary lineages but also that they
686 have been subject to frequent gene duplication, secondary loss, lateral transfer, and
687 functional co-option (Bianchet et al. 2002; Bishnoi et al. 2015).

688 In addition, FTLs are unique in the extraordinary sequence variability (isoforms)
689 that can be expressed in a single individual as a result of genetic mechanisms of
690 diversification in ligand recognition, characterized in detail in the so-called bindins,
691 proteins involved in gamete recognition in the Pacific oyster, *M. gigas* (Springer
692 et al. 2008; Moy et al. 2008; Moy and Vacquier 2008). In addition to their roles in
693 gamete recognition, oyster FTLs also mediate microbial recognition in innate
694 immune responses. FTLs can display single or tandemly arrayed CRDs of distinct
695 specificity in a single subunit (Odom and Vasta 2006; Bianchet et al. 2010), and can
696 potentially cross-link the recognized pathogens to the endogenous glycans on the
697 surface of the host’s phagocytic cells (Odom and Vasta 2006). In this regard, the
698 expression of CvFBL4 in *C. virginica* hemocytes is dramatically upregulated upon
699 LPS challenge, suggesting that FTLs may function in pathogen recognition in the

oyster's innate immune response (Saito and Vasta unpublished data). Moreover, PmF-lectin from the pearl oyster (*Pinctada fucata martensii*) is an FTL highly expressed in the hemocytes and gill that is significantly upregulated by experimental challenge with *Vibrio* sp. (Wang et al. 2011a). The identification of FTLs in both the shell matrix and mantle tissue proteins of the blunt-gaper clam, *Mya truncata*, has led to the proposal that during the shell biomineralization process, FTLs secreted by the mantle may carry out immune defense functions and are later incorporated into the shell matrix (Arivalagan et al. 2016). It is noteworthy that the highly diversified FTL repertoire found in the common periwinkle (*Littorina littorea*), a gastropod, has been rationalized as an immune defense system (Gorbushin and Borisova 2015). However, in contrast to other expanded lectin and lectin-like gene families, this connection has not been hypothesized yet in bivalves.

H-Type Lectins

H-type lectins (HTLs) are lectins initially identified in gastropods such as the Roman snail *Helix pomatia* as abundant proteins in the albumin gland secretion that coats the fertilized oocytes before the eggs are laid underground (Uhlenbruck and Prokop 1966). This unique localization as perivitelline active factors, their presence in the snail's hemolymph, and their strong binding to several streptococci strains and other potentially pathogenic bacteria led to the proposal that their role was to protect the snail eggs and adults from infection, as part of the innate immune defense (Uhlenbruck and Prokop 1966). Their shared specificity for N-acetylgalactosamine (GalNAc) and the human blood group A led to their use as typing reagents (Uhlenbruck and Prokop 1966). Recent structural studies revealed that HTLs are characterized by hexameric organization of peptide subunits that display a β -sandwich fold. Although other snail species from the genus *Helix* and the garden snail *Cepaea hortensis* also produce similar lectins (Sanchez et al. 2006), to date, no functional information has been collected yet about HTLs in bivalves, other than the fact that they do not represent an expanded gene family (Gerdol 2017).

Galectins

Galectins are β -galactosyl-binding lectins that require a reducing environment for binding activity but, unlike CTLs and some FTLs, do not require Ca^{2+} (Vasta and Ahmed 2008; Vasta et al. 2012b). Although galectins are structurally conserved and taxonomically widely distributed, they display a remarkable functional diversity by participating in developmental processes, cell adhesion and motility, regulation of immune homeostasis, and recognition of glycans on the surfaces of viruses, bacteria, and protozoan parasites (Vasta 2009). On the basis of their primary structure and subunit organization, mammalian galectins are classified as "proto," "chimera," and "tandem-repeat" types (Vasta and Ahmed 2008; Vasta 2009; Vasta et al. 2012b). Prototype galectins contain one CRD per subunit and are usually homodimers of noncovalently linked subunits. The chimera-type galectins have a single C-terminal CRD, like the prototype, and a non-CRD N-terminal domain that mediates the formation of trimers and pentamers. In contrast, the tandem-repeat galectins, in which two CRDs are joined by a linker peptide, are monomeric.

743 Molluscan galectins are less diversified than those in mammals but also show
744 different domain organizations, carbohydrate specificity for blood group oligosac-
745 charides, and upregulation of expression by infectious challenge, a feature that sup-
746 ports their proposed role in innate immune responses (Tasumi and Vasta 2007; Feng
747 et al. 2013, 2015; Kurz et al. 2013; Vasta et al. 2015). In contrast to vertebrates, the
748 identification and characterization of galectins in aquatic molluscs has been rela-
749 tively recent, with most of the studies being aimed at the identification of their
750 transcripts or proteins in diverse tissues and cell types, including hemocytes, and the
751 assessment of their expression upon environmental or infectious challenge (Yamaura
752 et al. 2008; Yoshino et al. 2008; Song et al. 2010, 2011; Zhang et al. 2011a; Bao
753 et al. 2013; Dheilly et al. 2015; Bai et al. 2016). In the eastern oyster, *C. virginica*,
754 however, the galectins CvGal1 and CvGal2 have been characterized in their detailed
755 molecular, structural, and functional aspects (Tasumi and Vasta 2007; Feng et al.
756 2013, 2015; Kurz et al. 2013). As a result, unique features of the galectin repertoire
757 of aquatic molluscs have become apparent, such as their domain organizations, as
758 well as structural and functional aspects (Vasta et al. 2015). CvGal1 and CvGal2
759 carry four canonical galectin CRDs (Tasumi and Vasta 2007; Feng et al. 2013,
760 2015), a domain organization that does not conform to any of the galectin types
761 described in vertebrates (Vasta and Ahmed 2008; Vasta et al. 2012b). Since then,
762 galectins have been identified in an increasing number of aquatic mollusc species,
763 including both bivalves and gastropods, and can be classified, in the vast majority of
764 cases, into the 2-CRD and 4-CRD types (Vasta et al. 2015). As revealed by a phylo-
765 genetic analysis, these galectin types are ancient, as they were already present in the
766 most recent common ancestor of both bivalves and gastropods (Vasta et al. 2015).
767 From the functional standpoint, CvGal1 can recognize microbial pathogens and
768 parasites and promote their phagocytosis, but it can also selectively bind to phyto-
769 plankton components, suggesting its participation in uptake of microalgae (Tasumi
770 and Vasta 2007). Furthermore, recent studies suggest that the protozoan parasite
771 *P. marinus* has adapted to subvert the oyster's innate immune/feeding recognition
772 mechanisms to gain entry into the host cells by being preferentially recognized by
773 CvGal1 and CvGal2 over algal food or bacterial pathogens (Tasumi and Vasta 2007;
774 Feng et al. 2013, 2015; Kurz et al. 2013; Vasta et al. 2015).

775 Fibrinogen-Related Domain-Containing Proteins

776 A class of proteins containing a C-terminal fibrinogen-related domain (FReD), and
777 similar to vertebrate ficolins, has gained a significant amount of attention in mol-
778 luscs. Because of their important role in the resistance of the snail *B. glabrata* to
779 trematode infection, together with their somatic sequence diversification (Adema
780 et al. 1997; Adema 2015; Gordy et al. 2015), a subclass of FReD-containing pro-
781 teins (which also contain one or two immunoglobulin-like domains), named
782 fibrinogen-related proteins (FREPs), have been studied as one of the first examples
783 in support of immune memory in invertebrates (Milutinović and Kurtz 2016).
784 Unlike fibrinogen chains, these lectin-like molecules are primarily involved in

immune recognition and are not linked to coagulation (Hanington and Zhang 2011). While these immune properties have been extensively documented in snails since the 1990s (as reported in detail in the section “Expansion and Diversification of Innate Immune Gene Families” in Chap. 12), the first studies of FReD-containing proteins in bivalve molluscs are quite recent.

The first indications pointing toward an involvement of bivalve FReD-containing proteins in immune recognition came from the upregulation of AiFREP in the scallop *Argopecten irradians* in response to *V. anguillarum* but not to *Micrococcus luteus* infections. The recombinant protein could agglutinate Gram-negative and Gram-positive bacterial cells, confirming AiFREP as a reasonable soluble PRR candidate (Zhang et al. 2009b). Years later, AiFREP-2 was functionally characterized in the same species, confirming and to some extent even extending the marked recognition properties of these two scallop proteins (Yang et al. 2014). Very similar results were obtained in *Magallana hongkongensis*, where the recombinant protein ChFCN could selectively bind different bacterial species, agglutinate *Escherichia coli* cells, and enhance hemocyte phagocytosis in vitro (Xiang et al. 2014b). Purified *M. galloprovincialis* transcripts encoding FReD-containing proteins were upregulated in mussels by multiple challenges and could similarly improve the phagocytic rate of hemocytes (Romero et al. 2011). Indirect indications supporting the immune involvement of FReD-containing proteins have been also collected from transcriptomic studies in QPX-infected *M. mercenaria* (Wang et al. 2016b) and *V. splendidus*-infected *Mytilus edulis* hemocytes (Tanguy et al. 2013).

Early sequence database mining approaches revealed that FReD-containing proteins are part of a large multigene family in *Mytilus* spp. (Gorbushin and Iakovleva 2011), and it is now well recognized that the genome of several bivalve species encodes more than 100 such genes, which are, for the most part, expressed in the hemocytes, gills, and digestive gland (Zhang et al. 2015; Huang et al. 2015a; Gerdol and Venier 2015). Bivalve FReD-containing proteins are characterized by a simpler domain organization than snail FREPs, as they lack N-terminal immunoglobulin domains, which are thought to play a fundamental role in somatic mutation (Gerdol 2017). Comparative genomics analyses have further revealed that the Ig-FReD domain combination is exclusively found in heterobranch gastropods (Gorbushin et al. 2010). In most cases, bivalve proteins contain a single FReD associated with a coiled coil region, which probably allows oligomerization (Skazina and Gorbushin 2016). In addition, while the process of somatic mutation in snail FREPs is supported by experimental evidence, no data have been provided yet to sustain a similar mechanism in bivalve FReD-containing proteins, which are however characterized by a relevant sequence diversity. This topic has been investigated in detail in *M. gigas*, where the occurrence of polymorphisms in five of these transcripts was originally attributed to allelic recombination or somatic diversification (Zhang et al. 2012b). However, the large number of FReD genes in bivalves suggest that some of these variants might be the result of recent duplications or interindividual sequence variability, mirroring the evolutionary patterns observed for C1q domain-containing (C1qDC) proteins and other expanded PRR families (Huang et al. 2015a).

The remarkable immune properties of FReD-containing proteins, together with their remote sequence similarity with vertebrate ficolins, suggest that these secreted

830 PRRs are somehow involved in the lectin pathway of the bivalve complement
 831 system (see section “[Evidence of an Ancient Complement System in Bivalves?](#)”)
 832 (Gerdol and Venier 2015; Wang et al. 2017b). However, definitive proof in support
 833 of this hypothesis remain to be collected, in particular for what concerns the identi-
 834 fication of mannose-binding protein-associated serine proteases (MASPs)—
 835 essential mediators of the complement system, which have not been identified yet
 836 in molluscs.

837 C1q Domain–Containing Proteins

838 Some Insights into the Massive Gene Family Expansion of C1q 839 Domain–Containing Proteins

840 Although the outstanding binding potential of the C1q domain allows high func-
 841 tional versatility in the recognition of different ligands, no metazoan taxa seem to
 842 have exploited these properties to the same extent as bivalve molluscs. The genomes
 843 of these animals encode several hundred secreted proteins containing this conserved
 844 domain at their C-terminal end, collectively known as C1q domain–containing
 845 (C1qDC) proteins. The immune properties of the C1q domain, whose structural fold
 846 is exemplified in Fig. 8, have been well documented from the study of the vertebrate

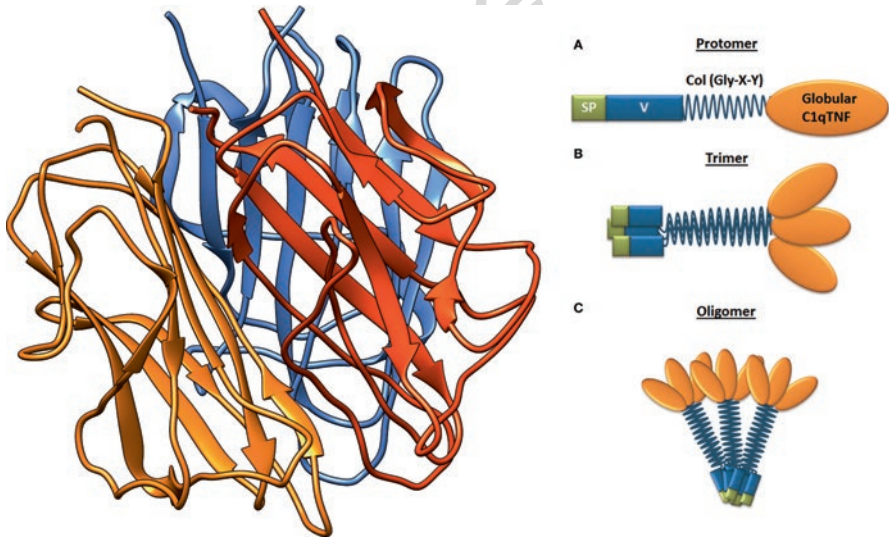


Fig. 8 Left: Three-dimensional structure of the three chains of the human C1q globular head (PDB accession ID: 2WNU; C1qa, C1qb, and C1qc chains are colored in orange, red, and blue, respectively). Right: Prototypical organization of vertebrate C1qDC proteins: **a** single protomer, comprising a signal peptide (SP), followed by a variable region (V, which might be absent in bivalve molluscs), a collagen region (usually replaced by a coiled coil domain in bivalve molluscs), and the globular C-terminal C1q domain. Promomers can assemble into trimers (**b**) and form higher-order bouquet-like structures (**c**). (Source: Thanasupawat et al. 2015)

[AU13](#)

[AU14](#)

complement system, where it is the major structural unit in the three chains of the C1q complex. However, the first indications pointing toward a similar role in molluscs only surfaced in 2004, with the isolation of a sialic acid-binding lectin from the garden snail *Cepaea hortensis* (Gerlach et al. 2004).

In bivalves, C1qDC proteins were first tentatively linked to pathogen recognition because of their high sequence diversity, exemplified by the identification of 168 different transcripts in *M. galloprovincialis* which, for the most part, strikingly displayed hemocyte specificity (Gestal et al. 2010; Gerdol et al. 2011), and the presence of over 300 genes in the Pacific oyster genome (Gerdol et al. 2015b). While most vertebrate C1qDC proteins, including those involved in the complement system, contain a central collagen region required for oligomerization (Fig. 8), about half of the oyster C1qDC proteins contain a coiled coil region, possibly exerting a function homologous to that of collagen. A relevant number of the other members of this gene family, however, lack oligomerization motifs and contain only an N-terminal signal for secretion followed by a globular head C1q domain, identifying the sgC1q subfamily. Surprisingly, just a few gene products have shown an association with additional domains; among these, the most notable example is provided by proteins containing multiple consecutive C1q domains (Gerdol et al. 2015b).

Another interesting finding was that such a massive expansion and diversification event occurred in Pteriomorphia and Heterodonta but not in the two other major subclasses, Palaeoheterodonta and Protobranchia, which possess only a few C1qDC genes, like most other protostomes (including nonbivalve molluscs). This lineage-restricted expansion event might have had important biological implications in mussels, clams, oysters, and scallops, providing these marine organisms with an unparalleled array of recognition molecules to be potentially used in microbe-associated molecular pattern (MAMP) recognition (Gerdol et al. 2015b). Another key piece in the puzzle of the evolution of bivalve C1qDC proteins was provided by the genome of the Manila clam, *Ruditapes philippinarum*. Indeed, most of the sequences 1589 C1qDC genes found in this clam appear to be unrelated to those found in oyster, thereby suggesting that the astounding molecular diversity in the two species derives from independent evolution (Mun et al. 2017).

Functional Studies Are Progressively Revealing the Immune Functions of C1q Domain-Containing Proteins

Genomic investigations are, however, insufficient in the absence of a functional characterization to link this expansion event to improved immune functions. Confirmations, in this sense, have been provided by different experimental approaches, i.e., gene expression studies that have evidenced the upregulation of oyster C1qDC transcripts in response to Rickettsia-like organisms and revealed their implication in the response to Brown Ring Disease, *P. olseni*, and QPX infections in clams (Xu et al. 2012; Leite et al. 2013; Allam et al. 2014; Wang et al. 2016b). Experimental challenges have further demonstrated that many bivalve C1qDC genes are induced by infection with various Gram-positive and

891 Gram-negative bacteria, as well as by fungi (Kong et al. 2010; Gestal et al. 2010;
892 Li et al. 2011a; Gerdol et al. 2011; Jiang et al. 2015), but also by direct stimulation
893 with LPS, PGN, β -glucan, and polyI:C (Wang et al. 2012a, b, 2015a; Yang et al.
894 2012), altogether reinforcing their role as PRRs. The indications collected from
895 gene expression studies were later confirmed by the binding properties of C1qDC
896 recombinant proteins toward LPS, PGN, polyI:C, mannan, β -1,3-glucan, and
897 yeast glucan (Wang et al. 2012a, 2015a; Jiang et al. 2015) as well as toward live
898 bacteria (Wang et al. 2015a; Zhao et al. 2016a; Huang et al. 2016).

899 From a functional point of view, an oyster recombinant C1qDC protein was
900 capable of significantly inhibiting the growth of Gram-positive and Gram-negative
901 bacteria (He et al. 2011), and others displayed strong agglutinating activity toward
902 Gram-positive bacteria, Gram-negative bacteria, and fungi, with a certain degree of
903 selectivity (Kong et al. 2010; Wang et al. 2012a). Some studies have tried to better
904 elucidate the mode of action of bivalve C1qDC proteins and their connection with
905 other molecular components of the immune system. For example, the bactericidal
906 properties of mussel hemolymph appear to be mediated by a C1qDC serum opsonin
907 that binds bacterial D-mannose, promoting the phagocytic action of hemocytes
908 (Pezzati et al. 2015). Similarly, a protein isolated from the scallop *Azumapecten far-*
909 *reri* is capable of enhancing the phagocytosis of invading *E. coli* cells (Wang et al.
910 2012b), and an oyster LPS-binding C1qDC protein could sensibly boost this activ-
911 ity toward *E. coli* and *V. splendidus* (Jiang et al. 2015). Furthermore, other recombi-
912 nant proteins are able to interact with heat-aggregated human IgGs and IgMs (Wang
913 et al. 2015a), providing novel and stimulating insights into the possible involvement
914 of these components in the activation of the prototypical complement system of
915 bivalve molluscs (see section “Evidence of an Ancient Complement System in
916 Bivalves?”).

917 Although bivalve C1qDC proteins were initially considered as hemocyte-specific
918 products, it is now clear that they are broadly expressed in all main tissues, with a
919 particular prevalence in the gills or in the digestive gland (Gerdol et al. 2015b), leav-
920 ing some open questions concerning their involvement in functions other than
921 immune recognition. In fact, the extreme diversification and binding properties of
922 these proteins would allow, in line of principle, additional physiological functions,
923 which are progressively starting to emerge.

924 Evidence of an Ancient Complement System in Bivalves?

925 A Brief Description of the Complement System

926 Despite the highly divergent evolutionary strategies adopted by metazoans to
927 develop an efficient immune system in highly diverse life environments, complex
928 molecular machinery of the utmost importance in pathogen recognition and clear-
929 ance is surprisingly conserved in nearly all animals. This protein complex, able to
930 enhance recognition and removal of microbial cells by recruiting the main players
931 of the vertebrate immune system (phagocytic cells and immunoglobulins), has been
932 named the “complement” system.

The complement system can be potentially activated by different biochemical pathways, which involve components of both innate and adaptive immunity, and has thereby been defined as a functional link between these two major branches of the immune system (Dunkelberger and Song 2009). In vertebrates, the different routes that can lead to complement activation involve either the binding of C1q to antigen-complexed M or G immunoglobulins (the classical pathway), the recognition of MAMPs by mannan-binding lectins (MBLs) and ficolins (the lectin pathway), or the direct recognition of MAMPs by C3b following spontaneous C3 hydrolysis (the alternative pathway) (Fig. 9). Overall, complement activation triggers, through a proteolytic cascade, the opsonization of invading microbes, their lysis by the action of the membrane attack complex (MAC), and the recruitment of phagocytic cells for their final elimination.

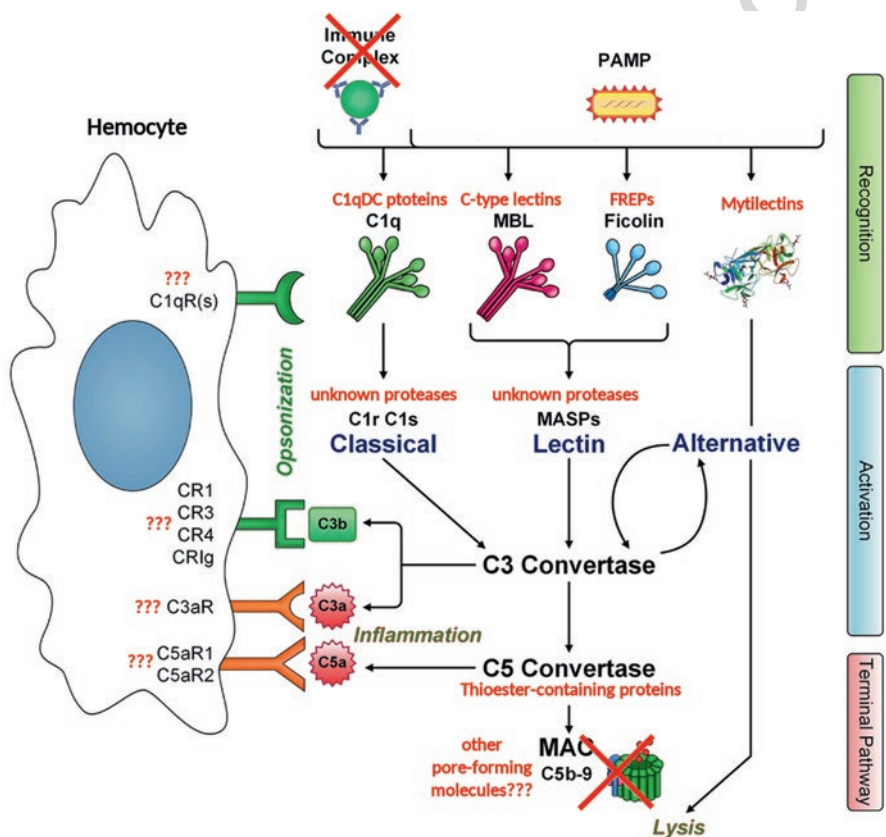


Fig. 9 Overview of the complement system in bivalves and comparison with vertebrates. The vertebrate molecular players are shown in black and the bivalve homologous components are indicated in red, whenever needed. Components that are absent in bivalves (namely, the membrane attack complex and antigen-complexed immunoglobulins) are struck through. (Edited from Bohlson et al. (2014))

945 **The Conserved “Core Components”: C2 and C3**

946 With the exception of Ecdysozoa, the near universal conservation of two core molecu-
947 lar components of the complement system—C3 and C2/factor B—suggest that a pro-
948 totypical complement system was present in the common ancestor of all metazoans
949 (Smith et al. 1999; Pinto et al. 2007). Accordingly, genes encoding these two highly
950 conserved elements are also readily identifiable in most bivalve genomes and tran-
951 scriptomes (Moreira et al. 2012a; Zhang et al. 2014c; Gerdol and Venier 2015). Their
952 first formal description was provided in the grooved carpet shell, *Ruditapes decussa-*
953 *tus* (Prado-Alvarez et al. 2009). The C3 component of the razor clam *Sinonovacula*
954 *constricta* was strongly upregulated in hemocytes and digestive gland upon bacterial
955 challenges. In addition, the serum of *S. constricta* was activated by LPS and bacteria,
956 confirming that the function of the bivalve protein was highly homologous to verte-
957 brates (Peng et al. 2016). Further confirmation was recently provided by the use of
958 polyclonal antibodies directed toward three distinct fragments of the Pacific oyster C3
959 protein, homologous to the α , β , and γ chains obtained in vertebrates from the proteo-
960 lytic cleavage of the C3 precursor. The observation of a single band recognizable in
961 serum under non-reducing conditions, as opposed to the presence of three distinct
962 bands of 110, 60, and 30 KDa under reducing conditions, pointed out that bivalve C3
963 molecules are processed by serum proteases in a similar fashion to what happens in
964 animals with a canonical complement system (Wang et al. 2017b).

965 The bivalve complement system might also involve thioester-containing proteins
966 (TEPs), accessory complement proteins that share a high degree of similarity with
967 C3/C4/C5 and promote opsonization of invading microbes and their elimination by
968 phagocytosis in other invertebrates (Blandin et al. 2008; Bou Aoun et al. 2010).
969 TEPs have been functionally characterized only in the scallop *A. farreri*, where they
970 possess a highly variable central region produced by the alternative splicing of six
971 mutually exclusive exons. This sequence variation appears to cover a key role in the
972 specificity of the immune response to be triggered, as the amount of the isoforms
973 produced largely varies on the basis of the type of challenge and the sex of the speci-
974 mens (Zhang et al. 2009c). A very recent study went into the subject in depth, evi-
975 dencing that like C3, scallop CfTEP undergoes fragmentation due to the action of
976 endogenous serum proteases (Xue et al. 2017b).

977 **Present Uncertainties and Future Directions**

978 The absence of immunoglobulins rules out the existence of the classical pathway
979 of the complement system in animals lacking an adaptive immune system, which
980 include bivalve molluscs. At the same time, the remote homology between vertebrate
981 C1q, ficolins, and MBLs, and similar sequences in invertebrate organisms, further
982 complicates the interpretation of the functional overlap between the lectin pathway
983 of the complement system between vertebrates and invertebrates. However, the high
984 diversification of C1qDC proteins might potentially provide a very broad potential
985 of recognition toward MAMPs, even in absence of immunoglobulins. At the same
986 time, while no bona fide sequence that is homologous to vertebrate MBLs or fico-
987 lins is present in molluscs, both C-type lectins and FReD-containing proteins (see
988 sections “[The Role of Lectins in Immune Recognition](#)” and “[Fibrinogen-Related](#)

Domain (FReD)-Containing Proteins”) underwent massive expansion and diversification events similar to C1qDC proteins. This further reinforces the idea that bivalves possess an astoundingly complex arsenal of soluble PRRs, which are possibly part of a complement lectin pathway. However, it is presently unclear how their recognition signals would converge to C3, as no clear homologs to MASP-1, MASP-2, C1r, and C1s serine protease, required for downstream activation of C3 in vertebrates, are present in bivalves (Gerdol and Venier 2015).

Altogether, these reports support the existence of a prototypical complement system in bivalve molluscs, therefore expanding the taxonomic distribution of this ancient immune defense system to Lophotrochozoa, in addition to echinoderms, horseshoe crabs, tunicates, and amphioxus. However, many uncertainties remain about the modes of activation of this system, and some of the hypothetical molecular players that are expected to be involved still remain to be identified. The mechanism of regulation of the complement system in oysters in response to LPS has been hypothesized in a recent study. The authors suggested that 12 serine protease domain-containing proteins might somehow play a key role in complement activation, and they further identified some possible C3 receptors containing integrin α/β domains and similar to ascidian C3 receptors (Wang et al. 2017b).

Finally, it is presently difficult to assess whether the final outcome of this process is simply the opsonization of pathogenic cells, which would facilitate their elimination by the recruitment of phagocytic cells, or whether it also involves lytic components functionally homologous to the membrane attack complex. As will be discussed in section “Lysozymes, BPIs and Other Pore-Forming Molecules,” while the constituents of the terminal pathway of the complement system appear to have been specifically developed in the vertebrate lineage, it is possible that other divergent pore-forming molecules function in a similar manner, sometimes combining MAMP-sensing and pore-forming properties within the same protein precursor.

Toll-Like Receptors 1016

Structure and Function of Toll-Like Receptors 1017

Toll-like receptors (TLRs) are metazoan immune receptors, which have found major evolutionary success. Because of their ability to recognize a broad range of ligands, TLRs are important players of the innate immune system of both vertebrate and invertebrate animals, functioning as MAMP sensors either on the plasma membrane or in endosomal compartments. The recognition properties of TLRs are provided by several extracellular leucine-rich repeats (LRRs), which can be organized either in a single cysteine cluster (scc) or in a multiple cysteine cluster (mcc) configuration, whereas the transduction of the immune signal occurs thanks to an intracellular TIR (Toll–interleukin receptor) domain (Fig. 10). This conserved signaling module is separated from the extracellular LRRs by a short transmembrane α -helical domain, which anchors TLRs to cell membranes.

The prototypical Toll protein of the fruit fly *Drosophila melanogaster*, after which all TLRs are named, is a multifunctional protein, acting both as a primary

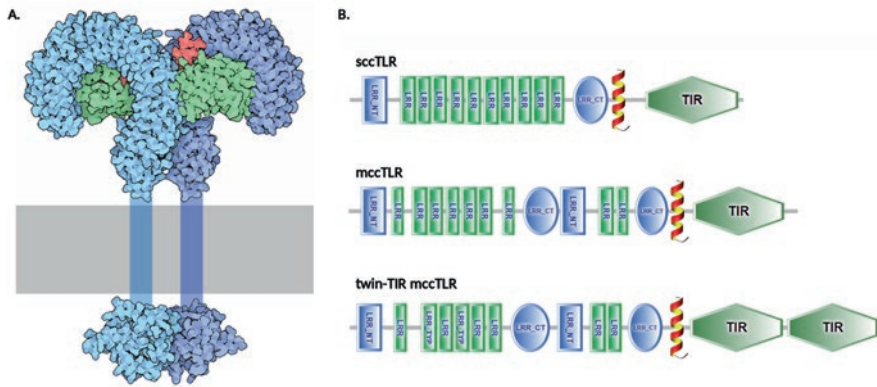


Fig. 10 (a) Structure of the human Toll-like receptor 4 dimer (blue) bound to bacterial lipopolysaccharide (red) through its extracellular LRR domains. The transmembrane region is shown schematically. (Image courtesy of RCSB PDB, <http://pdb101.rcsb.org/motm/143>). The intracellular TIR domain is shown on the inner side of the cell membrane. (b) Schematic domain organization of single cysteine cluster (scc), multiple cysteine cluster (mcc), and twin-TIR mcc Toll-like receptors found in bivalve molluscs

1031 determinant of embryonic dorsal–ventral polarity and as the receptor for the proin-
 1032 flammatory cytokine Spätzle. However, most of the TLRs described so far in verte-
 1033 brates function exclusively as immune receptors by directly recognizing LPS, PGN,
 1034 foreign nucleic acids, and other MAMPs without the mediation of cytokine-like
 1035 molecules. While the organization of TLRs has long been considered to be similar
 1036 to that of *Drosophila*, genomic studies have progressively unearthed some impor-
 1037 tant peculiarities that strikingly differentiate arthropods from all other animals. In
 1038 particular, echinoderms have developed an arsenal of immune receptors that are
 1039 potentially capable of recognizing a very broad range of invading microorganisms
 1040 (Buckley and Rast 2012).

1041 **The Emerging Role of Toll-Like Receptors in Bivalve Molluscs**

1042 Besides echinoderms, the massive expansion of the TLR repertoire by gene duplica-
 1043 tion involved other phyla, including molluscs (Gerdol et al. 2017), as most notably
 1044 evidenced by the identification of 83 TLR genes in the genome of the Pacific oyster
 1045 (Zhang et al. 2015). However, the genomic expansion of the bivalve TLR gene fam-
 1046 ily occurred independently from that of sea urchins, as it mostly targeted a group of
 1047 phylogenetically distinct genes. Because of the high molecular diversification of
 1048 bivalve TLR sequences, a novel uniform nomenclature has been recently suggested
 1049 to avoid confusion in the discussion of the functional properties of these receptors
 1050 (Zhang et al. 2015; Gerdol et al. 2017). Thus, it has been suggested that bivalve
 1051 receptors should be categorized as P-type, sPP-type, or twin-type (in the case of
 1052 mccTLRs), or as V-type or sP-type (in the case of sccTLRs) (Fig. 10). V-type TLRs,
 1053 present in hundreds of members in the sea urchin genome, include only a few
 1054 sequences in bivalve molluscs, where most TLRs are ascribable to the sP-type
 1055 expanded group (Gerdol et al. 2017).

CfToll-1 was the first TLR to ever be described in bivalve molluscs, providing the first pieces of evidence in support of the possible involvement of these receptors in bivalve immune recognition. Indeed this TLR, identified in the scallop *A. farferi* and pertaining to the P-type subfamily, is mildly upregulated by LPS challenges, pointing out a role in the detection of Gram-negative bacteria (Qiu et al. 2007). Following this initial report, several gene expression studies have implicated TLRs in the immune response to different types of microbes and associated pathologies. For example, a single TLR was strongly modulated in QPX-infected *M. mercenaria* (Perrigault et al. 2009) and in *P. marinus*-infected *C. virginica* oysters (Tanguy et al. 2004). Finally, TLRs have been also reported to be upregulated in response to *V. alginolyticus* challenges in different marine clam and mussel species (Moreira et al. 2012b; Martins et al. 2014).

These observations encouraged the design of targeted functional experiments aimed at identifying the microorganisms recognized by bivalve TLRs and their possible ligands. The most significant studies have been carried out in (1) *M. gigas*, where a TLR was found to be strongly induced by *V. anguillarum* challenges (Zhang et al. 2011c) and a second one (CgTLR6) displayed binding ability toward Gram-positive and Gram-negative bacteria, further revealing affinity to LPS and PGN but not to mannan (Wang et al. 2016b); (2) *Hyriopsis cumingii*, where three different TLRs, responsive to distinct microbial challenges, have been identified, pointing out a remarkable functional specialization (Ren et al. 2013, 2014; Zhang et al. 2017); (3) the noble scallop, *Mimachlamys nobilis*, where an sccTLR responded to *V. parahaemolyticus*, LPS, and PolyI:C challenges in hemocytes (Lu et al. 2016); and (4) *M. galloprovincialis*, where the upregulation of the P-type TLR MgTLR-i could be observed in response to *Vibrio* spp. and *M. luteus* but not to *Fusarium oxysporum* injection (Toubiana et al. 2013). The high selectivity of TLRs, in terms of both transcriptional responsiveness and binding potential, has been further confirmed by the transcriptional analysis of the entire complement of oyster TLR genes, which often responded to just a single pathogenic challenge in a highly specific manner (Zhang et al. 2015).

One of the most praiseworthy studies aimed at clarifying the placement of these receptors in the molecular networks of immune signaling targeted four different sccTLRs in *M. gigas* and permitted demonstration of their participation in the activation of nuclear factor kappa B (NF- κ B). The finding that oyster sccTLRs are localized both on the plasma membrane and in late endosomal vesicles was equally important, as it revealed a possible role of TLRs also in the modulation of immune response upon phagocytosis of invading microbes (Zhang et al. 2013a). Although only little effort has so far been put into the identification of the effector molecules whose production is controlled by TLRs, preliminary results clearly point toward a key role of TLR signaling in the regulation of AMP and lysozyme production through a MyD88-dependent pathway (see section “Canonical TLR Signaling”).

The experimental data collected so far confirm that the fundamental role of TLRs in the bivalve immune response to invading microorganism appears to be supported by overwhelming evidence. However, one might wonder whether this large family of receptors has acquired additional physiological roles due to neofunctionalization,

1101 as has been suggested for other bivalve recognition protein families. While evidence
1102 in support of this hypothesis still remains scarce, some reports hint that TLRs might
1103 be modulated by other stimuli, i.e., biotoxins (Detree et al. 2016b), abiotic stress
1104 (Zhang et al. 2015), and variations of pH (Xing et al. 2017).

1105 Other Membrane-Bound Immune Receptors

1106 Peptidoglycan Recognition Proteins

1107 Peptidoglycan recognition proteins (PGRPs) are a class of well-characterized PGN-
1108 binding molecules that, in the fruit fly *D. melanogaster*, comprises both membrane-
1109 bound and secreted members. Membrane-bound PGRPs are directly involved in
1110 MAMP recognition during infections by Gram-negative bacteria and activate the
1111 Immune deficiency (IMD) signaling cascade (Royet and Dziarski 2007). On the
1112 other hand, secreted PGRPs cooperate with Gram-negative Binding Proteins
1113 (GNBPs) in the extracellular environment, triggering the prophenoloxidase cas-
1114 cade, which leads to the activation of Toll signaling (see section “Canonical TLR
1115 Signaling”) and melanization (see section “The Phenoloxidase Cascade”). While
1116 PGRPs are also present in vertebrates, they are not anchored to the plasma mem-
1117 brane and they mostly exert bactericidal/bacteriostatic activity in the extracellular
1118 environment (Montaño et al. 2011).

1119 PGRPs have been functionally characterized in detail in arthropods and verte-
1120 brates, but nearly no information is available for the other major animal phyla. In
1121 bivalve molluscs, genome and transcriptome screenings show the presence of both
1122 membrane-bound and secreted PGRPs, even though large margins of uncertainty
1123 remain about their functional overlap with arthropods and vertebrates. First, there is
1124 no evidence in support of an extracellular pathway homologous to that of the
1125 *Drosophila* prophenoloxydase proteolytic cascade, and the absence of Spätzle-like
1126 proteins make it highly doubtful that secreted PGRPs participate in TLR activation
1127 in bivalves (see section “The Phenoloxidase Cascade”). Second, the high sequence
1128 divergence between bivalve PGRPs and those from other organisms does not allow
1129 similarity-based functional inference (Gerdol and Venier 2015).

1130 The first report of PGRPs in bivalve molluscs, in the form of a short secreted
1131 protein, dates back to 2007, when an inducible gene product was identified in the
1132 scallop *A. farreri* following Gram-positive and Gram-negative bacterial challenges
1133 (Su et al. 2007). This finding was later confirmed in *M. galloprovincialis*,
1134 *Bathymodiolus azoricus* (Martins et al. 2014), and *H. cumingii*, where broad-
1135 spectrum antibacterial activity and lytic activity toward both Lys-PGN and DAP-
1136 PGN were demonstrated (Yang et al. 2013c). Furthermore, another study reported
1137 the modulation of the expression of two secreted short PGRPs in *Solen grandis*, in
1138 particular, in response to PGN but not LPS (Wei et al. 2012), confirming previous
1139 results concerning PGN specificity obtained in the bay scallop (Ni et al. 2007).
1140 Finally, another secreted PGRP molecule from *M. gigas* displays a unique domain
1141 architecture, as it combines the PGN-binding domain with a G-type lysozyme
1142 domain, which could potentially enable the coexistence of bacterial recognition and

killing properties in the same molecule (Itoh and Takahashi 2009) (see section “Lysozymes, BPIs and Other Pore-Forming Molecules”). Overall, the vast majority of the studies that have targeted bivalve secreted PGRPs so far are seemingly concordant in attributing to them functional properties more similar to those of vertebrate PGRPs than to those of arthropods. Their cooperation with GNBP and their involvement in the activation of TLRs seem unlikely at this point.

Interestingly, while no membrane-bound PGRP has been functionally characterized yet in bivalves, at least two proteins of this type are present in the Mediterranean mussel transcriptome. Together with the contemporary identification of some conserved intracellular mediators, this prompted researchers to hypothesize the possible existence of an IMD-like pathway (see section “Other Immune Signaling Pathways”) (Gerdol and Venier 2015). While this hypothesis still awaits experimental confirmation, a recent study carried out in *B. azoricus* identified five paralogous PGRP genes, which were connected to the regulation of bacterial endosymbiosis in gills (Détrée et al. 2017).

Recently Discovered Receptors

Besides TLRs and PGRPs, only a very few other cases of PRRs anchored to the extracellular surface of bivalve immune cells have been studied so far. The most relevant are the Nimrod-like receptor (CgNimC) and LRR and Ig domain-containing proteins (LRRIGs), both identified in *M. gigas*. The former receptor has been implicated in the recognition of Gram-negative bacteria because of its relevant upregulation in response to *Vibrio* spp. challenges and LPS binding. Further functional assays established that CgNimC plays a fundamental role in regulating the phagocytic rate of hemocytes toward invasive Gram-negative bacteria (Wang et al. 2015d). On the other hand, the two LRRIGs genes identified in the genome of *M. gigas* encode large proteins bearing extracellular LRRs (like TLRs), coupled with an immunoglobulin-like domain, a transmembrane domain, and a short uncharacterized cytosolic C-terminal domain. Immunoglobulin-like domains are abundant in bivalve genomes, and their marked immunological properties have been well defined in vertebrates and, partly, also in invertebrates (e.g., gastropod FREPs; see Chap. 12, section “Defense-Associated Humoral Components”). LRRIGs can bind a broad range of MAMPs and are upregulated in hemocytes in response to various types of challenges. Furthermore, they can modulate the expression of cytokine-like factors (i.e., TNF and IL-17) and promote hemocytic phagocytosis of *Vibrio* cells, thereby reinforcing their position as key regulators of immune response in oysters (Wang et al. 2017a; Huang et al. 2018).

Cytosolic Pattern Recognition Receptors

In comparison with the impressive amount of literature produced about soluble and membrane-bound PRRs, it is perhaps surprising that only a handful of studies have so far taken into account the possible involvement of cytosolic receptors in the immune system of bivalves. Most of the molecular players described below have

1184 been identified just at the sequence level and therefore emerge as interesting targets
1185 for future functional investigations.

1186 Different intracellular PRRs are potentially capable of recognizing MAMPs
1187 present in the cytosol. These receptors have a dual function in: (1) directly detecting
1188 the presence of pathogens (e.g., viruses) in the cellular space; and (2) indirectly
1189 detecting microbes in the extracellular environment from their degradation products
1190 (e.g., peptidoglycan components). In summary, this system works in a synergistic
1191 manner with membrane-bound PRRs, thereby reinforcing the immune response
1192 through the combination of converging signaling routes derived from the intracel-
1193 lular and extracellular environments.

1194 **NACHT–Leucine-Rich Repeat Proteins and Bacterial Sensing**

1195 NACHT–leucine-rich repeat (NACHT-LRR) proteins (NLRs) act as sensors of the
1196 two major peptidoglycan-derived bacterial components, muramyl dipeptide (MDP)
1197 and γ -D-Glu-meso-diaminopimelic acid (iE-DAP) in the cytosol (Fritz et al. 2006).
1198 These MAMPs can be translocated inside the cytoplasm whenever bacteria pres-
1199 ent in the extracellular environment are attacked by antimicrobial effectors, or
1200 they can be released as a consequence of the digestion of phagocytosed bacterial
1201 cells. Activated NLRs oligomerize, recruiting adaptor molecules that can modulate
1202 immune response, cell death, or survival. Vertebrate NLRs are also responsible for
1203 the assembly of inflammasomes—large macromolecular complexes involved in the
1204 modulation of inflammation—which are however unlikely to exist in invertebrate
1205 animals (Latz et al. 2013).

1206 In spite of the great expansion of NLRs in many metazoans, no such receptor has
1207 ever been functionally characterized in molluscs. The typical tripartite domain
1208 architecture of NLRs comprises C-terminal leucine-rich repeats required for ligand
1209 binding, a central NACHT domain, which regulates oligomerization, and an
1210 N-terminal death fold domain (DFD), whose type (DEATH, DED, CARD, or PYD)
1211 determines the recruitment of specific downstream signaling adaptors. Although the
1212 single NLR-like protein identified in *M. galloprovincialis* displays a CARD/
1213 NACHT/LRR domain combination, it bears limited sequence homology with bona
1214 fine vertebrate NLRs, leaving its possible involvement in immunity a matter of
1215 speculation (Gerdol and Venier 2015).

1216 **RIG-Like Receptors: Fundamental Receptors of Viral Infection**

1217 While NLRs are mainly employed in bacterial sensing, a series of other receptors
1218 collectively known as RIG-like receptors (RLRs) cover an analogous function in the
1219 sensing of viruses. Upon activation, these helicase-like molecules trigger the antiviral
1220 response through the their N-terminal caspase recruitment domain (CARD)
1221 (Yoneyama and Fujita 2007). RLRs are capable of recognizing a broad range of
1222 dsDNA viruses, thanks to the mediation of DNA-dependent RNA polymerase III,
1223 which uses viral DNA as a template for the generation of 5' triphosphate single-
1224 stranded RNAs, which are efficiently recognized by the helicase domain of RLRs.

1225 Consistently with the expected rapid evolution of antiviral defense mecha-
1226 nisms in the continuous race to arms between the host and the pathogen, this

molecular machinery diverged significantly among animal groups (Paro et al. 2015). Bona fide RLRs were long thought to be exclusively present in vertebrates. However, following early reports of RLR-like genes in the genomes of cnidarians (Zou et al. 2009), a RLR highly responsive to poly(I:C) stimulation was also identified in *M. gigas* (Zhang et al. 2014e). Definitive proof about the involvement of RLRs in antiviral immunity was provided in a study demonstrating that the RLR CgRIG-I-1 was upregulated in response to OsHV-1 infection in Pacific oyster larvae, and that it could directly bind poly(I:C). The identification of the key adaptor protein IPS-1/MAVS (see section “Other Immune Signaling Pathways”), brought convincing evidence in support of the existence of an RLR-mediated signaling pathway activated in response to dsDNA viruses, closely matching that of vertebrates.

Another important aspect in the context of viral sensing is the possible involvement of Dicer, the main antiviral molecule in the cytosol of insect cells, which lack RLRs. In particular, only one out of the two Dicer gene copies present in the genome of *Drosophila* (Dicer-2) can process dsRNAs to produce siRNA (Lee et al. 2004), whereas the single mammalian Dicer gene is mostly involved in the production of miRNAs and only in some cell types can it generate siRNAs (Maillard et al. 2013). While the preferential substrates of this catalytic helicase in bivalves are presently unknown, all molluscs bear a single-copy Dicer gene (Rosani et al. 2016).

Stimulator of Interferon Genes: A Major Hub for Microbial Sensing in the Cytosol

The third major intracellular sensor of microbial infections is the Stimulator of Interferon Genes (STING). Unlike NLRs and RLRs, STING is a multifunctional protein, which can act either as a direct MAMP sensor or as a signaling adapter collecting infection signals derived from several pathogenic agents (Burdette and Vance 2013). This broad spectrum of recognition is guaranteed by the interaction with different cytosolic cofactors, whose presence in molluscs is mostly unconfirmed and sometimes even unlikely due to lineage-specific gene losses and high sequence divergence (Gerdol and Venier 2015).

In vertebrates, the dimerization and migration of STING from the endoplasmic reticulum membrane to the perinuclear region is a fundamental step for the subsequent activation of interferon response and inflammation (Ishikawa et al. 2009) (see section “Lysozymes, BPIs and Other Pore-Forming Molecules”). Although only a few reports have documented the existence of STING in bivalve molluscs (Gerdol and Venier 2015; He et al. 2015), the peculiar domain architecture of this molecule suggests a different subcellular localization and mode of action. Indeed, all lophotrochozoan STING molecules lack transmembrane domains and present a duplicated STING globular domain associated with a TIR domain; this structure could potentially enable self-dimerization upon ligand binding and the activation of downstream immune signaling through TIR–TIR heterotypic interactions. At the same time, it might imply important functional differences in comparison with vertebrates, including the interaction with different and presently unknown alternative MAMP cosensors.

1271 In any case, the main functional property of the STING globular domain, i.e.,
1272 the ability to bind cyclic dinucleotides in the cytosol, is expected to be retained.
1273 The most relevant ligands of STING are cyclic diguanylate (c-di-GMP) and cyclic
1274 guanosine monophosphate–adenosine monophosphate (cGAMP). While the former
1275 is a second messenger directly produced by bacteria, the latter is synthesized by
1276 cyclic GMP–AMP synthases (cGAS) whenever foreign DNA is detected in the
1277 cytoplasm, playing a fundamental role in the detection of both bacterial and viral
1278 nucleic acids (Ablasser et al. 2013). Although the importance of the cGAS/STING
1279 complex in activating the antiviral response has been only recently uncovered, it is
1280 certainly noteworthy that bivalve genomes display a significant expansion of cGAS-
1281 like genes in comparison with gastropods, which would suggest improved compe-
1282 tence for viral detection (Gerdol 2017).

1283 Signaling and Regulatory Pathways

1284 MAMPs of various natures, such as glycoproteins, components of cell walls and
1285 membranes, and exogenous nucleic acids can be recognized by the broad array of
1286 bivalve PRRs described in the previous sections, activating a cascade of intracellu-
1287 lar events that eventually result in cell response to the perceived stimulus. Multiple
1288 signal transduction pathways, mostly based on protein–protein interactions and
1289 modifications (e.g., kinase-mediated phosphorylation), regulate the timing and
1290 intensity of the immune response, as well the cellular fate (death or survival).

1291 Canonical Toll-Like Receptor Signaling

1292 The Essential Role of MyD88 in Immune Signal Transduction

1293 The main signal transduction pathway reported to mediate the immune responses of
1294 bivalve species is TLR/NF- κ B signaling (Fig. 11), which is deeply intertwined with
1295 other accessory networks that will be described in the section “[Other Immune](#)
1296 [Signaling Pathways](#).” The recognition of ligands by the extracellular LRR domains
1297 of TLRs leads to their dimerization, which in turn activates key transcription fac-
1298 tors, enabling the production of AMPs, lysozymes, interleukins (ILs), and other
1299 immune effectors against bacterial, fungal, and viral pathogens. The first essential
1300 step of TLR-mediated signal transduction involves the recruitment of TIR-DC
1301 adaptor proteins, which in vertebrates are primarily the myeloid differentiation
1302 primary response protein 88 (MYD88) and the TIR-domain-containing adapter-
1303 inducing interferon- β (TRIF) (O’Neill and Bowie 2007).

1304 Because of the lack of a TRIF homolog, the TLR signaling in bivalves is essen-
1305 tially a MyD88-dependent pathway, even though the possible involvement of
1306 alternative evolutionarily conserved TIR-DC adapters cannot be excluded (Gerdol
1307 et al. 2017). The fundamental signaling mediator MyD88 is characterized by an
1308 N-terminal death domain, required for perpetrating signal transduction, and by a
1309 C-terminal TIR domain that interacts upstream with the cytosolic TIR domain of

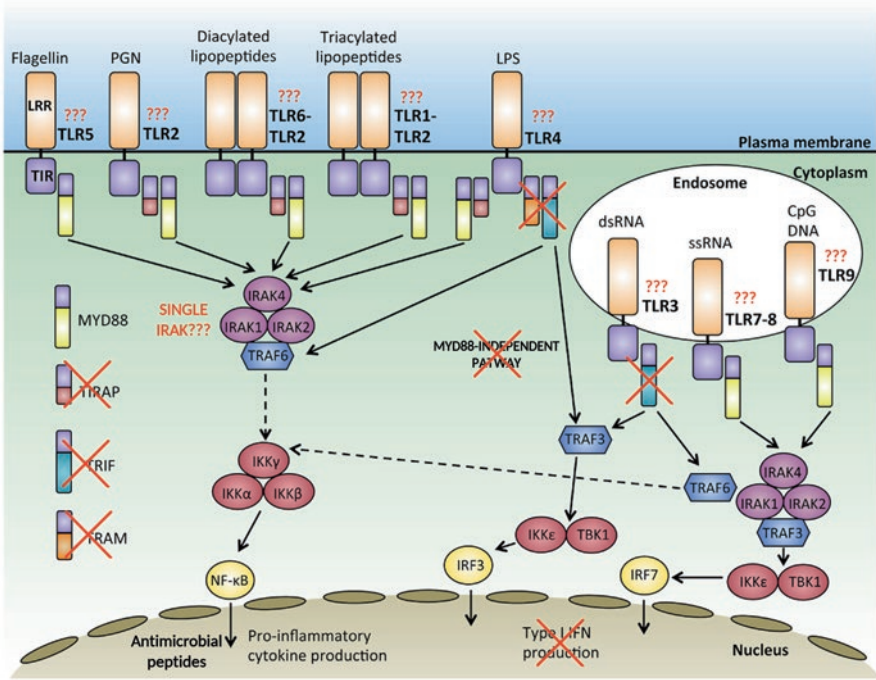


Fig. 11 Vertebrate canonical Toll-like receptor (TLR) signaling and comparison with that of bivalve molluscs. Unidentified components in bivalves are struck through and elements whose presence is uncertain are indicated by question marks. In particular, the low similarity between vertebrate and molluscan TLRs leaves the binding specificity of bivalve TLRs, for the most part, unknown. The homo- or heterodimerization of TLRs following ligand binding, either in the extracellular environment or in the endosomal compartment, recruits adaptor proteins, which propagate immune signals. Only MyD88, among the vertebrate adaptors, has been identified so far in bivalves. The recruitment and activation of IRAK kinases and the IKK complex results in the migration of the NF- κ B and possibly IRF transcription factors to the nucleus, where they regulate the production of proinflammatory cytokines and antimicrobial peptides. (Edited from Wang et al. 2014c)

TLRs. The upregulation of MyD88 transcripts has been documented in different bivalve species in response to various bacterial MAMPs (Toubiana et al. 2013; Ren et al. 2016; Xin et al. 2016a) and OsHV-1 infection in oysters (Renault et al. 2011; Du et al. 2013). The multiple MyD88 genes identified in the genomes of *M. gigas* and *M. yessoensis* indicate an expanded gene family (Zhang et al. 2015; Ning et al. 2015), possibly linked with the diversification of TLRs (see section “Toll-Like Receptors”). Some MyD88-like proteins lack the N-terminal death domain and are therefore thought to function as negative regulators (Xu et al. 2015b), together with the sterile alpha and armadillo motif containing protein (SARM), an evolutionarily conserved negative regulator of TLR signaling, as well as an intermediary of apoptosis and antiviral innate response (Belinda et al. 2008; Panneerselvam and Ding 2015).

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1322 Toll-Like Receptor–Mediated Signal Transduction: From the Cell 1323 Membrane to the Nucleus

1324 All of the expected elements of canonical MyD88-dependent TLR signaling have
1325 been identified in the transcriptomes of *M. gigas* (Zhang et al. 2011c), *M. gallopro-*
1326 *vincialis* (Toubiana et al. 2014), and *Saccostrea glomerata* (Ertl et al. 2016), and
1327 even physically mapped to *A. farreri* bacterial artificial chromosomes by fluores-
1328 cence in situ hybridization (Wang et al. 2011b; Zhao et al. 2015). These approaches
1329 highlighted a remarkable similarity with the immune signaling system of deuterostomes and a less significant overlap with arthropods. The immune role of such
1330 molecules has been confirmed by the assessment of their upregulation following
1331 immune stimulation trials and a detailed functional characterization in several
1332 bivalve species. While many accessory factors take part in this elaborate signaling
1333 network, either as positive or negative regulators, or as molecular switches to activate
1334 connected pathways, we will discuss below only the main molecular players
1335 (Fig. 11).
1336

1337 The second intracellular step of the MyD88-dependent TLR signaling involves
1338 the interaction between MyD88 and the *Interleukin-1 receptor-associated kinases*
1339 (IRAK)-1/-4 complex, with the subsequent recruitment of the *TNF receptor-*
1340 *associated factor 6* (TRAF6). The two IRAK proteins identified in mussels (both
1341 homologous to IRAK-4) were strongly overexpressed in hemocytes following bacterial
1342 challenges (Toubiana et al. 2014), similarly to the soft shell clam *Mya aren-*
1343 *aria* IRAK-4-like transcript, responsive to *V. splendidus* challenges (Mateo et al.
1344 2010). The turnover of IRAK kinases is regulated by the Toll interacting protein
1345 TOLLIP, characterized as an acute phase protein in *M. yessoensis* (Zhang et al.
1346 2015) but present with steady expression levels in *M. galloprovincialis* (Toubiana
1347 et al. 2014). TRAF6 is one of the key components of the pathway, as it regulates the
1348 activation of the IKK complex together with the *Transforming growth factor acti-*
1349 *vated kinase-1* (TAK1). TRAF6 responds to Gram-positive and Gram-negative, as
1350 well as to LPS challenges in the scallop *A. farreri* and in the mussel *M. galloprovin-*
1351 *cialis* (Wang et al. 2011b; Toubiana et al. 2014). Very limited functional information
1352 has been collected so far about TAK1, the associated proteins TAB1/2, and the com-
1353 ponents of the *Inhibitor of kappa-B kinase* (IKK) complex, in bivalves. Most nota-
1354 bly, an IKK-like sequence has been characterized in oyster and connected to the
1355 activation of NF- κ B (Escoubas et al. 1999). As a major difference with vertebrates,
1356 only a single IKK α/β homolog is present in *M. galloprovincialis*. The IKK complex
1357 finally phosphorylates the *Inhibitor of nuclear factor kappa-B* (IK β), which is then
1358 ubiquitinated and targeted for proteasomal degradation. This process allows the
1359 entering of the NF- κ B or Rel transcription factors in the nucleus, ultimately enabling
1360 the transcription of the target effector genes.

1361 After the initial characterization of an IK β homolog in *M. gigas* (Montagnani
1362 et al. 2008), three paralogous genes were identified in this species. All of them were
1363 positively regulated by MAMP and heat-killed bacteria stimulation (Zhang et al.
1364 2011e; Xu et al. 2015a). Similarly, *M. galloprovincialis* possesses at least two IK β
1365 genes, which both experienced moderate to strong upregulation in response to bac-
1366 terial challenges (Toubiana et al. 2014). IK β homologs were also found to be

responsive to various types of challenges in *A. farreri*, *Cyclina sinensis*, *Meretrix meretrix*, *P. fucata*, *R. philippinarum*, *S. glomerata*, and *S. grandis* (Zhang et al. 2009a; Green and Barnes 2009; Wang et al. 2011b; Yang et al. 2011b; Moreira et al. 2012a; Lee et al. 2013; Liu et al. 2014; Gao et al. 2016). In this respect, a contrasting result was obtained in *A. irradians*, as IK β was downregulated following *V. anguillarum* challenges (Mu et al. 2010). The consensus of studies further seems to indicate widespread expression of these inhibitors in all adult tissues, even though most experimental studies have been focused on expression dynamics in hemocytes.

Nuclear Factor Kappa B: A Key Regulator of Immune Response

Nuclear factor kappa B (NF- κ B) family members, sharing a domain architecture similar to human p100/p105 or to p65, have been identified in multiple bivalve species, where they are present as single-copy genes (Li et al. 2015b). The first functional confirmation of the involvement of bivalve NF- κ B homologs in immune response came from the observation that the overexpression of the oyster gene in *Drosophila* cell lines was able to induce the expression of a NF- κ B reporter gene (Montagnani et al. 2004). This molecule could be further placed within the TLR-mediated MyD88-dependent circuitry thanks to RNAi studies in *C. sinensis* (Gao et al. 2016). Furthermore, the *A. farreri* homolog controls the expression of AMPs, providing direct evidence in support of its involvement in the production of effector molecules (Oyanedel et al. 2016). Overall, compelling evidence demonstrates the MyD88-dependent inducibility of NF- κ B in the acute phase of response to various bacterial and viral MAMPs in bivalves, supporting the role of these transcription factors in regulating the expression of proinflammatory factors, effector molecules, and cytokines involved in fundamental aspects of bivalve immunity (Wang et al. 2011b; Huang et al. 2012; Toubiana et al. 2014; Li et al. 2015b; Gao et al. 2016). However, significant differences in the magnitude of this response exist among species which might, to some extent, even explain the different interspecies susceptibility to disease, as evidenced by the comparative analysis of shallow-water and deepsea mussels (Martins et al. 2014).

Other Immune Signaling Pathways

Role of the Mitogen-Activated Protein Kinase Cascade in Immune Signaling

While the processes outlined above cover the main signaling pathway from MAMP sensing to the activation of nuclear factors, some components of the TLR/NF- κ B signaling found in vertebrates and invertebrates alike represent a bridge to other signaling pathways (O'Neill and Bowie 2007; Brown et al. 2011). Most notably, TRAF6 can interact with MEKK1 thanks to mediation by the *Evolutionarily conserved signaling intermediate in Toll pathways* adapter (ECSIT), which is also found in bivalves (Toubiana et al. 2014; Lin et al. 2017), activating the mitogen-activated protein kinase (MAPK) cascade. In essence, the MAPK signaling is a phosphorylation cascade activated by many immune and nonimmune signals (e.g.,

1408 growth factors, cytokines, bacteria, viruses, oxidative stress), which modulates vari-
1409 ous cell processes. This important signaling cascade activates classical MAP kinases
1410 (ERK, p38, JNK), whose concerted action can determine alternative cellular fates,
1411 including cell survival and proliferation, differentiation, or death. The successful
1412 use of commercial antibodies targeting MAPK components evidenced the remark-
1413 able conservation of this pathway in all animals (Canesi et al. 2002; Bettencourt
1414 et al. 2009). Sequences denoting MAPK proteins have been identified in different
1415 mussel and oyster species (Martins et al. 2014; Zou et al. 2015; Gerdol and Venier
1416 2015; Wang et al. 2017a) and p38, JNK, and ERK kinases in particular have been
1417 specifically linked to bivalve immune response (Sun et al. 2016; Qu et al. 2016,
1418 2017a). Ultimately, MAPK signaling results in the activation of AP-1, a heterodi-
1419 meric transcription factor composed of Jun and Fos subunits. The immune role of
1420 bivalve AP-1 has been so far mostly inferred from gene expression data collected in
1421 *C. hongkongensis* and *R. philippinarum* (Xiang et al. 2014a; Wu et al. 2015; Qu
1422 et al. 2015a). Regardless of the alternative activation of the IKK complex or of the
1423 MAPK cascade downstream of MyD88, the two signaling branches extensively
1424 communicate with each other, as TAK1 can phosphorylate (and activate) MAPKs,
1425 and MEKK1 can phosphorylate (and activate) the IKK complex (Moustakas and
1426 Heldin 2003).

1427 **Interferon-Responsive Factors**

1428 Another alternative signaling route potentially activated upon the interaction
1429 between TLRs and intracellular adaptors would lead to the activation of *Interferon-*
1430 *Responsive Factors* (IRFs), a class of transcription factors that enable the expres-
1431 sion of interferons and other proinflammatory cytokines. However, this typical
1432 vertebrate pathway implies the mediation of TRIF (instead of MyD88) and RIP
1433 kinase 1 (instead of IRAKs) which both lack convincing homologs in bivalves
1434 (Meylan et al. 2004). Bivalve IRFs have been linked to resistance to infections in
1435 *H. cumingii* (Wang et al. 2013a) and to the transcriptional activation of genes with
1436 ISRE elements in the pearl oyster *P. fucata* and the mussels *Bathymodiolus plati-*
1437 *frons* and *Modiolus modiolus* (Huang et al. 2013b, 2017a). However, since the exist-
1438 ence of MyD88-independent TLR signaling seems unlikely in bivalves, these
1439 IRF-like molecules are probably related to other signaling routes originated from
1440 cytosolic PRRs, which will be described in detail below.

1441 **Is an Immune Deficiency–Like Pathway Present in Bivalve Molluscs?**

1442 The possible presence of a bivalve immune deficiency (IMD)–like pathway involved
1443 in the recognition of Gram-negative bacteria and homologous to that found in
1444 *Drosophila* (Lemaitre and Hoffmann 2007) has been long hypothesized. In this
1445 case, the immune signals would originate from membrane-bound PGRPs and be
1446 transduced in the cytosol by signaling molecules that are partially shared with the
1447 vertebrate tumor necrosis factor receptor (TNFR) signaling pathway. These include
1448 dFADD and DREDD/Caspase-8, which are both present in bivalves (Gerdol and
1449 Venier 2015), but also the IKK complex and MAPK pathway, which can be acti-
1450 vated by the cross talk between TNFR and TLR signaling. Crucially, however, the

key IMD adaptor molecule is lacking and no functionally homologous component has been identified yet in bivalves (Gerdol and Venier 2015). Taking into account the relevant sequence divergence between the intracellular domain of arthropod and molluscan membrane-bound PGRPs (see section “[Other Membrane-Bound Immune Receptors](#)”), the identity of the hypothetical key mediator of the IMD-like pathway in these animals remains presently unknown.

Signaling Pathways Activated in Response to Microbial Sensing in the Cytosol

The interconnected signaling pathways presented so far act at the crossroads with the cytosolic PRRs described in section “[Cytosolic Pattern Recognition Receptors](#),” which share several signal transducers with the TLR/NF- κ B/MAPK/IRF circuitry, thereby resulting in the activation of the same transcription factors and in the production of similar effector molecules. Among these, the signaling by NLRs would hypothetically involve the mediation of *receptor-interacting serine/threonine protein kinase 2* (RIPK2) for the recruitment of TAK1 and the consequent activation of the IKK complex (Nembrini et al. 2009). However, the lack of a bivalve RIP2K homolog points out that a bivalve NLR-based cytosolic MAMP-sensing system, if it exists, should be based on molecules that are divergent from their vertebrate functional homologs.

In vertebrates, STING stimulates the phosphorylation of IRF3 through the action of the TANK-binding kinase 1 (TBK1) (Tanaka and Chen 2012), the gene of which has been recently characterized in *M. gigas*. The oyster homolog was strongly upregulated in response to *V. alginolyticus* and OsHV-1 infections and, most importantly, its direct interaction with STING was demonstrated by co-IP studies, thereby confirming a mode of signal transduction similar to those in vertebrates (Tang et al. 2016).

RLRs, key sensors of viral nucleic acids (see section “[Cytosolic Pattern Recognition Receptors](#)”), require the *IFN-beta promoter stimulator* (IPS-1, also known as *CARD adaptor inducing IFN-beta*, or CARDIF, and *Virus induced signaling adaptor*, or VISA) to induce the expression of interferon and inflammatory cytokines via IRFs or NF- κ B (Fredericksen et al. 2008). This adapter has remained elusive for a long time in invertebrates, until the very recent discovery of the *M. gigas* homolog CgMAVS. The functional characterization of the oyster protein confirmed its primary role in antiviral response, as (1) CgMAVS could be strongly upregulated in response to viral infections; (2) the interaction between the CARD domain of CgRIG-I-1 and CgMAVS was demonstrated by yeast two-hybrid and co-IP; (3) an interaction was similarly demonstrated with the downstream signaling adapter TRAF6; and (4) the inactivation of CgMAVS by RNAi in infected oyster spat determined a remarkable increase in mortality (Huang et al. 2017b). The demonstrated interaction with TRAF6 would imply the activation of NF- κ B. However, the most important MAVS interactor in vertebrates is another member of the TRAF family, TRAF3, which can recruit TBK1, activating IRF3. The first molluscan TRAF3 homolog was recently identified in the freshwater mussel *Anodonta woodiana*. Although the physical interaction with MAVS and RLRs has not been

1495 demonstrated yet, bacterial and viral challenges triggered the overexpression of this
1496 molecule, supporting its involvement in RLR-mediated signaling (Qu et al. 2017c).

1497 Altogether, these functional studies, supported by the identification of nearly all
1498 of the required signaling molecules in sequence databases (Philipp et al. 2012;
1499 Green et al. 2015; Ren et al. 2017b), as well as by the observation of their significant
1500 upregulation in response to experimental OsHV-1 infection in oysters (He et al.
1501 2015), highlight that bivalve molluscs are equipped with a well-developed molecular
1502 system for viral sensing in the cytosol.

1503 **Production of Cytokines**

1504 **Elusive Regulators of the Molluscan Immune System**

1505 The complex signaling machinery described in detail in the previous sections ulti-
1506 mately leads to the production of effector molecules that are used to kill or to reduce
1507 the pathogenicity of invading microbes (see section “[Humoral Immune Effectors](#)”)
1508 or to regulate immune response at a cellular level (see section “[Cellular Immune](#)
1509 [Responses](#)”) and at a systemic level. Cytokines are small glycoproteins with regula-
1510 tory immune functions, which are the most important regulators of metazoan immu-
1511 nity, as they activate signaling elements leading to the expression of other cytokines,
1512 antiviral effectors, and other immune-related genes. Their action is very fast and
1513 powerful in the amplification of the immune response despite an extremely low
1514 concentration in body fluids. Furthermore, many cytokines have a pleiotropic effect
1515 and a somewhat redundant function (Nicola 1994). Despite the essential and long-
1516 known role of cytokines in vertebrates, their existence in invertebrate animals was
1517 long debated until the first molecules with a cytokine-like activity were first identi-
1518 fied (Beschlin et al. 2001; Herpin et al. 2004). Moreover, as explained in section
1519 “[Other Membrane-Bound Immune Receptors](#),” one of the most studied cytokines in
1520 the *D. melanogaster* model, Spätzle (Parker et al. 2001), is not present in bivalves
1521 and therefore TLRs are likely to act in a vertebrate-like fashion, by directly binding
1522 MAMPs with their extracellular LRR domains. Despite the availability of genomic
1523 sequence data, interferon-like factors remain elusive in all invertebrates, seemingly
1524 supporting the idea that vertebrate and invertebrate cytokines have a different evo-
1525 lutionary origin, despite sharing a similar mode of action and a quite conserved
1526 intracellular signaling machinery. For the most part, molecular studies on molluscan
1527 cytokines are limited to evolutionarily conserved factors, readily identifiable by
1528 sequence similarity.

1529 **Structurally Conserved Cytokines: Interleukin-17, Macrophage** 1530 **Migration Inhibitory Factor, and Allograft Inflammatory Factor-1**

1531 The first bivalve cytokine to be identified was interleukin-17, produced at signifi-
1532 cant levels in oyster hemocytes in response to bacterial exposure (Roberts et al.
1533 2008). IL-17 sequences have been subsequently isolated in many bivalve species or
1534 detected as highly responsive transcripts to bacterial challenges and abiotic stimuli

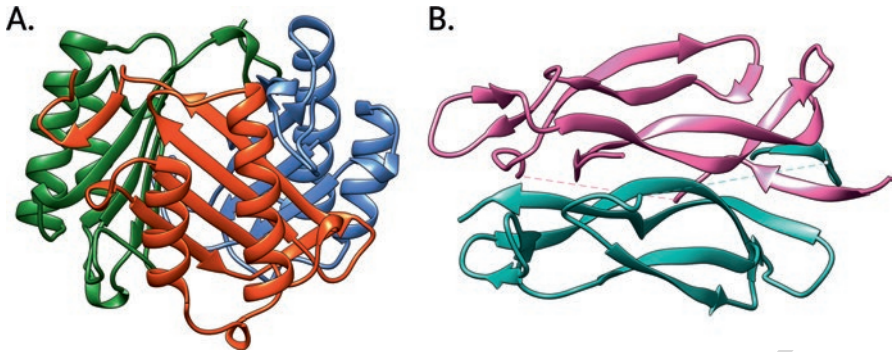


Fig. 12 Structure of two of the evolutionarily conserved cytokines found in bivalve molluscs. (a) Human macrophage migration inhibitory factor (MIF) trimer (PDB accession ID: 1MIF). (b) Human interleukin-17 dimer (PDB accession ID: 4HR9)

(Wu et al. 2013; Moreira et al. 2014; Xin et al. 2015, 2016b). Genomic studies have further revealed that oyster IL-17 proteins are the product of a multigenic family, which comprises at least five members (Li et al. 2014). Although IL17 signaling requires further study in bivalves, homology-based inference suggests that because of its conserved structure (Fig. 12), the binding of IL17 to its receptor stimulates downstream CIKS/CIKSL proteins via SEFIR–SEFIR domain interactions and, subsequently, TRAF proteins related to both MAPK and NF- κ B signaling (Rosani et al. 2015).

The macrophage migration inhibitory factor (MIF) and the allograft inflammatory factor-1 (AIF-1) are two other proinflammatory cytokines that have been identified in bivalves by sequence similarity. The former is a CD74 ligand, which stimulates the acute phase response. Despite the clear difference between bivalve and vertebrate circulating immune cells, the *M. galloprovincialis* MIF displays a well conserved three-dimensional fold (Parisi et al. 2012) (Fig. 12). In contrast with expression data collected in mussels, the *A. farreri* MIF sequence was upregulated upon bacterial challenges in a study that also provided an important confirmation about the functional conservation this molecule, as the recombinant protein could induce fibroblast migration (Li et al. 2011b). In addition, single nucleotide polymorphisms of MIF have been connected with increased resistance to *Vibrio* spp. infections in *M. meretrix* (Zou and Liu 2016). AIF-1, on the other hand, is activated in macrophages upon tissue injury. In *O. edulis*, AIF-1 was upregulated in the hemocytes and mantle of oysters affected with heavy bonamiosis (Martín-Gómez et al. 2014), and its expression could be induced in *M. gigas* with multiple immune challenges (Zhang et al. 2013b). From a functional point of view, the similarity between vertebrate and bivalve AIF-1 proteins is remarkable. Indeed, the oyster homologs could stimulate phagocytosis in the granulocyte hemocyte subpopulation and a clear involvement in tissue damage could be also established (Li et al. 2013a).

1563 **Tumor Necrosis Factor- α : A Cytokine Acting at the Crossroads** 1564 **Between Immunity and Apoptosis**

1565 Following the identification of a *tumor necrosis factor* α (TNF- α) in disk abalone
1566 (De Zoysa et al. 2009), this multifunctional immune modulator was also described
1567 in *M. gigas*, *C. hongkongensis*, and *O. edulis* (Martín-Gómez et al. 2014; Sun et al.
1568 2014; Qu et al. 2017b). Oyster TNF- α transcripts are upregulated in response to
1569 immune challenges and bonamiosis and modulate phagocytosis and apoptosis in
1570 hemocytes. Furthermore, TNF- α recombinant proteins could induce the expression
1571 of NF- κ B reporter genes in human cell lines. In bivalve molluscs, the conserved
1572 function of this cytokine, which acts at the crossroads between the immune system
1573 and the apoptotic machinery, is supported by the identification of conserved acces-
1574 sory factors, i.e., TTRAP (Yang et al. 2011a) and *lipopolysaccharide-induced TNF*
1575 *factor* (LITAF), a positive regulator of TNF- α transcription (Zhu and Wu 2012;
1576 Yang et al. 2013a). As mentioned in section “Other Immune Signaling Pathways,”
1577 TNF- α would exert its function through a signaling pathway partially shared with
1578 the arthropod IMD pathway, which includes the key evolutionarily conserved com-
1579 ponents dFADD and DREDD (Gerdol and Venier 2015). The transduction of
1580 immune signal inside the cell is enabled by the binding of TNF-like molecules to
1581 their receptors, collectively known as TNFRs. Functional tests carried out in many
1582 bivalve species support the involvement of bivalve TNFRs in the establishment
1583 of immune response, despite their limited homology with vertebrate receptors (Li
1584 et al. 2009; Su et al. 2011; Xing et al. 2016; Xiang et al. 2016). Another cytokine
1585 involved in the regulation of cell death, the *TNF-related apoptosis-inducing ligand*
1586 (TRAIL), is ubiquitously expressed in various tissues in *H. cumingii* and *Magallana*
1587 *ariakensis*. The few experimental pieces of evidence collected so far point toward
1588 the involvement of the MAPK pathway in the activation of this cytokine and also
1589 suggest the involvement of caspase 3 as a downstream effector (Yang and Wu 2010;
1590 Yang et al. 2013b).

1591 **New Opportunities for Cytokine Studies in Bivalves**

1592 Many divergent molecules with a cytokine-like function in bivalve molluscs have
1593 only been recently identified or still remain to be uncovered. An important example
1594 is provided by myticin C, a long-known mussel antimicrobial peptide, which has
1595 also been shown to bear chemotactic properties, stimulating hemocyte migration
1596 and morphological changes (Balseiro et al. 2011). The discovery of a class II helical
1597 cytokine in *M. gigas* with remote homology with vertebrate IFN-like molecules
1598 further stimulates research efforts directed at the discovery of novel cytokines in
1599 bivalves. CgIFNLP was upregulated in response to poly(I:C) stimulation and the
1600 recombinant protein could sensibly enhance both apoptosis and phagocytosis in
1601 oyster hemocytes (Zhang et al. 2015).

1602 In the vertebrate IFN signaling, the activation of IFN receptors stimulates the
1603 activity of downstream Janus kinases (JAK) and, consequently, the migration of the
1604 Signal transducer and activator of transcription (STAT) to the nucleus, with the con-
1605 sequent expression of *IFN-stimulated genes* (ISGs). This signaling pathway, whose
1606 presence in bivalves had been already assessed by a number of transcriptomic studies

(Philipp et al. 2012; Green et al. 2014, 2015), has been conclusively implicated in the regulation of immune response by CgIFNLP through its newly isolated receptor (Zhang et al. 2016b). 1607
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Connections with the Neuroendocrine System 1610

The neuroendocrine immunomodulation (NEI) regulatory network encompasses 1611
the complex cross talk between the nervous system, the endocrine system, and the 1612
immune system to maintain homeostasis and to modulate innate immune response 1613
in all animals (Fig. 13). Although NEI appears to be simpler in invertebrates than in 1614
vertebrates, it is highly conserved and represents an efficient regulatory mechanism 1615
(Hartenstein 2006). From this point of view, molluscs are of particular interest, as 1616
they are the most primitive animals with a complete NEI system there is evi- 1617
dence that points to hemocytes as a connecting link between the immune and the 1618
nervous system (Liu et al. 2017b). While cephalopods have long been considered as 1619
privileged molluscan models for the study of NEI because of their well-developed 1620
nervous system and amenability for laboratory research (Di Cosmo and Polese 1621
2016), in recent years bivalve molluscs have been the subject of an increasing num- 1622
ber of studies (Song et al. 2015; Wang et al. 2017a). The main components of the 1623
NEI are the cholinergic, catecholaminergic, and nitric oxidase systems, together 1624
with the action of the neuropeptides. 1625

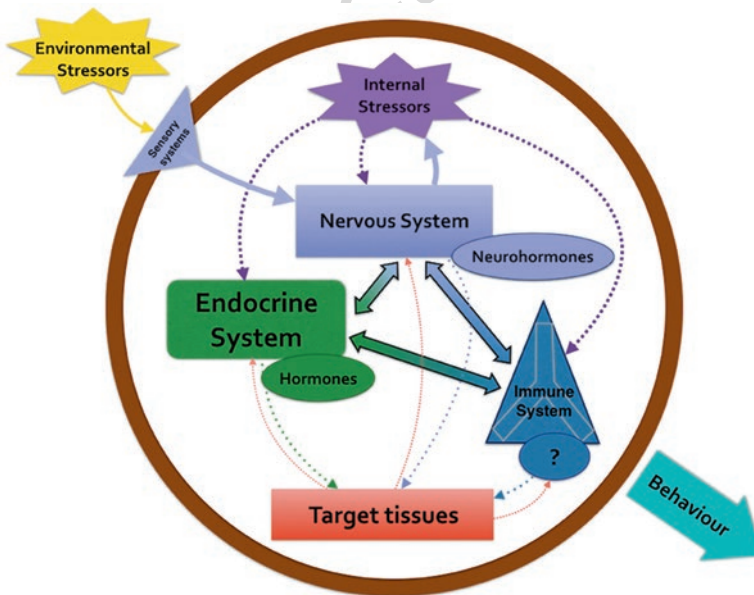


Fig. 13 Cross talk between the nervous, endocrine, and immune systems in response to an external stimulus. (Original Source: Di Prisco and Polese 2015)

1626 **The Cholinergic and Catecholaminergic Neuroendocrine Systems**

1627 The cholinergic neuroendocrine system can be activated by pathogens and tends to
1628 negatively regulate the immune response on a long time scale. The main component
1629 of the cholinergic nervous system is acetylcholine (ACh), whose concentration has
1630 been shown to significantly increase in the hemolymph of scallops upon stimulation
1631 with LPS or TNF- α (Shi et al. 2014). Acetylcholinesterase-like enzymes and mus-
1632 carinic receptors of ACh have been detected in the hemocytes and other tissues of
1633 bivalve molluscs. Strikingly, the *A. farreri* acetylcholinesterase is thought to con-
1634 tribute to the rebalancing of the immune system following immune response in
1635 *A. farreri* (Shi et al. 2012). As a further confirmation in support of the existence of
1636 the cholinergic anti-inflammatory pathway in this animal group, the expression of a
1637 novel muscarinic acetylcholine receptor was regulated by LPS stimulation in
1638 *M. gigas*. The activation of this receptor seems to be crucial for the production of
1639 TNF and for the regulation of apoptosis in hemocytes (Liu et al. 2016b). Moreover,
1640 the subunits of the nicotinic acetylcholine receptor of *A. farreri* were subjected to a
1641 similar induction in response to LPS and TNF- α (Shi et al. 2015).

1642 The catecholaminergic neuroendocrine system is mainly composed of catechol-
1643 amines (dopamine, norepinephrine and epinephrine), their metabolic enzymes, and
1644 receptors. Catecholamines are among the first neurotransmitters to appear during the
1645 ontogenesis of molluscs to regulate cell proliferation, differentiation, and neurogen-
1646 esis. In adults, the synthesis and release of catecholamines has been reported in the
1647 hemocytes, mantle, and gills. The first important evidence supporting the involve-
1648 ment of this system in the modulation of both the cellular and humoral immune
1649 response was provided by the observation of the induction of the alpha-1 norepineph-
1650 rine receptor in response to LPS in *M. gigas*. This receptor could in turn modulate the
1651 expression of TNF and induce phagocytosis and apoptosis of hemocytes (Liu et al.
1652 2016c). Furthermore, the catecholaminergic system is markedly activated after acute
1653 heat and bacterial stress in oyster larvae (Liu et al. 2017a).

1654 **Nitric Oxide, Neuropeptides, and Open Challenges** 1655 **in Neuroendocrine Immunomodulation Studies**

1656 NO synthase (NOS) is a fundamental enzyme for the production of nitric oxide
1657 (NO), a key signaling molecule involved in multiple processes, including immune
1658 defense. Unlike vertebrates, molluscs display only a single NOS isoform, point-
1659 ing toward the existence of a unique prototypical enzyme that combines the func-
1660 tions of the three vertebrate isoforms. Recently, the mutual modulation between
1661 norepinephrine and nitric oxide during immune response has been demonstrated
1662 in scallops (Jiang et al. 2014), showing the intimate linkage among all of these
1663 regulatory systems.

1664 Neuropeptides include a diverse class of cell signaling molecules. These mole-
1665 cules are produced and released by neurons, and their mechanism of action occurs
1666 through a regulated secretory pathway. As in vertebrates, various neuropeptides
1667 identified in molluscs could potentially play important roles in immune regulation.
1668 Although 74 possible neuropeptide genes have indeed been identified in the oyster

genome (Zhang et al. 2012a), neuropeptide studies in the context of immunity are still lacking in bivalves. 1669
1670

As a final consideration about the regulation of NEI function in molluscs, the action of microRNAs also needs to be taken into account. In fact, several miRNAs (named NeurimmiRs) are highly responsive to acetylcholine and norepinephrine stimulation in oyster hemocytes. The in silico-predicted targets for NeurimmiRs comprise over 300 genes with functions in cell death, immunity, and response to stimulus, which might therefore explain the observed decrease in phagocytosis and late apoptosis/necrosis in stimulated hemocytes (Chen et al. 2015). One of the identified miRNAs was subjected to further studies, which evidenced its role in repressing acetylcholine production and choline uptake in hemocytes (Chen et al. 2016). 1671
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Humoral Immune Effectors 1680

Antimicrobial Peptides 1681

Because of their fundamental role as a first line of defense in the molluscan innate immune system and potential biotechnological applications, antimicrobial peptides (AMPs) have been the subject of a considerable number of molecular studies. The first pioneer studies, targeting the hemolymph of mussels, provided the impetus for the characterization of novel antimicrobial compounds, using classical biochemical methods. This field of research is growing thanks to the application of in silico data-mining approaches, and bivalves have been one of the most extensively exploited sources of AMPs in the animal kingdom over the past 20 years. 1682
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Defensins, Mytilins, and Myticins: Main Players in Hemocyte-Mediated Immune Response 1690

The story of antimicrobial research in bivalve molluscs dates back to 1996, when several novel cysteine-rich peptides similar to arthropod defensins were extracted from the active fraction of hemolymph in the marine mussels *M. edulis* and *M. galloprovincialis* (Hubert et al. 1996; Charlet et al. 1996). Two novel peptides, containing eight cysteine residues arranged in a slightly different pattern, were named mytilins and displayed significant activity mostly directed against Gram-positive bacteria (Charlet et al. 1996). Mytilins and defensins exert their antimicrobial action following the recruitment of a specialized subpopulation of circulating hemocytes to the site of infection, where they are intracellularly released from granules (Mitta et al. 2000b, c). Although these AMPs are clearly involved in the intracellular killing of bacterial cells phagocytosed by hemocytes, they also appear to secondarily participate in the systemic immune response when released in the hemolymph (Mitta et al. 2000d). A few years later, a new class of AMPs named myticins was identified in *M. galloprovincialis* plasma and hemocytes. These peptides displayed only limited antimicrobial properties in comparison with defensins and mytilins but shared eight conserved cysteine residues and high hemocyte 1691
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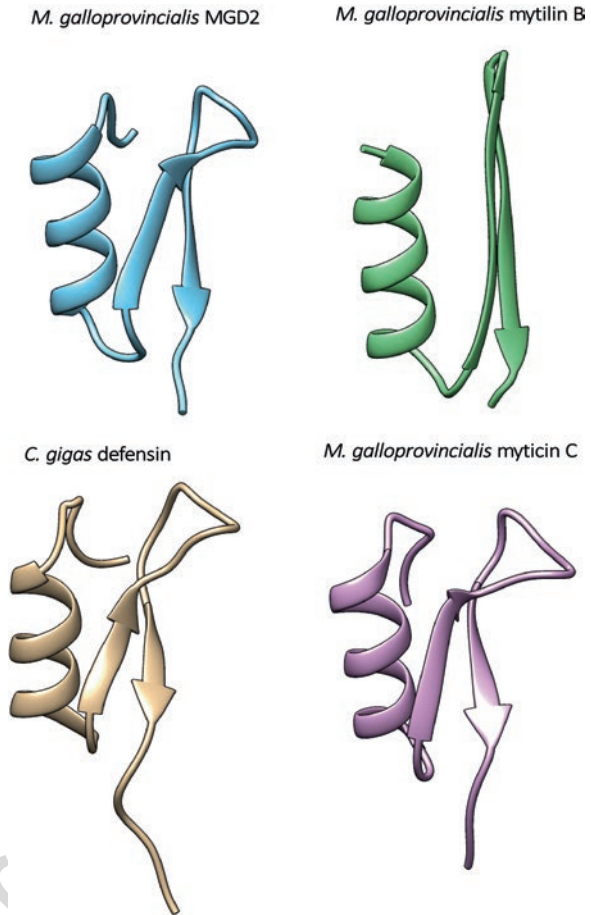
1708 specificity (Mitta et al. 1999). Although the antimicrobial activity of myticins is
1709 rather weak and it can be only attained at acidic pH (Martinez-Lopez et al. 2013;
1710 Domeneghetti et al. 2015), they might have alternative potential roles both as anti-
1711 viral agents and as chemokine/cytokine-like molecules (Balseiro et al. 2011;
1712 Novoa et al. 2016).

1713 Molecular and genetic studies revealed that these mussel AMPs are produced as
1714 secreted pre-propeptides. The highly cationic charge of the central mature peptide
1715 region is balanced by an acidic C-terminal extension of the precursor protein,
1716 which is likely removed after its release from hemocyte granules. It was also
1717 revealed that these AMPs pertain to multigenic families that share a similar archi-
1718 tecture, as they all comprise four exons and three introns, with fixed exon/intron
1719 boundaries (Mitta et al. 2000a). An aspect of mussel hemocyte-specific AMPs that
1720 has revealed somewhat counterintuitive patterns concerns unpredictable fluctua-
1721 tions in gene expression in response to bacterial challenges (Mitta et al. 2000a) and
1722 significant intraspecific variation, suggesting that genome–environment interactions
1723 play a major role in regulating AMP production (Li et al. 2010).

1724 A few years after the original discovery of AMPs in mussel hemocytes, defensin-
1725 like AMPs with eight cysteines were also identified in circulating immune cells in the
1726 Pacific oyster, together with a second isoform mainly expressed in the mantle edge
1727 (Gueguen et al. 2006; Gonzalez et al. 2007a). Over the years, many other sequences
1728 labeled as “defensin” or “defensin-like” AMPs have been isolated in different bivalve
1729 species. Besides their structural differences, summarized by the presence of either
1730 three or four disulfide bonds, these AMPs are also often characterized by different
1731 spectra of activity, preferential tissues of expression, and accessory functions.
1732 For example, a foot-specific defensin-like peptide has been linked to byssogen-
1733 esis in zebra mussels (Xu and Faisal 2010), whereas a gill-specific peptide with
1734 marked activity against Gram-positive bacteria has been isolated from gills extracts
1735 of *C. virginica* (Seo et al. 2005). Clam and freshwater mussel defensins display a
1736 spectrum of activity and tissue specificity similar to those of *Mytilus* AMPs, but they
1737 are reportedly upregulated following bacterial challenges (Peng et al. 2012; Wang
1738 et al. 2015c). These reports suggest that different cysteine-rich peptides currently
1739 classified with the same label could have slightly different biological properties
1740 depending on the species of origin.

1741 From a structural point of view, all of the aforementioned defensin-like AMPs
1742 (including mytilins and myticins) share a common structural motif, the cysteine-
1743 stabilized α -helix β -sheet (CS- $\alpha\beta$) fold (Fig. 14). This conserved and successful
1744 compact domain consists of an α -helix and two antiparallel β -sheets, whose orienta-
1745 tion and reciprocal position in the 3D space are fixed by intramolecular disulfide
1746 bridges (Yang et al. 2000; Gueguen et al. 2006). Crystallographic studies revealed
1747 that, in spite of a negligible primary sequence homology and a slightly different
1748 position of cysteine residues, defensins and mytilins share not only the same struc-
1749 tural fold but also similar hydrophobic and hydrophilic areas (Roch et al. 2008).
1750 Although the 3D structure of myticins has not been experimentally determined yet,
1751 modeling approaches have unequivocally evidenced that they are also likely to
1752 adopt a CS- $\alpha\beta$ fold (Domeneghetti et al. 2015).

Fig. 14 Experimentally determined three-dimensional structures of *M. galloprovincialis* MGD2 defensin, mytilin B, and *M. gigas* defensin. The in silico-predicted structure of *M. galloprovincialis* myticin C is also reported. The conserved cysteine-stabilized α -helix β -sheet fold, comprising an α -helix followed by two antiparallel β -sheets, is easily detectable



Other Cysteine-Rich Antimicrobial Peptide Families

In recent years, data-mining approaches have led to the identification of macins, an additional group of bivalve AMPs in the CS- $\alpha\beta$ peptide superfamily. Originally identified in other metazoan phyla, macins were first described as a multigenic family in *M. galloprovincialis* (Gerdol et al. 2012) and later reported in other bivalve species. Although the functional significance of the complex cysteine array of macins is still poorly understood, these peptides are of great interest because of their role in wound healing, in addition to bacterial killing, and their widespread expression across all main tissues.

In comparison with canonical defensins, big defensins pertain to a structurally different but evolutionarily widespread class, also comprising vertebrate β -defensins. The characterizing six-cysteine array of big defensins is located in the C-terminal domain of these AMPs, and it is coupled with an N-terminal α -helical domain whose presence is also required for antimicrobial action. Big defensins have been isolated in many different bivalve species and, while all studies have evidenced the

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1768 inducible expression of these AMPs, contrasting reports have been produced
1769 concerning the main tissues of expression (Zhao et al. 2010; Rosa et al. 2011;
1770 Gerdol et al. 2012; Li et al. 2012; Wang et al. 2014a; Yang et al. 2016). A more
1771 precise indication concerning the localization of big defensins has been provided by
1772 immunofluorescence studies carried out in *A. irradians*, which have evidenced a
1773 prominent abundance in the gill and mantle epithelia, strongly implicating a role in
1774 mucosal immunity (González et al. 2017).

1775 The remarkable diversity of bivalve cysteine-rich AMPs is not limited to pep-
1776 tides with a known structure but also involves novel cysteine arrays and unknown
1777 disulfide connectivities. The first example is that of mytimycin, an antifungal pep-
1778 tide identified in mussel hemolymph extracts (Charlet et al. 1996). Like the other
1779 AMPs stored in granules, this peptide is produced as an inactive precursor, whose
1780 C-terminal extension contains an EF-hand domain. The mature peptide region can
1781 vary in terms of both the number and the arrangement of cysteine residues (Sonthi
1782 et al. 2011). More recently, three additional plausible AMP families—myticusins
1783 (Liao et al. 2013), mytichitins (Qin et al. 2014), and CRP-I (Gerdol et al. 2015a)—
1784 have been identified in *Mytilus* spp. but promising preliminary results still await a
1785 detailed functional characterization.

1786 **Improved Strategies Are Required to Discover Novel Antimicrobial** 1787 **Peptide Families**

1788 Although different molecules with heterogeneous evolutionary origins, amino acid
1789 compositions, and three-dimensional structures can act as antimicrobial agents, nearly
1790 all known bivalve AMPs pertain to a single large category, i.e., AMPs rich in cysteine
1791 residues engaged in disulfide bonds. This reflects the overwhelming prevalence of
1792 the scientific literature on the subject, as very scant information is available about
1793 AMPs devoid of disulfide bonds in Bivalvia. As a striking example, no AMP with an
1794 amphipathic α -helical secondary structure has ever been isolated, despite their wide-
1795 spread distribution and the important role these peptides cover in the innate immune
1796 system of other protostomes (Giangaspero et al. 2001). While it is possible that this
1797 lack of information mirrors a major shift toward the use of cysteine-rich AMPs in
1798 molluscs compared with other metazoans, other explanations are possible. For exam-
1799 ple, *in silico* similarity-based discovery methods are biased toward conserved disul-
1800 fide arrays, whereas α -helical or linear AMPs do not necessarily present a primary
1801 sequence similarity significant enough to allow BLAST- or profile-based detection.

1802 Some evidence supporting the involvement of peptides enriched in particular
1803 amino acids in bivalve immune response first surfaced with the report of short,
1804 secreted proline-rich peptides (CgPrp), which were found to be coexpressed with
1805 defensins in circulating hemocytes in *M. gigas*, synergistically enhancing their
1806 activity (Gueguen et al. 2009). A second, unrelated AMP was constitutively
1807 expressed in multiple tissues of the same species, and it was named molluscidin.
1808 This cationic peptide, similar to an AMP isolated in abalones, contained a series of
1809 dibasic repeats and exhibited broad-spectrum antimicrobial activity (Seo et al.
1810 2013). The third and most recent case of linear cationic AMPs comprises myticalins
1811 and modiolalins from marine mussels pertaining to the *Mytilus* spp. and *Modiolus*

spp. genera, respectively. These AMPs, identified thanks to an *in silico* approach, display a broad spectrum of activity against Gram-positive and Gram-negative bacteria. Myticalins are produced as pre-propeptides and display a gill-specific pattern of expression, suggesting a possible function as modulators of the microbial communities associated with this important filtrating tissue (Leoni et al. 2017).

The last major category of AMPs comprises peptides generated by fragmentation of larger precursors with various nonantimicrobial functions. Two important examples are provided by an antibacterial peptide isolated from *Anadara kagoshimensis*, which is a fragment of hemoglobin I (Chen et al. 2017b) and by the N-terminal highly cationic fragment of the histone H2B (named molluscin), which appears to modulate the bacterial community in the gills of oysters and possibly other bivalves (Seo et al. 2011). Histone H4 may also have a role in bivalve immunity (Nikapitiya et al. 2013).

Sequence Hyperdiversity as an Effective Weapon to Fight Microbial Infection

In addition to interspecies variability, several bivalve AMPs are characterized by an unusually high degree of intraspecific diversity. For example, the diversity of myticin C was first observed by denaturing gradient gel electrophoresis (DGGE), because of the presence of unique characteristic band patterns in individual mussels (Costa et al. 2009a). It was later found out that this variability also matched nucleotide variation at the mRNA level and that about 8% of the codons within the myticin C sequence evolved under strong positive selection (Pallavicini et al. 2008; Padhi and Verghese 2008). This high level of polymorphisms has been also observed in other (but curiously not in all) mussel AMPs with targeted massive parallel sequencing (Rosani et al. 2011). Similar considerations are also valid for oyster and clams defensins, whose sequence variability can be linked to relevant directional selection pressures (Schmitt et al. 2010; Wang et al. 2015c). It is still not entirely clear whether this remarkable sequence diversity is due to a high number of paralogous genes, high allelic variability, RNA editing, or all of these factors combined. Furthermore, evidence collected from both oysters (Rosa et al. 2015) and mussels (Leoni et al. 2017) strongly hints that complex phenomena of gene presence/absence variability might partially explain the extreme diversification of antimicrobial effectors. Certainly, the presence of such a diversified arsenal of AMPs, apparently driven by selective forces, suggests that amino acid variations might have been evolutionarily exploited to broaden the spectrum of action of these molecules, endowing bivalve populations with effective weapons to face the challenge of microbial infection.

Lysozymes, Bactericidal/Permeability-Increasing Proteins, and Other Pore-Forming Molecules

Lysozymes

The term “lysozymes” is used to collectively describe a group of heterogeneous and widespread proteins involved in the animal innate immune system, which display

1853 strong lytic action against bacteria. Although all lysozymes share a similar structural
1854 fold, they largely diverge in their primary sequence, which can therefore be used for
1855 classification purposes within three main classes: chicken-type (C-type), goose-
1856 type (G-type), and invertebrate-type (I-type) lysozymes (Callewaert and Michiels
1857 2010). From a genomic perspective, it is now clear that genes encoding all three
1858 major lysozyme types can be simultaneously present in the same species, some-
1859 times with several different variants, which might cover slightly different biologi-
1860 cal functions (Gerdol and Venier 2015). In spite of their remarkable primary
1861 sequence divergence, all lysozymes share the same glycoside hydrolase enzymatic
1862 activity, which catalyzes the hydrolysis of peptidoglycan and, to a lesser extent,
1863 chitin. As PGN is a main component of the bacterial cell wall in Gram-positive
1864 bacteria but not in Gram-negative bacteria, lysozymes display stronger activity
1865 against the former.

1866 The first studies on bivalve lysozymes were conducted on I-type sequences, with
1867 the purification of chlamysin in the Arctic scallop, *Chlamys islandica* (Nilsen et al.
1868 1999). Highly similar sequences, implicated either in immune response or in diges-
1869 tive processes, were later reported in several other bivalve species (Matsumoto et al.
1870 2006; La Peyre et al. 2010; Yue et al. 2011; Ren et al. 2012). The isolation of the
1871 complete gene sequence of bivalve I-type lysozymes allowed in-depth phylogenetic
1872 analyses, which revealed a remote homology between this class of enzymes and
1873 vertebrate C-type lysozymes, hinting at an evolutionary origin from a common
1874 ancestor (Bachali et al. 2002). The discovery that different I-type paralogous genes
1875 in hydrothermal vent mussels play a crucial role not just in antimicrobial response
1876 but also in the management of symbiotic communities (Detree et al. 2016a) is one
1877 of the most significant recent developments in bivalve lysozyme research.

1878 In comparison, bivalve C-type lysozymes have been the subject of little scientific
1879 attention, with only a handful of studies reported so far. Following its initial identi-
1880 fication in *M. galloprovincialis* (Venier et al. 2009), this enzyme was characterized
1881 as an inducible gene product, capable of targeting a broad range of bacteria (Wang
1882 et al. 2013c).

1883 The presence of G-type lysozymes, previously thought to be taxonomically
1884 restricted to vertebrates, was demonstrated in 2007 in the scallop *A. farreri* (Zhao
1885 et al. 2007). In the following years, G-type lysozymes have been genetically and
1886 partly also functionally characterized in scallops and mussels (He et al. 2012a;
1887 Wang et al. 2013c; Li et al. 2013b), evidencing that paralogous gene copies might
1888 have acquired a specialized function in either digestive or immune functions. As a
1889 unique known case in nature, a chimeric protein combining a C-terminal G-type
1890 lysozyme domain with an N-terminal PGRP domain has been identified in *M. gigas*.
1891 This protein, which might combine bacteria binding and lytic properties, was induc-
1892 ible in hemocytes in response to *Marinococcus halophilus* and *V. tubiashii* exposure
1893 (Itoh and Takahashi 2009).

1894 More recently, a fourth type of lysozyme was identified in veneroid clams. This
1895 novel antibacterial protein surprisingly shared significant similarity with lysozymes
1896 produced by bacteriophages to break the PGN chains of the infected bacterial cell
1897 walls and release mature phages (Ding et al. 2014). An interesting comparative

study shed some light on the origin of this gene, revealing its co-option from viruses by horizontal gene transfer in two major bivalve groups, Heterodonta and Palaeoheterodonta. Following this event, the newly acquired sequences underwent complex genomic rearrangements, which overall contributed to increased antibacterial potential (Ren et al. 2017a).

Bactericidal/Permeability-Increasing Proteins

While lysozymes mainly target Gram-positive bacteria, a similar antibiotic action is exerted toward Gram-negative bacteria by Bactericidal/permeability-increasing proteins (BPIs), strong pore-forming agents found in nearly all metazoans. The specificity of action of BPIs is given by the recognition of LPS. The biological properties of *M. gigas* BPI (reminiscent of its vertebrate homologs) and its pattern of expression (broad distribution in different epithelia) suggested a role as a first line of defense in oyster mucosal immunity (Gonzalez et al. 2007b). Further genetic investigations revealed the presence of a second oyster gene copy, which displayed a slightly different expression pattern and functional specialization (Zhang et al. 2011d). Although the expression of BPIs can be positively regulated by LPS and bacterial challenges in oysters and ark shells (Zhang et al. 2011d; Mao et al. 2013), the molecular networks underlying this mechanism are still unknown. However, they are likely to be dissimilar to those involved in the production of lysozymes, which appear to be mostly downregulated under the same experimental conditions (Li et al. 2008; Ren et al. 2012), with some notable exceptions (He et al. 2012a; Wang et al. 2013c).

Might Pore-Forming Molecules Provide a Connection with the Complement System?

The possible connections with MAMP sensing by secreted and membrane-bound PRRs and maybe even with the primitive bivalve complement system remain to be fully elucidated. Because of the absence of convincing homologs of the molecular components of the terminal lytic pathway of the complement system, other pore-forming molecules are likely to cover a similar function in bivalve molluscs. While both lysozymes and BPIs could be involved, other options remain to be investigated.

A fascinating possibility is provided by different recently described cases. The first one, described so far only in the Mediterranean mussel, involves a protein containing a Membrane Attack Complex/Perforin (MACPF) domain structurally similar to that of C6/C7/C8/C9 proteins (Estévez-Calvar et al. 2011). Despite the negligible primary sequence similarity with these complement components, its upregulation strongly suggested an involvement in innate immune response. This observation gained even more importance with the report of over a dozen different similar gene products in the mussel transcriptome, which in some cases encode proteins where the perforin-like domain is associated with a PGN-binding ApeC domain (Gerdol and Venier 2015). The second class of molecules that might act as functional homologs to the complement terminal pathway are mytilectins (see section “The Role of Lectins in Immune Recognition”). Indeed, some mytilectins display a C-terminal aerolysin-like

1941 pore-forming domain, which could be employed in the lysis of microbial cells
1942 (Gerdol and Venier 2015). While both the Ricin B/aerolysin and ApeC/MACPF
1943 domain combinations could potentially result in highly efficient and concerted rec-
1944 ognition and killing of invading pathogens, further functional assays will clearly be
1945 needed to investigate the possibility that these molecules are involved in pathogen
1946 recognition and clearance in mussels and other bivalve species.

1947 **Proteases and Protease Inhibitors**

1948 **An Overview on the Role of Proteases and Their Inhibitors** 1949 **in the Bivalve Immune System**

1950 Several important immune processes are regulated by the concerted action of protea-
1951 ses and their inhibitors, which might act either on endogenous proteins, by cleav-
1952 ing regulatory subunits and enabling their biological activity of their targets, or on
1953 exogenous proteins produced by invading microbes and parasites, leading to their
1954 inactivation and degradation. Some of the fundamental immune processes described
1955 in other sections, such as the complement system (see section “[Evidence of an](#)
1956 [Ancient Complement System in Bivalves?](#)”), the prophenoloxidase cascade leading
1957 to melanization (see section “[The Phenoloxidase Cascade](#)”), and apoptosis (see
1958 section “[Apoptosis and Autophagy](#)”), are essentially governed by a cascade of pro-
1959 teolytic activations, initially triggered by the recognition of MAMPs by PRRs.
1960 Although the molecular players involved in such cascades have been comprehen-
1961 sively characterized in some invertebrates, such as in the case of melanization in
1962 insects (Tang 2009) or hemocyte clotting in horseshoe crabs (Iwanaga et al. 1998),
1963 the nature of such proteases has not been entirely clarified in bivalve molluscs.

1964 This can be partly explained by the lack of specific studies on the subject, but
1965 also finds a justification in the fact that these molecules pertain to large and multi-
1966 functional families of proteases involved in a multitude of other cellular processes,
1967 often not linked with immune response. As an example, while the core components
1968 of the bivalve complement system, as well as a remarkable number of lectin-like
1969 molecules, have been characterized in bivalves, no MASP-like proteases has been
1970 identified with certainty (see section “[Evidence of an Ancient Complement System](#)
1971 [in Bivalves?](#)”), leaving a huge gap of knowledge about the link between MAMP
1972 recognition in the extracellular environment and the activation of C3, even though
1973 several similar uncharacterized serine proteases are present in bivalve genomes
1974 (Wang et al. 2017b). Similarly, the nature and specificity of action of the bivalve
1975 prophenoloxidase-activating enzymes (see section “[The Phenoloxidase Cascade](#)”)
1976 and the identity of the proteases involved in the process of activation of AMPs (see
1977 section “[Antimicrobial Peptides](#)”) still remain uncertain. Big defensins, CRP-I,
1978 mytimycins, and myticalins, for example, possess a dibasic cleavage site, which
1979 could be potentially cleaved off by proprotein convertases (Gerdol et al. 2012,
1980 2015a; Leoni et al. 2017). However, other mussel AMPs such as defensins, mytilins,
1981 and myticins lack a clear consensus motif for propeptide cleavage and are therefore
1982 expected to be the substrates of other, still unknown, proteases.

Cathepsins

While all of the aforementioned proteases mainly exert their biological action in the extracellular environment, others are typically present in lysosomal compartments, where they aid the phagocytic processing of heterophagic and autophagic material. Among these, cathepsins have been the subject of multiple studies and inked to immune functions in bivalves, consistently with the well-known role these proteases have in the regulation of vertebrate immune and cell death processes (Zavasnik-Bergant and Turk 2006; Repnik et al. 2012). In particular, multiple cathepsins have been characterized in the Chinese razor clam, *S. constricta*, where B-, C-, and L-type cathepsin were upregulated following *V. anguillarum* challenges in the mantle and, in particular, in the digestive gland (Niu et al. 2013a, b, 2014). Similar observations concerning tissue specificity and responsiveness to bacterial challenges have been also collected for a cathepsin L in *Cristaria plicata* (Hu et al. 2014), in contrast with a report from the Sidney rock oyster *S. glomerata*, where cathepsin B and L transcripts were mostly detected in hemocytes (Ertl et al. 2016).

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Serine Protease Inhibitors: The Case of Oyster Perkinsosis

The infection process of many animal pathogens is also aided by a number of proteases, which target and inactivate host defense proteins and sometimes have more profound effects on the modulation of the host immune system (Armstrong 2006; Donnelly et al. 2011). In bivalve molluscs, this system has been best characterized in response to the parasite *P. marinus*, which produces proteases that specifically target defense plasma proteins, thereby impairing the immune response and creating favorable conditions for the establishment of infections by bacterial pathogens (Oliver et al. 1999; Tall et al. 1999). As a consequence, many bivalve species have developed large gene families of protease inhibitors to counteract the action of exogenous proteases produced by protozoans and other parasites (Romestand et al. 2002).

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The serine protease inhibitors of the eastern oyster, *C. virginica* (CvSI) (Xue et al. 2009), pertain to the I84 family of serine protease inhibitors. These molecules have been implicated in resistance to *P. marinus* infections because of their high activity in oysters selected for increased survival in comparison with susceptible specimens (La Peyre et al. 2010) and their ability to inhibit the perkinsin pathogenic protease (Xue et al. 2006). Furthermore, a polymorphism located in the promoter region of the CvSI-1 gene was conclusively linked to its increased transcription and, consequently, to improved resistance to *P. marinus* (He et al. 2012b), and the expression levels of CvSI could also explain the interspecies differences in susceptibility to infection between *C. virginica* and the more resistant oyster species *Crassostrea corteziensis* (Gutiérrez-Rivera et al. 2015). Altogether, I84 serine protease inhibitors are part of a highly expanded and still rapidly evolving molluscan gene family (Xue et al. 2017a).

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Kazal-Type Serine Protease Inhibitors and Tissue Inhibitors of Metalloproteinases

Kazal-type serine protease inhibitors are another large and widespread class of molecules that have been connected to immune functions in marine bivalves. These

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2026 molecules were reportedly upregulated in the hemocytes of the scallop *A. irradians*
2027 following tissue injury and bacterial challenges (Zhu et al. 2006). Another Kazal-
2028 type protease inhibitor from *A. farreri* contained 12 tandemly repeated Kazal-
2029 domains and was upregulated upon *V. anguillarum* challenges (Wang et al. 2008),
2030 and two similar but shorter proteins could be similarly induced in the hepatopan-
2031 creas of *R. philippinarum* and in multiple tissues of the clam *Mesodesma donacium*
2032 under similar experimental conditions (Maldonado-Aguayo et al. 2013; Yu et al.
2033 2017). Like I84 inhibitors, Kazal-type inhibitors are produced by a multigenic fam-
2034 ily, whose members display different substrate specificity and sensitivity to stimula-
2035 tion (Zhang et al. 2014a).

2036 The third large class of immunity-related protease inhibitors that has been stud-
2037 ied in bivalves comprises the tissue inhibitors of metalloproteinases (TIMPs).
2038 Cg-TIMP, first identified in *M. gigas* because of its accumulation in hemocytes fol-
2039 lowing shell injury and bacterial challenges (Montagnani et al. 2001), is activated
2040 through a DAMP-dependent pathway and is possibly regulated by NF- κ B binding
2041 elements located in its promoter (Montagnani et al. 2007). The immune properties
2042 of TIMPs have not been investigated in other bivalve species, with the exception of
2043 the blood cockle *Tegillarca granosa*, where TgTIMP-4 is responsive to LPS, PGN,
2044 and *V. parahaemolyticus* challenges (Wang et al. 2012c).

2045 These and other protease inhibitors might be involved in the management of
2046 microbial infections, as suggested by multiple reports of their upregulation from
2047 transcriptomic studies (Feng et al. 2010; Moreira et al. 2012a; Allam et al. 2014;
2048 Nikapitiya et al. 2014). However, the mode of action of just a few of these mole-
2049 cules has been properly functionally characterized. Therefore, protease inhibitors
2050 remain attractive targets for the study of host–pathogen interactions, in particular in
2051 the context of viral infections.

2052 The Phenoloxidase Cascade

2053 The recognition of MAMPs by PRRs, as well as various types of environmental
2054 stress, can trigger an extracellular proteolytic cascade, which leads to the conver-
2055 sion of prophenoloxidases (ProPO) to their active form, phenoloxidases (PO),
2056 copper-binding metalloproteins that catalyze the oxidation or hydroxylation of phe-
2057 nols. Different enzyme classes (tyrosinases, catecholases, and laccases) with low
2058 substrate specificity and similar activity exist in invertebrates, leading to a certain
2059 confusion in their unambiguous identification by biochemical tests on tissue extracts
2060 (Luna-Acosta et al. 2017). However, the activity of PO leads to the synthesis of the
2061 melanin pigment. This process, unique to a few invertebrate phyla, including arthro-
2062 pods and molluscs, enables the deposition of melanin on invading microbes, limit-
2063 ing the spread of infection. While the molecular players involved in the regulation
2064 of the melanization proteolytic cascade have been extensively studied and charac-
2065 terized in arthropods (Christensen et al. 2005; Tang 2009), limited information is
2066 available in molluscs (Luna-Acosta et al. 2017).

Secreted PGRPs are the main PRRs responsible for the activation of the ProPO cascade in *Drosophila* and other arthropods (Schmidt et al. 2008). However, as explained in section “Other Membrane-Bound Immune Receptors,” while extracellular proteins with an N-acetylmuramoyl-L-alanine amidase domain are encoded by molluscan genomes, they seem to share closer similarities to those of vertebrates, where they play a direct bactericidal role. This divergence is in line with the major differences between arthropods and molluscs, which involve the interconnected TLR (with the lack of Spätzle; see section “Canonical TLR Signaling”) and IMD pathways (see section “Other Immune Signaling Pathways”).

In bivalves, the melanization process has been known for a very long time as a normal physiological process linked to shell deposition in pallial mantle epithelia (Waite and Wilbur 1976). However, increased melanization, usually followed by a massive rearrangement of extracellular matrix deposition and alterations in shell mineralization, is also among the most distinctive features of some common pathologies of the bivalve mantle tissue (see section “Major Infectious Diseases Affecting Bivalve Molluscs”) (Ford and Borrero 2001; Paillard 2004). Further evidence supports the involvement of the ProPO cascade in response to parasitic, bacterial, and viral infection, as PO activity appears to be strongly altered in *M. sydneyi*-infected Sydney rock oysters (Raftos et al. 2014; Luna-Acosta et al. 2017). Melanization is probably not merely an extracellular event, as it might also be implicated in the intracellular killing of encapsulated microbes (Butt and Raftos 2008). Moreover, the different rates of inhibition of PO activity in the hemocytes of *M. gigas* and *Geukensia demissa* in response to *P. marinus* infections could be linked to the different degree of susceptibility of the two species to infection (Jordan and Deaton 2005). These observations support the important role of the ProPO cascade as a system of defense against microbial infections in bivalve molluscs.

The existence of an extracellular ProPO cascade linked to components of the hemolymph has been conclusively demonstrated in *M. gigas* and *Perna viridis*, where it could be induced by LPS, zymosan, and laminarin (Asokan et al. 1997; Hellio et al. 2007). However, a proper functional characterization of POs is still lacking in most bivalve species and the sequences of very few PO genes have been identified. This is ascribable in part to the broad distribution of PO activity in different tissues and life stages, including the digestive gland, the mantle and shell, and the foot, where POs are likely to cover specific functions that are yet to be fully unveiled (Luna-Acosta et al. 2011b). For example, tyrosinases pertain to a gene family which underwent significant expansion in bivalves and has been implicated in the shell mineralization process (Huang et al. 2017c; Chen et al. 2017a). However, a tyrosinase-like protein significantly contributes to PO activity in *S. glomerata* hemocytes (Aladaileh et al. 2007) and a tyrosinase-like transcript whose expression level was significantly overexpressed in response to bacterial challenges has been reported in *A. farreri* (Zhou et al. 2012). In the same species, a 576-kDa protein with PO activity, selectively inhibiting the growth of *Vibrio* spp. and *Aeromonas salmonicida*, has been purified from hemocytes (Xing et al. 2012). Interestingly, a protein with a similar molecular weight (555 kDa), displaying

2111 *p*-diphenoloxidase activity, has been obtained from the hemocytes of a different
2112 scallop species, *A. irradians* (Jiang et al. 2011). Other studies have identified the
2113 hemocyte-specific PO enzyme as a laccase in *M. gigas* (Luna-Acosta et al. 2010,
2114 2011a) and *R. philippinarum*, where only minor tyrosinase-like activity could be
2115 detected (Le Bris et al. 2013).

2116 While the function of the ProPO cascade in the bivalve immune response has
2117 been fully established in relation to different diseases, this topic has been the subject
2118 of limited molecular studies and therefore still awaits detailed investigations to clar-
2119 ify which PRRs enable the melanization of invading microbes, both in the extracel-
2120 lular matrix and within phagocytic cells.

2121 Cellular Immune Responses

2122 Phagocytosis

2123 Hemocytes Are the Main Cell Type Involved in the Phagocytic 2124 Process

2125 Phagocytosis, encapsulation, and cell-mediated cytotoxicity have been extensively
2126 described in bivalves at a functional level and, more recently, at a genomic level
2127 (Schmitt et al. 2012; Soudant et al. 2013; Allam and Raftos 2015; Zannella et al.
2128 2017; Schultz and Adema 2017) (Fig. 15).

2129 During the early 1900s, the pathologist Metchnikoff used marine organisms,
2130 among other models, to describe and hypothesize the role of phagocytosis in diges-
2131 tion, immune defenses, and clearing of damaged cells (Gordon 2016; Schultz and
2132 Adema 2017). A dual role for bivalve hemocytes in digestion and immunity may be
2133 especially important during larval stages in bivalves, as suggested by evidence of
2134 phagocytic activity in early stages of larval development (Song et al. 2016).
2135 Moreover, hemocytes concentrate particulate material in the connective tissues sur-
2136 rounding the digestive glands in bivalve larvae (Dyachuk 2016). A more specific
2137 role for phagocytosis and encapsulation in disease resistance in bivalves has been
2138 hypothesized for Brown Ring Disease in clams, summer mortality in Pacific oys-
2139 ters, and QX disease (*M. sydneyi*) in Sydney rock oysters, based on in vitro observa-
2140 tions of increased phagocytic function and/or upregulation of transcripts for genes
2141 putatively involved in phagocytosis in resistant bivalves compared with susceptible
2142 individuals (Allam and Ford 2006; Samain et al. 2007; Kuchel et al. 2010; Raftos
2143 et al. 2014).

2144 Hemocytes are, by far, the best-studied phagocytic cells in bivalves. Flow cytom-
2145 etry has allowed for the development of high-throughput assays for the evaluation
2146 of hemocyte immune parameters in bivalves, including characterization of the pop-
2147 ulations of cells involved in phagocytosis of inert and biological particles and the
2148 subsequent stimulation of the oxidative burst response. Of the two major types of
2149 hemocytes described in bivalves on the basis of morphology, granulocytes in gen-
2150 eral seemed to be responsible for the majority of the phagocytic response and pro-
2151 duction of radical oxygen/nitrogen species (ROS/RNS), but this is highly dependent

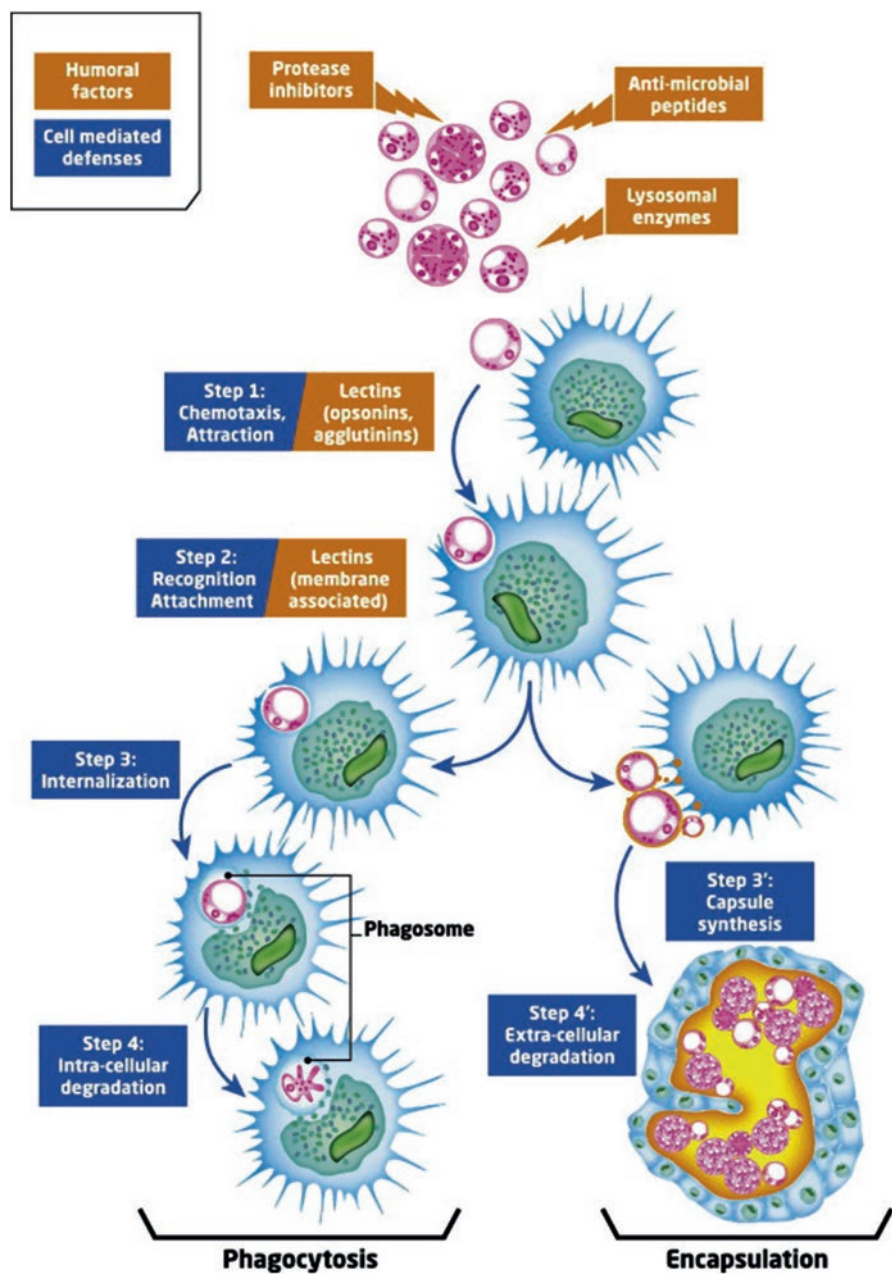


Fig. 15 The main humoral and cellular components of the bivalve immune response to microbial infection. The different steps of phagocytosis and encapsulation are shown in blue. Invading pathogens are indicated in purple, and humoral effectors (see section “Humoral Immune Effectors”) are shown in green. (Source: Soudant et al. 2013)

2152 on the bivalve species and the nature of the stimuli (Schmitt et al. 2012; Soudant
2153 et al. 2013; Allam and Raftos 2015; Zannella et al. 2017; Schultz and Adema 2017).
2154 Moreover, differences in the timing of phagolysosome fusion between eosinophilic
2155 and basophilic hemocytes in deepwater mussels indicate that these two types of
2156 granulocytes may play different roles in phagocytosis, suggesting further definition
2157 of phagocytic capabilities within hemocyte populations (Tame et al. 2015). An addi-
2158 tional type of hemocyte, a hemoblast-like cell, may be involved in phagocytosis,
2159 composing a small percentage of all phagocytic cells in a hemocyte population and
2160 showing low levels of oxidative burst and lysosomal enzyme activity. Differences in
2161 the rates of phagocytosis by hemocytes also depend on the source of hemocytes
2162 within an individual (i.e., circulating hemocytes versus those present in the pallial
2163 or extrapallial spaces). Hemocytes have the ability to migrate through the epithelia
2164 into these cavities and then go back into the tissues, and those collected from the
2165 pallial cavity appear to have higher phagocytic activity than circulating hemocytes
2166 (Allam and Pales Espinosa 2016). These observations indicate that different popula-
2167 tions of hemocytes may respond to selected stimuli and show different mechanisms
2168 of action (Evariste et al. 2016; Bettencourt et al. 2017; Vieira et al. 2017).

2169 Other cells thought to have phagocytic capabilities are epithelial cells, with an
2170 ability that may be exploited by intracellular bacteria such as the Chlamydia- and
2171 Rickettsia-like organisms commonly seen in the gill and mantle epithelia of marine
2172 bivalves and gastropods (Allam and Pales Espinosa 2016). Development of specific
2173 cell markers will help us to understand if differences in phagocytic activity between cell
2174 populations within bivalves are due to the presence of specialized cell populations
2175 and/or the context in which these responses are occurring.

2176 **Phagocytosis in Detail: Chemotaxis, Opsonization, and Endocytosis**

2177 The process of phagocytosis involves the steps of chemotaxis, opsonization, endo-
2178 cytosis, formation of phagosomes, phagosome–lysosome fusion, respiratory burst,
2179 and exocytosis. Upon infection and injury, hemocytes migrate to the site of injury
2180 through the process of chemotaxis. Examples of bivalve pathogens causing massive
2181 focal infiltration of hemocytes at the site of infection include *V. tapetis* (Brown Ring
2182 Disease), *P. marinus*, and QPX. A chemotactic and/or chemokinetic response of
2183 hemocytes has been observed in response to several PAMPs, including bacterial
2184 endotoxins and extracts from trematodes and *P. marinus*. The nature of the chemo-
2185 taxis/chemokinetic response depends on the type of PAMP (Schmitt et al. 2012;
2186 Soudant et al. 2013; Allam and Raftos 2015; Zannella et al. 2017; Schultz and
2187 Adema 2017).

2188 Chemotaxis is followed by opsonization and phagocytosis. Transcriptomic anal-
2189 ysis of Pacific oysters in response to LPS and other immune stimuli indicates that
2190 phagocytosis is promoted by a variety of opsonins (Zhang et al. 2012a). Several
2191 PRRs have been functionally demonstrated to mediate phagocytosis induction by
2192 immune stimuli through several signaling pathways (see sections “[Recognition,](#)
2193 [Agglutination, and Opsonization](#)” and “[Signaling and Regulatory Pathways](#)”).
2194 For example, an extracellular superoxide dismutase (Cg-EcSOD), highly abun-
2195 dant in oyster cell-free hemolymph, induces phagocytosis mediated by a β -integrin

(Duperthuy et al. 2011). Lectins from Manila clams (MCL and MCL4) stimulate the opsonization of *P. olseni* parasite and *V. tubiashii* bacterial cells and subsequent phagocytosis by clam hemocytes in vitro (Soudant et al. 2013; Zannella et al. 2017). Competitive inhibition of a sialic acid-binding immunoglobulin-type lectin (CgSiglec-1) inhibits the stimulation of phagocytosis and apoptosis by LPS in oyster hemocytes, consistent with the role of siglecs as regulators of immune responses (Liu et al. 2016a). Expression of genes involved in signaling pathways associated with integrin signaling and phagocytosis (PI3K, Rho J, MAPPK, PKC), phagosome maturation (Rab32), and respiratory bursts (NADPH oxidase) were upregulated upon secondary exposure to live *V. splendidus* after a primary challenge with killed *V. splendidus* (Zhang et al. 2014d).

Phagocytosis in Detail: Respiratory Burst and Exocytosis

The process of phagosome-lysosome fusion has been functionally observed in deepwater mussels (Tame et al. 2015). After phagosome-lysosome fusion, a respiratory burst ensues, followed by secretion of antimicrobial proteins (see section “Antimicrobial Peptides”) (Soudant et al. 2013). On the basis of studies using enzyme activity measurements and the use of inhibitors, it appears that the mechanisms for production of reactive oxygen species (ROS) and reactive nitrogen species (RNS) are in general homologous to the ones observed in vertebrates (Soudant et al. 2013; Schultz and Adema 2017). The timing and extent of the respiratory burst in bivalve hemocytes, however, differs from those of the respiratory burst in vertebrate models. Moreover, bivalves and other marine invertebrates also show some differences from vertebrates in terms of the basal (not pathogen stimulated) generation of ROS as part of energy metabolism in organelles such as the mitochondria, endoplasmic reticulum, and peroxisomes (Donaghy et al. 2015). Sequencing studies indicate that, in addition to NADPH oxidase, bivalves contain genes similar to dual oxidase (DUOX, involved in immunity in *Drosophila*), which are upregulated in response to pathogenic vibrios. Bivalve hemocytes also show myeloperoxidase (MPO) activity (Schmitt et al. 2012; Donaghy et al. 2015). Radical nitrogen species, such as nitric oxide and peroxynitrite, also have an important role against pathogens in bivalves (Villamil et al. 2007). Nitric oxide also acts as an immune regulator (see section “Connections with the Neuroendocrine System”), enhancing phagocytosis, antibacterial activity, and apoptosis in bivalve hemocytes (Song et al. 2015). Expression of the single nitric oxide synthase (NOS) described in bivalve molluscs is modulated by immune stimuli (Song et al. 2015). In oyster hemocytes stimulated with zymosan, the NOS pathway is more active in hyalinocytes, while NADP oxidase activity is more prevalent in granulocytes (Lambert et al. 2007).

Antioxidant and detoxification enzymes are produced to protect cells from the toxicity of ROS and maintain redox homeostasis. Genome and transcriptome studies have led to the identification of the genes for five superoxide dismutases (SODs) in the Pacific oyster genome (He et al. 2015), two functional catalase genes in the oyster *M. hongkongensis*, and the genes coding for several glutathione peroxidases (GPxs) and glutathione transferases (GSTs) (Sui et al. 2017; Wang et al. 2017a). Of the six known groups of superoxide dismutases, only manganese and copper/zinc

2240 have been characterized so far in bivalves. Little is known, however, about the
2241 specific roles of these enzymes in immunity and disease resistance. An extracellular
2242 SOD from Pacific oysters, CgEcSOD, a major component of oyster plasma, shows
2243 both antioxidant and PRR activities and is able to promote the phagocytosis of
2244 the bacterial pathogen *V. splendidus* (Wang et al. 2017a). The expression of Mn and
2245 Cu/Zn SODs is upregulated with both viral and bacterial challenge, and alleles in
2246 the intracellular and extracellular Cu/Zn SOD have been associated with disease
2247 resistance to *Vibrio* infection in bay scallops (Wang et al. 2013b; Song et al. 2015;
2248 Wu et al. 2017).

2249 **Accessory Factors and Mechanisms of Regulation of Cell-Mediated** 2250 **Cytotoxicity**

2251 Other molecules shown to be involved in intracellular killing in the phagolysosome
2252 in bivalves include hydrolytic enzymes (β -glucuronidase, esterases, phosphatases,
2253 sulfatases, lipases), including unique versions of lysozymes showing tissue-specific
2254 patterns of gene expression (see section “[Lysozymes, BPIs and Other Pore-Forming](#)
2255 [Molecules](#)”) and other antimicrobial molecules (phenoloxidases, antimicrobial
2256 peptides; see section “[Antimicrobial Peptides](#)”) (Tanguy et al. 2013; Zannella
2257 et al. 2017). Phagocytosis and encapsulation are also aided by the prophenoloxidase
2258 system, a complex biochemical cascade occurring mainly in the hemolymph of
2259 bivalves, which is activated by microbial MAMPs, exogenous proteases, and envi-
2260 ronmental stress, leading to the formation of the antimicrobial molecule melanin
2261 (see section “[The Phenoloxidase Cascade](#)”)

2262 Little is known about the process of regulation of cell-mediated cytotoxicity in
2263 bivalves. A potential regulator of hemocyte function, thymosin beta-4, has been
2264 characterized in the oysters *M. hongkongensis* and *M. gigas*, and in the gastropod
2265 *Haliotis discus discus*. Treatment of oysters with recombinant protein led to
2266 increased numbers of circulating hemocytes, increased bacterial clearing, reduction
2267 of ROS production, and increased production of antioxidant enzymes, suggesting a
2268 potential role in wound healing (Li et al. 2016a). Dysregulation of the oxidative
2269 burst, on the other hand, may be involved in the pathogenesis of several diseases
2270 affecting marine bivalves. For example, oxidative stress resulting from a strong oxi-
2271 dative burst response, characterized by a strong upregulation of oxidase genes and
2272 downregulation of antioxidant genes, may contribute to the pathology seen in larval
2273 and juvenile oysters experimentally challenged with OsHV-1 μ Var (He et al. 2015;
2274 Young et al. 2017) or infected with the bacterial pathogen *A. crassostreae* (McDowell
2275 et al. 2014).

2276 **Mechanisms of Evasion Adopted by Invading Pathogens**

2277 Several pathogenic and nonpathogenic vibrios, *Chlamydia* and Rickettsia-like
2278 organisms, and the protozoan parasites *B. ostreae*, *P. marinus*, and *P. olseni* appear
2279 to have evolved mechanisms to evade cell-mediated cytotoxicity in bivalves,
2280 exploiting that ability to survive within host tissues. Potential mechanisms used to
2281 evade phagocytosis and encapsulation include dysregulation of immune signaling
2282 through phosphorylation of p38-MAPK and induction of apoptosis of hemocytes

(Ciacci et al. 2017; Burgos-Aceves and Faggio 2017). Other microbes can avoid intracellular killing by respiratory burst pathways in bivalve molluscs (Schmitt et al. 2012; Soudant et al. 2013; Allam and Raftos 2015). The enzymes arginase, alkaline phosphatase, ascorbate-dependent peroxidase, and superoxide dismutase are several of the factors potentially involved in the ability of *P. marinus* to inhibit ROS production in oyster hemocytes and survive in vitro exposure to ROS (Schott and Vasta 2003; Schott et al. 2003; Fernández-Robledo et al. 2008) (Fig. 16). The parasite is also resistant to high concentrations of nitric oxide (Villamil et al. 2007). The natural resistance-associated macrophage protein (NRAMP) in *P. marinus*, involved in iron uptake in *P. marinus* trophozoites, is hypothesized to deplete iron in hemocytes, limiting the ability of hemocytes to mount an effective respiratory burst (Lin et al. 2011). Moreover, the wall of parasites such as *P. olseni* appears to be resistant to proteolysis (Montes et al. 2002). Extracellular products from a pathogenic strain of *V. splendidus* inhibit phagocytic activity in mussel *M. edulis* hemocytes, while those of a nonpathogenic strain do not (Ben Cheikh et al. 2016). Some metazoan parasites such as the digenean trematodes *Bucephalus* sp. and

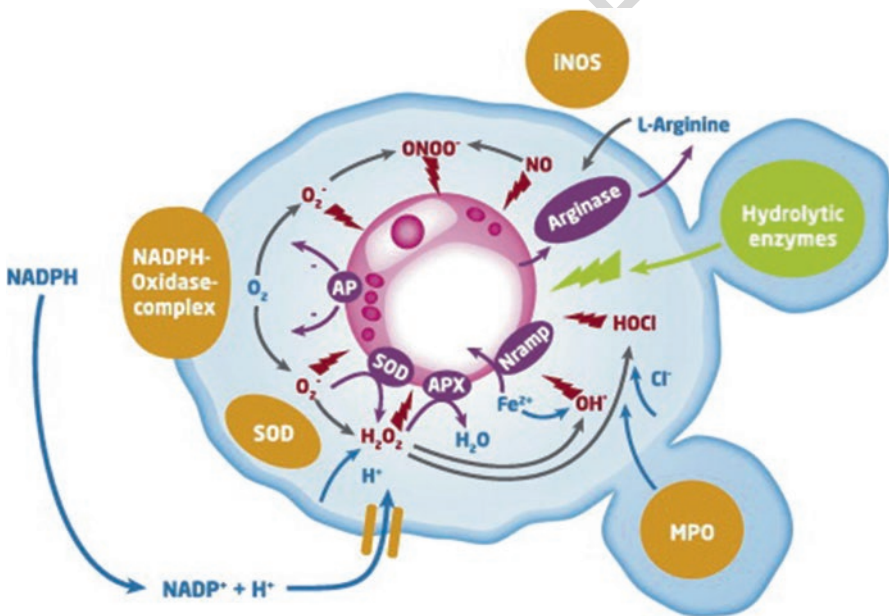


Fig. 16 Interaction between prooxidant (orange) and antioxidant (purple) activities in the phagosome of an hemocyte from the eastern oyster, *Crassostrea virginica* (blue), upon phagocytosis of the protozoan parasite *Perkinsus marinus* cell (purple). Prooxidant activities are exerted by hemocytes to kill the invading microbe by exposure to ROS (red), whereas antioxidant activities are used by *P. marinus* to escape these defensive measures. AP acid phosphatase, APX ascorbate-dependent peroxidase, HOCl hypochloride, iNOS inducible nitric oxide synthase, MPO myeloperoxidase, NO nitric oxide, Nrapm Natural Resistance-Associated Macrophage Protein, O₂⁻ superoxide anion, ONOO⁻ peroxynitrite, SOD superoxide dismutase. (Source: Soudant et al. 2013)

2299 *Proctoeces maculatus* may also modulate hemocyte function in bivalve hosts, leading
2300 to decreased hemocytic infiltration in infected tissues (Carella et al. 2015).

2301 **Encapsulation and Granuloma Formation**

2302 The processes of encapsulation and granuloma formation occur when particles or
2303 pathogens are too large to be engulfed by hemocytes (e.g., in infection by trema-
2304 todes) or the phagocytosis response is unsuccessful (e.g., in infection by *Perkinsus*
2305 spp. or *Nocardia* spp.). In the process of encapsulation, hemocytes recruited to the
2306 site of infection surround and encapsulate the invading pathogen, secreting extra-
2307 cellular matrix products to prevent dissemination of the pathogen to other tissues
2308 and a variety of lysosomal enzymes and antimicrobial molecules to attempt to kill
2309 it (Soudant et al. 2013; Allam and Raftos 2015; Carella et al. 2015). This process
2310 can occur within the tissues, leading to granuloma-like formation, or within the
2311 extrapallial space between the mantle and the inner side of the bivalve shell, leading
2312 to conchiolin or pearl formation (Carella et al. 2015). Examples of diseases leading
2313 to granuloma formation include trematode infestations, Perkinsosis in *Ruditapes*
2314 clams, QPX in the quahog *M. mercenaria*, and fungal infections in Sydney rock
2315 oysters (Soudant et al. 2013; Allam and Raftos 2015). Diseases characterized by
2316 conchiolin formation include Roseovarius or Juvenile Oyster Disease and Brown
2317 Ring Disease in *Ruditapes* clams (Allam and Pales Espinosa 2016). On the basis
2318 of morphological differences it has been hypothesized that specialized popula-
2319 tions of hemocytes may be responsible for encapsulation (Allam and Raftos 2015).
2320 In *Ruditapes* clams infected by *P. olseni*, granulocytes secrete (from membrane-
2321 bound granules) a polypeptide named p225, which surrounds encapsulated para-
2322 sites and restricts parasite proliferation (Montes et al. 2002). Consistent with the
2323 importance of hemocytic infiltration in diseases characterized by granuloma-like
2324 formations, transcriptomic studies have shown differential expression of genes
2325 involved in hemocyte migration, pathogen recognition and binding, and inflamma-
2326 tion (McDowell et al. 2014; Allam et al. 2014; Wang et al. 2016a, b).

2327 The process of shell formation aids in encapsulation in the extrapallial cavity,
2328 playing an important role in immune defenses by preventing the penetration of
2329 pathogens through the mantle of bivalves. The process of shell formation in bivalves
2330 involves the secretion of organic molecules by secretory cells in the epithelium of
2331 the mantle outer fold, which provide a matrix for the deposition of calcium carbon-
2332 ate in a variety of structures, depending on the bivalve species. Hemocytes also
2333 play an important role in shell formation. A population of granulocytes containing
2334 calcium carbonate stored in granules migrate into the extrapallial space upon shell
2335 injury, forming aggregates at the biomineralization edge, which are incorporated
2336 into the shell as it forms (Mount et al. 2004; Zhang et al. 2012a; Li et al. 2016a). The
2337 fact that about 45% of the domains identified in the shell proteome of bivalves are
2338 related to immune function indicate the importance of the shell in bivalve immune
2339 defenses (Arivalagan et al. 2017). Among the organic compounds (1–5% of the
2340 total shell) that are embedded in the calcium carbonate structure that makes the
2341 shell, many immune-related molecules are worthy of mention, including PRRs
2342 such as galectin, scavenger receptor and C1q-related proteins, and effectors such as

phenoloxidases, proteases, and protease inhibitors (Zhang et al. 2012a; Arivalagan et al. 2017; Calvo-Iglesias et al. 2017). Moreover, genes coding for the shell proteins are differentially expressed in oysters challenged with *A. crassostreae* and in Manila clams infected with *V. tapetis*. These two bacterial pathogens preferentially attach to the inner side of the shell in bivalves, and the diseases they cause are characterized by the formation of conchiolin (McDowell et al. 2014; Allam et al. 2014).

Apoptosis and Autophagy

The Profound Implications of Apoptosis in Bivalve Physiology and Pathology

Apoptosis, a form of programmed cell death, is a highly evolutionarily conserved process involving two major distinct but converging pathways, the death-receptor-mediated pathway (an extrinsic pathway) and the mitochondrial pathway (an intrinsic pathway). Apoptosis plays an important role in immune responses by preventing the proliferation of intracellular pathogens, limiting inflammation, and being involved in the activation of certain immune cells, such as neutrophils in vertebrates (Poon et al. 2014; Creagh 2014). On the basis of changes in apoptosis levels in response to a variety of environmental stimuli, apoptosis is thought to play key physiological roles in molluscs, such as maintenance of tissue homeostasis; processing and clearing of environmental pollutants; combating of bacterial, viral, and protistan pathogens; and adjustment to exposure to insecticides, herbicides, and pharmaceuticals (Kiss 2010; Moreau et al. 2015; Romero et al. 2015; Carella et al. 2015; Zhang et al. 2016a). The functional relevance of apoptosis modulation by pathogens and environmental stressors in bivalves, however, is still unclear, since the effect of challenge/exposure on apoptosis levels is not always consistent (Soudant et al. 2013). For example, exposure to *Perkinsus* spp. modulates apoptosis in oyster and clam hemocytes and tissues, but the nature of the modulation depends on the bivalve species and the stage of infection. Advanced stages of *P. marinus* infection in *C. virginica* are generally characterized by suppression of apoptosis, which is, on the other hand, enhanced at early stages of infection (Sunila and LaBanca 2003; Goedken et al. 2005; Hughes et al. 2010; Wang et al. 2017a). Interestingly, the protozoan parasite of eastern oysters *P. marinus* expresses many antiapoptotic genes in response to exposure to oyster pallial fluid, suggesting that this parasite may be able to regulate apoptosis in the host (Pales Espinosa et al. 2014). Basal rates of apoptosis in oysters also differ between the source of hemocytes, ranging from 5–25% in hemocytes in hemolymph to up to 50% in hemocytes within tissues (Sunila and LaBanca 2003; Goedken et al. 2005; Cherkasov et al. 2007; Sokolova 2009)

Main Molecular Players in the Apoptotic Process

Although the major molecules and pathways of apoptosis appear to be conserved between bivalves and other species on the basis of genomic studies (Fig. 17), only a few of them have been characterized functionally. These include the executioner caspase-3 and caspase-1 (caspase-7-like) from *M. gigas*, which appear

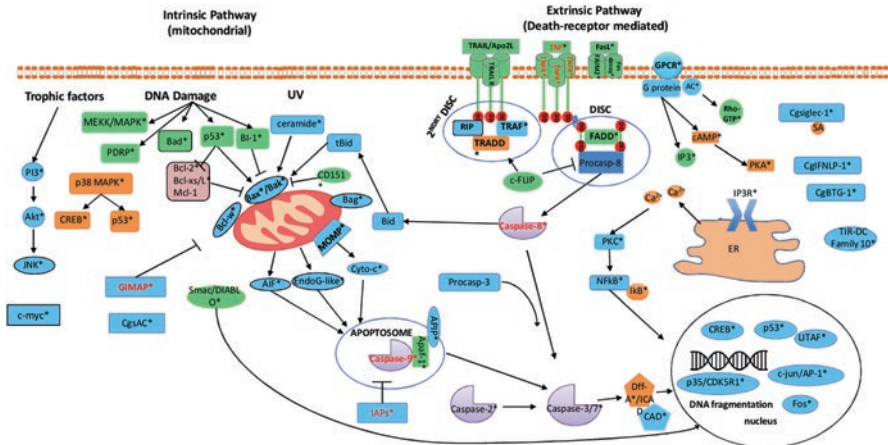


Fig. 17 Apoptosis pathway molecules, with those identified in molluscs indicated with asterisks. Genes identified only in *M. gigas* are prefixed by “Cg” and expanded gene families are shown in red *text*. Molecules that have been only preliminarily identified in molluscs via the eastern oyster genome annotation are denoted with “-like” and genes that been implicated in caspase-independent mechanisms are outlined in black. (Kögel et al. 2013)

2384 to act as intracellular LPS receptors (Xu et al. 2016b; Wang et al. 2017a).
 2385 Interestingly, bivalves may possess a caspase-independent apoptotic pathway,
 2386 hypothesized to be involved in apoptosis induced by the protozoan parasite *P. marinus*
 2387 (Wang et al. 2017a).

2388 Several gene families involved in the apoptotic process have experienced lineage-
 2389 specific expansions, including tumor necrosis factors (TNF), tumor necrosis factor
 2390 receptors (TNFRs), caspase 8, inhibitor of apoptosis proteins (IAPs), cysteine-
 2391 aspartic proteases (caspases), and GTPase of the immune-associated proteins
 2392 (GIMAPs) (Zhang et al. 2012a; Qu et al. 2015b; McDowell et al. 2016; Li et al.
 2393 2016b; Wang et al. 2017a). Enhanced genetic diversity of these apoptosis pathway
 2394 gene families may allow for more diverse but also pathogen-specific functional
 2395 responses to disease and therefore increase the ability of apoptosis pathways to aid
 2396 in stress mitigation and increase survival. For example, while oyster *M. hongkon-*
 2397 *gensis* Chcaspase8s is upregulated with bacterial challenge, *M. gigas* Cgcaspase8–2
 2398 responds to viral challenge but not bacterial challenge (Wang et al. 2017a).

2399 Two of these gene families, coding for IAPs and GIMAPs (also known in plants
 2400 as immune-associated nucleotide-binding genes, or IANs), are of particular interest
 2401 because of their known critical apoptosis regulatory roles in other organisms, their
 2402 high level of transcript diversity in bivalves, and their demonstrated differential
 2403 expression in bivalves after immune challenge. The GIMAP/IAN family has 26
 2404 annotated members in *M. gigas*, similar to the predicted 26–28 GIMAPs in the east-
 2405 ern oyster, several of which are downregulated in eastern oyster juveniles after chal-
 2406 lenge with *Roseovarius* Oyster Disease (ROD), suggesting an upregulation of
 2407 apoptosis (McDowell et al. 2016). The functional significance of this expansion in
 2408 bivalves is unknown, but GIMAPs are known to play key roles in regulation of

lymphocyte survival, T-cell selection and homeostasis, phagolysosomal processing and membrane trafficking in vertebrates, and pathogen resistance in the model plant system *Arabidopsis* (Weiss et al. 2013; Webb et al. 2016).

The CgIAP family represents another expanded apoptosis-related family in oysters, with 48 gene members, likely the result of tandem gene duplications (Qu et al. 2015b; Zhang et al. 2016a; Wang et al. 2017a). IAP proteins have known roles in apoptosis inhibition by interacting with caspases, and direct evidence of this interaction has been shown for CgIAP2, where its characteristic BIR2 domain directly interacts with Cgcaspace-2 (Zhang et al. 2011b; Qu et al. 2015b). Bacterial challenges of the Pacific oyster with the bacterial pathogen *V. anguillarum* have shown increased gene expression over time (Zhang et al. 2011b; Qu et al. 2015b). When two families of Pacific oyster with different susceptibility to ostreid herpesvirus-1 (OsHV-1) were exposed to this virus, CgIAP expression was significantly upregulated in both families though with higher levels of expression in the family most sensitive to OsHV-1 (Zhang et al. 2016a). Another gene family with potential roles in apoptosis worth mentioning here is the TIR-DC family 10, characterized by the presence of two baculovirus inhibitor of apoptosis protein repeat (BIR) domains. This gene family has been found only in bivalves (Gerdol et al. 2017).

Potential Involvement of Autophagy in Immune Response

Not much is known about the role of other forms of programmed cell death in innate immune responses in bivalves. Autophagy, which is involved in innate immunity against intracellular pathogens in vertebrates, is induced in oysters in response to bacterial and viral challenge, as well as environmental stimuli such as changes in salinity, hypoxia, toxins, or lack of nutrition (Carella et al. 2015; Wang et al. 2017a). Genes in the autophagy (ATG) pathway have been described in Pacific oysters, and autophagy is involved in survival after challenge with OsHV-1 and *V. aestuarianus*, two pathogens commonly associated with summer mortality in the Pacific oyster, *M. gigas*. Interestingly, while challenge with OsHV-1 led to induction of autophagy, challenge with *V. aestuarianus* resulted in inhibition of autophagy (Moreau et al. 2015).

Overview of the Immune System of Other Molluscan Classes

We have so far outlined the main molecular and cellular components of the immune system of Bivalvia, the second largest molluscan class. Bivalves have been the subject of extensive immunological research over the past few decades, motivated by the high socioeconomic importance of edible species, their widespread distribution, and their amenability for laboratory research. The largest molluscan class in terms of the number of species, gastropods, has attracted considerable attention for similar reasons. These animals—adapted to the freshwater, marine, and terrestrial environments—present astounding morphological diversification, including snails, slugs, limpets, nudibranchs, and others. This diversity can be correlated with the adaptation of lineage-specific strategies for immune defense, which in some cases has led

2450 to the acquisition of unique traits and advanced mechanisms, such as the somatic
2451 diversification of FREPs. The main features of the gastropod immune system are
2452 presented in detail in Chap. 12.

2453 Unfortunately, very little information is available concerning several aspects of
2454 the basic biology of the other molluscan classes, such as aplacophorans, monoplacophorans, polyplacophorans, and scaphopods. Consequently, the immune systems of these animals and the possible peculiar survival strategies that might have been developed in these taxa during their evolution are presently unknown. The few data collected so far concern cellular immunity of chitons, where phagocytic cells located in circulating hemolymph, as well as in connective tissue, seem to bear remarkable immune recognition properties (Crichton et al. 1973; Crichton and Lafferty 1975).

2462 The exception is represented by cephalopods, which have historically attracted
2463 major scientific attention, in particular due to their complex nervous system, intelligence, and learning skills. However, immune studies are also emerging, as evidenced by the conspicuous amount of literature produced on this subject over the past few years. The following sections will review the most distinctive peculiarities of the cephalopod immune system of these fascinating animals.

2468 **A Short Journey in the Immune System of Cephalopods**

2469 Cephalopods (i.e., nautilus, cuttlefish, squid, and octopuses) comprise over 800
2470 living species (Sweeney and Roper 1998), about 300 belonging to Octopodidae
2471 (Jereb and Roper 2016) and including several species complexes (Allcock et al.
2472 2011; Amor et al. 2014; Cheng et al. 2014; Sales et al. 2017). They are considered
2473 to rival vertebrates (Packard 1972) for physiological adaptations, complex neural
2474 organization, and behavior (Jereb and Roper 2005, 2010; Huffard 2013; Jereb and Roper 2016; Marini et al. 2017). The immune system of cephalopods consists of innate mechanisms and includes cellular and humoral defenses (Ford 1992; Castillo et al. 2015; Pila et al. 2016).

2478 **The Highly Complex Circulatory System of Cephalopods**

2479 This molluscan taxon is the sole group of animals, other than vertebrates, to enjoy a
2480 fully enclosed high-pressure blood system, an example of convergent evolution
2481 (Wells 1983). Three hearts (one systemic and two branchial) move blood through an
2482 extraordinarily complex network of arteries, veins, and capillaries (Fig. 18), thus
2483 representing “a triumph of engineering over design” (Wells and Smith 1987). An
2484 overview on the physiology of the circulatory system and its development is available in a number of works (Naef 1928; Boletzky 1968; Wells 1983; Budelmann et al. 1997).

2487 **Morphology and Function of Cephalopod Hemocytes**

2488 In contrast to bivalves, the circulating blood (hemolymph) in cephalopods turns
2489 blue when oxygenated (Wells 1983) because of the presence of hemocyanin. The

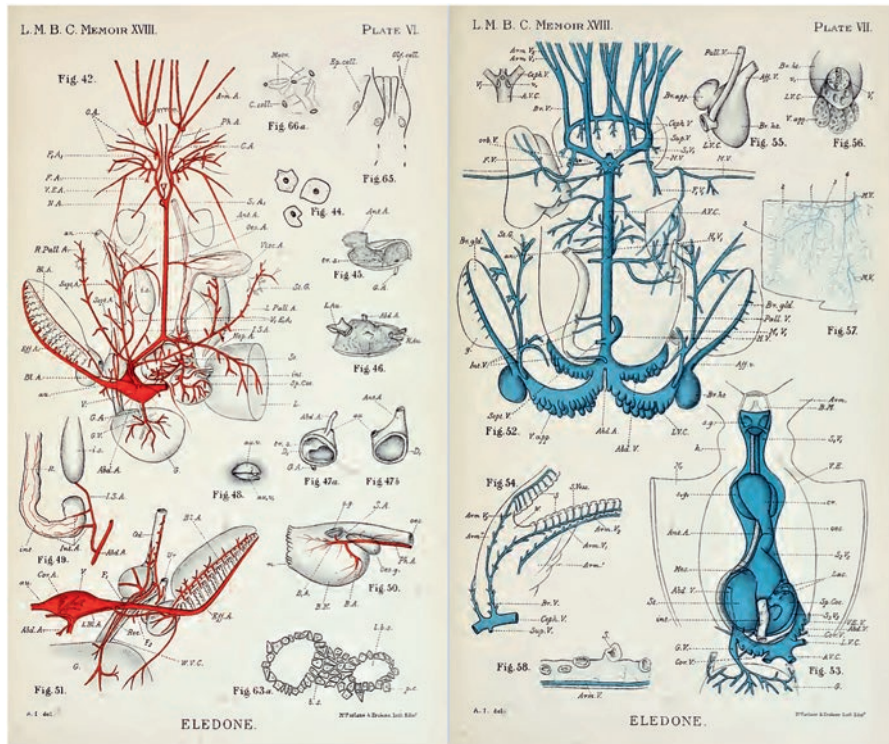


Fig. 18 General outline of the cephalopod circulatory system as exemplified for *Eledone cirrhosa* by Isgrove (1909). In coleoids (cuttlefish, squid, and octopuses), three hearts exist: the systemic heart pumps oxygenated blood (red); the two branchial hearts move blood through the capillaries of the gills (Wells 1983). An extraordinary network of arteries (red), veins, and capillaries exist in cephalopods. The venous system (blue, right) is shown with the principal cephalic vein, pallial veins, three venae cavae, and a large perivisceral blood sinus. In *Nautilus* the circulatory system (not shown) is characterized by large venous spaces, i.e., the pericardium (Owen 1832), differently from what occurs in coleoids

hemocytes—also named leukocytes (Bolognari 1949, 1951), amoebocytes, or granulocytes (Budelmann et al. 1997)—are the “key” cellular components of the immune system of cephalopods. In an analogy to other molluscs, the identification of cellular types in cephalopods and their characterization is often contradictory, since their classification may be biased by the technique that is utilized (Vieira et al. 2017). Furthermore, the variability in observed cells may reflect the physiological status of the animals (Bolognesi and Fenech 2012; Locatello et al. 2013; Castellanos-Martínez et al. 2014b). Attempts to develop a consensus on the nomenclature of hemocytes have been made for some molluscan species (Cheng 1984) but are still lacking for cephalopods. However, we outline their general description on the basis of the few reports available (Fig. 19).

Budelmann et al. (1997) described two types of cells in cephalopod hemolymph. The first type of hemocytes are round or oval cells, with an elongated V-shaped

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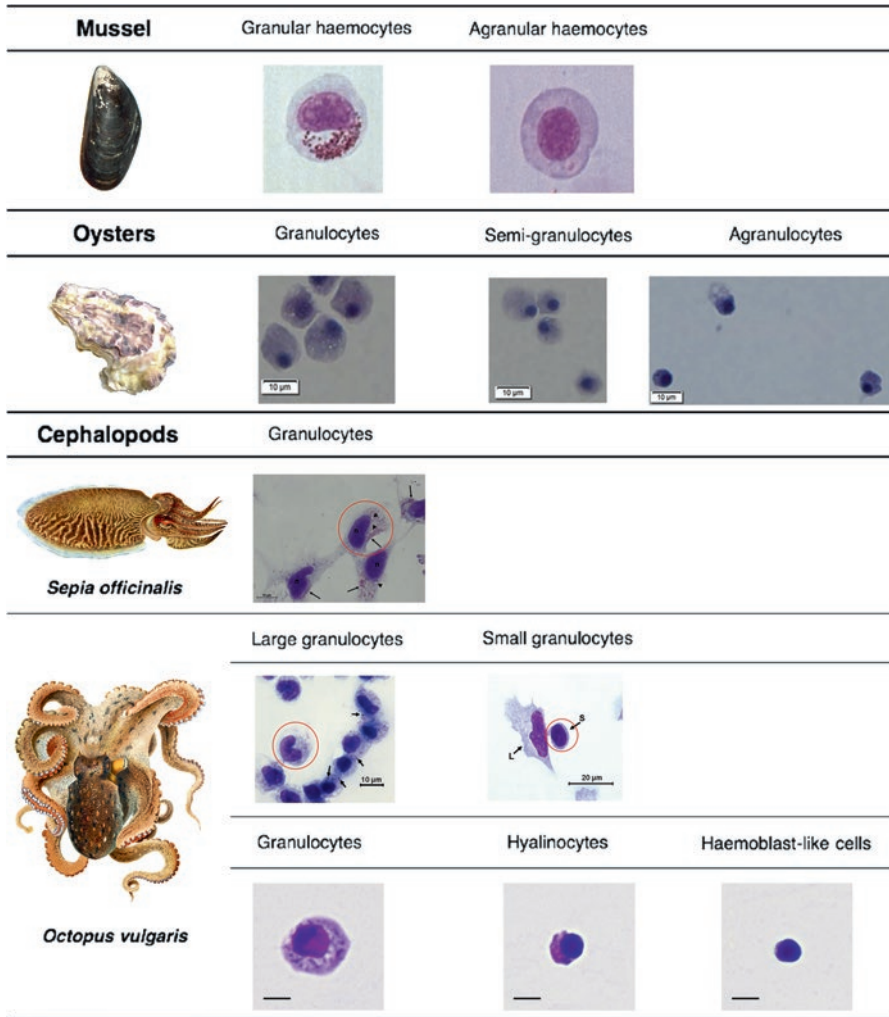


Fig. 19 The different types of hemocytes identified in cephalopod molluscs. Examples from bivalves are provided for comparison. See also Table 2 for further detail. The drawings are based on the original descriptions provided for mussels by Bolognesi and Fenech (2012), for oysters by Wang et al. (2017a), and for the cephalopods *Sepia officinalis* and *Octopus vulgaris* by Le Pabic et al. (2014a) and by Castellanos-Martínez et al. (2014b) and Troncone et al. (2015), respectively

AUT7

2503 nucleus, known to extend large pseudopods producing amoeboid locomotion and
 2504 capable of a phagocytic response and the secretion of pore-forming lysins and cyto-
 2505 toxic oxygen radicals by exocytosis of small granules (Budelmann et al. 1997). The
 2506 second type include vacuolized round cells, which are relatively sessile (they do not
 2507 display pseudopods), accumulate into large agglomerates, and are similar in size
 2508 and shape to hemocytes. Each cell has either numerous small lysosomes or a single
 2509 large lysosome. They are able to incorporate particles through micropinocytosis.

Vacuolized round cells are thought to correspond to the pore cells of other molluscs and to the monocyte–macrophage system of vertebrates (Budelmann et al. 1997). 2510
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Troncone and colleagues (2015) recognized three types of hemocytes in *Octopus vulgaris*: hemoblast-like cells, hyalinocytes, and granulocytes. According to those authors, the hemoblast-like cells are the smallest ones, not motile and without pseudopodia. Hyalinocytes are described as variable in size, with a rounded or oval nucleus, and no or few granules and vacuoles of different diameters in the cytoplasm. The cells are capable of amoeboid movement and can form pseudopodia. Granulocytes are variable in size, highly amoeboid, and able to form many long filopodia. Granulocytes are described as being characterized by an eccentric oval nucleus and numerous cytoplasmic granules of different sizes (endoplasm), while no granules are found in the ectoplasm (Troncone et al. 2015). 2512
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In coleoids (cuttlefish, squid, and octopuses) the hemocytes originate from the white body (Bolognari 1949, 1951; Cowden 1972; Cowden and Curtis 1973), a multilobed organ covered by a thin layer of connective tissue surrounding, as cushions, the optic lobes and located in the “orbits” in the head of the animal. White bodies extend between the medial external surfaces of the eyes and the skull, and encapsulate the “central brain.” The morphology, structure, and function of this organ were originally described by Bolognari (1949, 1951). A pioneering attempt to isolate the cellular components and to estimate their mitotic activity and culturing in vivo was carried out by Necco and Martin (1963). Further characterization of this organ in the octopus was provided by Cowden (1972), including ultrastructural analysis (Cowden and Curtis 1974). A functional description of the white bodies is also available for *S. officinalis* (Claes 1996) and for sepiolids (see below), while no analogous structures are known in *Nautilus*, to the best of our knowledge. 2522
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After histological examination, the white bodies appear as a network of connective fibers, blood vessels, and vascular varicosities in which a mass of cellular strings is observed. These are believed to be precursors of the hemocytes (Bolognari 1949, 1951; Cowden 1972). Leukocytes at different stages of “maturity” are identified in the white bodies of *O. vulgaris* (Cowden 1972). According to the classical ultrastructural description, the hemocytoblasts (or reticulum cells of the white bodies) are characterized by an abundant “rough” endoplasmic reticulum, mitochondria, and Golgi, and an irregular large vesicle reported to “contain some internal fibrillar material condensed” in some areas (Cowden and Curtis 1974). These authors also provided a thorough description of other cellular characteristics, and of the transformation of hemocytoblasts to form primary and secondary leukoblasts, and finally mature leukocytes. This last cell type appears to have a folded nucleus containing an abundance of condensed chromatin ... and dense extrachromosomal aggregates. The cytoplasm contains a number of electron-dense, rounded inclusions,” possibly derived from the reduction of vesicles characterizing the hemocytoblasts (Cowden and Curtis 1974). 2535
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Two main groups of hemocytes are recognized in cephalopods: cells containing many granules (granular hemocytes or granulocytes), and cells with few or no granules (agranular hemocytes, agranulocytes, or hyalinocytes). These correspond to the two types of cells described by Budelmann et al. (1997). 2551
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2555 The octopus hemocytes (sensu lato) act as immunocompetent cells in the hemo-
2556 lymph (Ford 1992). They are involved in the recognition and elimination of poten-
2557 tial pathogens through phagocytosis, encapsulation, infiltration, and production of
2558 reactive agents with oxidizing capacity (i.e., reactive oxygen species (ROS) and
2559 reactive nitrogen species (RNS)). Hemocytes are also involved in scar formation,
2560 wound healing, and tissue repair by migrating to the site of injury, increasing in
2561 number and activity and forming plugs at the wound site to prevent hemolymph loss
2562 (Polglase et al. 1983; Féral 1988; Shaw et al. 2016; Imperadore et al. 2017).

2563 The composition and number of hemocytes are highly variable both among spe-
2564 cies (Le Pabic et al. 2014a) and between individuals (Malham et al. 1998, 2002;
2565 Locatello et al. 2013; Roubledakis et al. 2017) in an analogy to other molluscs
2566 (Anisimova et al. 2017). The number of circulating hemocytes appears variable
2567 among different individuals following “stressors” such as handling (Malham et al.
2568 1998, 2002), immune challenge (Locatello et al. 2013), or life stages (Roubledakis
2569 et al. 2017). Phagocytosis is known as the primary immune response of hemocytes
2570 and has been reported in various species, e.g., *Sepia officinalis* (Le Pabic et al.
2571 2014a), *O. vulgaris* (Novoa et al. 2002; Rodríguez-Domínguez et al. 2006), and
2572 *Eledone cirrhosa* (Malham et al. 2002).

2573 **Molecular Immunology Studies Are Still at Their Embryonal Stage** 2574 **in Cephalopods**

2575 The humoral defense is achieved through soluble molecules (Castillo et al. 2015)
2576 such as opsonins, agglutinins, proteolytic enzymes, protease inhibitors, antimicro-
2577 bials or cytotoxic compounds, phenoloxidase, and its intermediate synthesis prod-
2578 ucts, which are in part similar to those described in detail for bivalve molluscs in the
2579 previous sections (Rögener et al. 1985; Lacoue-Labarthe et al. 2009; Alpuche et al.
2580 2010; Le Pabic et al. 2014b; Roubledakis et al. 2017). However, as evidenced by
2581 recent transcriptomic approaches, a relevant fraction of lineage-specific genes with
2582 unknown function exists in cephalopods. This observation is particularly relevant
2583 considering large high number of unknown mRNAs identified in the transcriptomes
2584 obtained from *O. vulgaris* hemocytes (Castellanos-Martínez et al. 2014a) and the
2585 white bodies of the sepiolid *Euprymna tasmanica* (Salazar et al. 2015).

2586 Salazar and colleagues (2015) also provided a description of putative *Euprymna*
2587 immune-related genes, identifying—for example—NF- κ B and components of the
2588 Toll signaling pathway, pattern recognition proteins, TNF-receptor-associated factors,
2589 and proteins denoting membrane attack complex/perforin domains, which in large
2590 part mirror those described in bivalves (see sections “Recognition, Agglutination,
2591 and Opsonization,” “Signaling and Regulatory Pathways,” and “Humoral Immune
2592 Effectors”).

2593 Although the cellular and “humoral” components of cephalopods have been
2594 studied extensively (Castillo et al. 2015), our knowledge of cephalopod immunity is
2595 still in its infancy. In brief, evidence exists for (1) a possible role of the white bod-
2596 ies as a hematopoietic and immune organ, and (2) the presence of different types
2597 and numbers of circulating cells after challenges. Molecular fingerprints for the
2598 immune response have been so far explored only in a limited way (Collins et al.

2012b; Castellanos-Martínez et al. 2014a; Salazar et al. 2015). Preliminary evidence collected over the past few years suggests that cephalopod immunity, like that of other molluscs (see Chap. 12, section “Molluscs Exhibit Immune Priming with Intermediate Degrees of Specificity, and Involving a Plethora of Mechanisms” for a detailed discussion), may show some form of memory. The analysis of the plasticity of innate immune responses in these fascinating organisms is one of the most important future avenues for cephalopod science and, in particular, for immunological studies.

Bobtail Squid as a Model for the Study of Bacterial Symbiosis

The capacity of an animal's immune system to recognize and remove nonself is crucial for its survival and, by tradition, this has been the context in which we have defined immune components, and even how we have designed experiments to understand their roles. This is easy to envision when one considers the detrimental presence of microorganisms to the host, either because of nutrient competition or tissue damage. This kind of association is, by definition, usually considered pathogenic, but this is just one of the three types of symbiotic relationships an animal can establish with another species. The other two types are commensalism (where one species benefits and the other neither benefits nor gets harmed) and mutualism (a type of beneficial relationship between two species, in which both obtain some type of benefit). An animal can establish any one of these associations with the immense variety of microorganisms that share its ecological niche, i.e., bacteria, protozoans, helminths, fungi, or viruses. This section focuses on the major findings resulting from 30 years of study of one of these beneficial interactions, the *Euprymna scolopes*–*Vibrio fischeri* symbiosis. This model has somewhat challenged our vision on the role of the immune system in metazoans.

The squid–*Vibrio* symbiosis is one of the most studied and better understood binomial associations between an animal and its bacterial symbionts (McFall-Ngai 2008; Castillo et al. 2015; McAnulty and Nyholm 2017; Stabb and Visick 2013; Norsworthy and Visick 2015; Mandel and Dunn 2016). In addition, modern sequencing and proteomic technologies have recently allowed the identification of several molecular players participating in the squid's immune system (Chun et al. 2006; Wier et al. 2010; Collins et al. 2012a, b; Kremer et al. 2013; Salazar et al. 2015). The next paragraphs contain a brief description of this symbiosis, followed by specific information on the molecular players involved, with emphasis on the squid host immune components.

Main Features of the Squid–*Vibrio* Symbiosis

This mutualistic symbiosis involves the squid *E. scolopes*, also known as the bobtail squid, a relatively small (adult mantle length ~3–4 cm), nocturnal sepiolid species, native to the Hawaiian archipelago (Berry 1912) (Fig. 20, panel b1). The symbionts are Gram-negative marine Proteobacteria members of the Vibrionaceae family, capable of producing bioluminescence by means of luciferase activity under

2640 quorum-sensing conditions. The bacteria reside in the squid in a specialized bilobed
2641 structure called the light organ (LO) (McFall-Ngai and Montgomery 1990). The LO
2642 is localized on the ventral side of the animal and inside the muscular mantle, just
2643 above the funnel or siphon (Fig. 20, panels a1-2, b1-2). In this location, the LO is
2644 flushed with ocean water during regular breathing or swimming movements of the
2645 mantle. Microorganisms present in the water, including *V. fischeri*, come in direct
2646 contact with the LO surface which, in response to bacterial compounds such as
2647 lipopolysaccharide (LPS) and peptidoglycan (PG), secretes mucus to which bacte-
2648 ria attach and start aggregating (Nyholm et al. 2000; Foster et al. 2000) (Fig. 20,
2649 panel a2). Several studies have found that the mucus contains chemoattractants
2650 (N-acetylgalactosamine and N-acetylneuraminic acid) (Altura et al. 2011; Mandel
2651 et al. 2012), as well as soluble antimicrobials and nitric oxide (Davidson et al. 2004;
2652 Kremer et al. 2013). Together, these host-derived products are thought to favor
2653 *V. fischeri* attachment while discouraging nonsymbiont organisms from collecting at
2654 the site. In addition, the LO of juvenile *E. scolopes* is characterized by having on
2655 either side a pair of appendages made from densely ciliated epithelial cells where
2656 the mucus is held (Fig. 20, panel b2). The beating cilia help to move aggregated
2657 bacteria and particles toward the three open pores that serve as the entrance to the
2658 internal part of the LO (Nyholm et al. 2000). As *V. fischeri* cells enter the LO through
2659 a pore, they encounter a narrow, ciliated duct that eventually opens into a series of
2660 branched and closed-ended spaces known as crypts. Here, the bacteria reach their
2661 final place of residence. The lumen of the crypts is covered by epithelial cells with
2662 multiple microvilli that secrete mucus and other host molecules, and that, once the
2663 squid is colonized, will be in close contact with the bacterial symbionts. Not many
2664 *V. fischeri* cells are necessary to seed the LO, as it has been estimated that as few as
2665 3–6 cells can start the colonization of each lobe of this organ (Wollenberg and Ruby
2666 2009). If the bacteria colonizing the LO are capable of producing light, about 12 h
2667 after their arrival in the crypts, the combination of light and microbial products is
2668 recognized by the host and a developmental signal for a series of programmed mor-
2669 phological changes is initiated. This program includes the following events:
2670 (1) apoptosis of the ciliated appendages; (2) fusion of the three pores and ducts into
2671 a single one; and (3) an increase in microvilli and swelling of the crypt epithelia
2672 (McFall-Ngai and Ruby 1991; Nyholm and McFall-Ngai 2004). The overall result
2673 is irreversible loss of the lateral appendages from the LO surface and physiological
2674 changes in internal structures over the next 4 weeks that will ensure the maintenance
2675 of the newly acquired symbionts (Koch et al. 2014) (Fig. 20, panel b2).

Fig. 20 (continued) on the ciliated appendages. **(b1)** Adult female *E. scolopes* squid side view; the transparent window allows us to see the light organ and accessory nidamental gland locations. **(b2)** Adult light organ with crypts. **(b3)** Host–symbiont interaction zone in adult squid consisting of crypt epithelial cells with microvilli and migrating hemocytes. AE appendage epithelia, ANG accessory nidamental gland, BS blood sinus, CE crypt epithelia, Cr crypts, Hc hemocyte, IS ink sac, Le lens, LO light organ, Mu mucus, NA nidamental gland, P pore, Vf *Vibrio fischeri* bacteria, YF yellow filters

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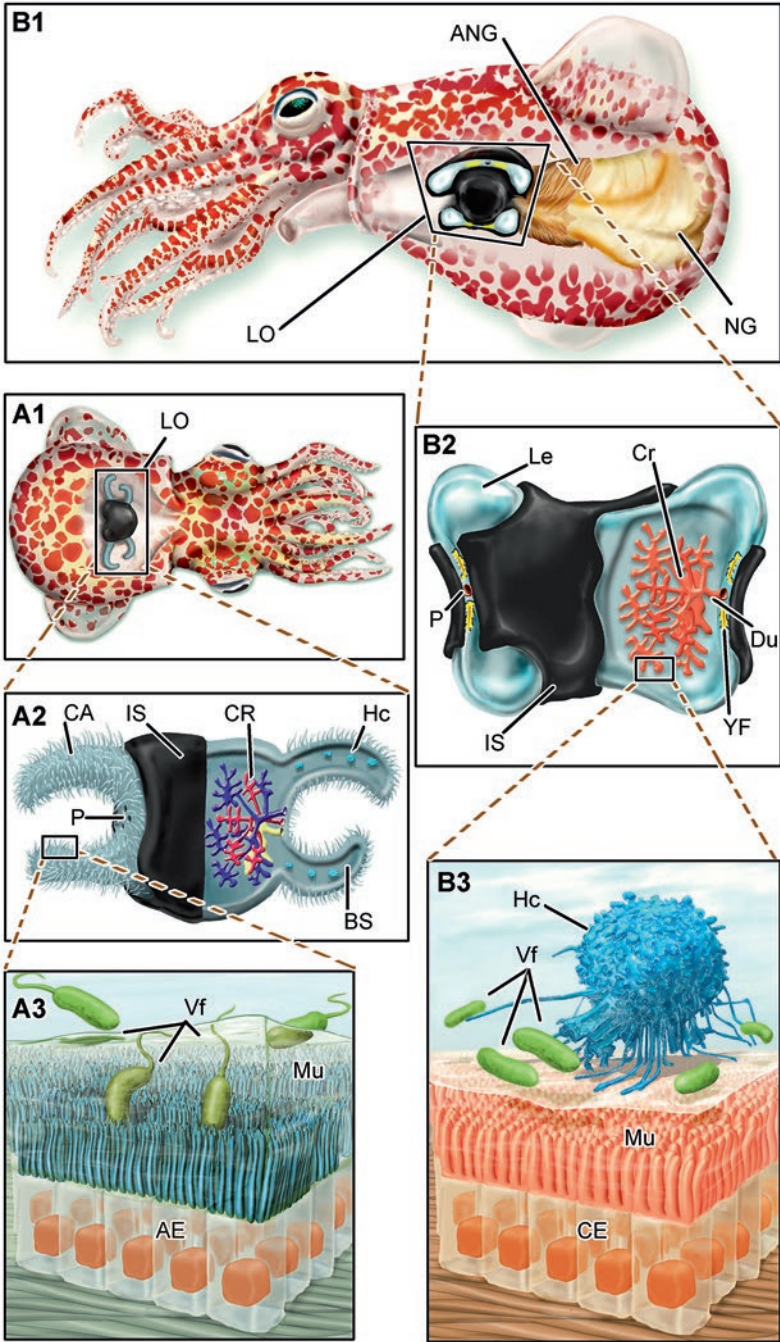


Fig. 20 *Euprymna scolopes* squid and tissues associated with bacterial symbiosis. (a1) Juvenile *E. scolopes* squid ventral view. (a2) Juvenile light organ with crypts and ciliated appendages. (a3) Host–symbiont interaction zone in juvenile squid, consisting of the surface of epithelial cells

2676 Once this association between *E. scolopes* juvenile squid and bacteria is estab-
2677 lished, the symbiosis will be maintained for the duration of the animal's life (Nyholm
2678 and McFall-Ngai 2004). An important characteristic of this symbiosis is the diel
2679 rhythm, which consists, among other things, of daily expulsion of the majority (90–
2680 95%) of the bacterial population from the LO at dawn (Lee and Ruby 1994;
2681 Boettcher et al. 1996; Nyholm and McFall-Ngai 1998). This thick exudate contains
2682 live and dead *V. fischeri* cells and also some host hemocytes and epithelial cells
2683 (Graf and Ruby 1998; Nyholm and McFall-Ngai 1998). In the 8 h following the
2684 emptying of the LO, the remaining population of symbionts quickly grows and
2685 divides inside the crypts, until they reach a density high enough to enable quorum
2686 sensing, thereby becoming luminescent again at night (Nyholm and McFall-Ngai
2687 1998). It is suggested that the squid uses this light to camouflage itself from poten-
2688 tial predators and preys. This is suggested by the presence of several tissues in the
2689 LO, including a lens and a reflector, that allow the animal to control the amount of
2690 light emitted, with the purpose of replicating down-welling light from the moon and
2691 stars. This behavior is known as counterillumination and prevents the production of
2692 a shadow during swimming in the water column. (Ruby and McFall-Ngai 1992;
2693 Jones and Nishiguchi 2004).

2694 The *Euprymna scolopes*–*Vibrio fischeri* mutualism offers advantages over other
2695 animal model systems for understanding of the physiology and molecular mecha-
2696 nisms of animal–bacterial beneficial associations (Ruby 1999; McFall-Ngai 2008;
2697 Lee et al. 2009). This is mainly because this it is a binary association (Ruby and Lee
2698 1998; Mandel 2010), where both organisms can be cultured separately, thereby
2699 allowing manipulation of the bacterial introduction, and because the bacterial sym-
2700 biont is genetically tractable and introductions of mutations and markers are modi-
2701 fications relatively easy to achieve (Ruby 1999; McFall-Ngai 2008; Lee et al. 2009).
2702 Moreover, the direct contact and interaction between the two players (host and bac-
2703 teria) in this symbiosis occur extracellularly, meaning that the bacteria never breach
2704 the epithelial integrity of the host tissues. Thus, their interaction occurs via secreted
2705 molecules and by means of cell surface molecules both at the level of juvenile squid
2706 ciliated appendages (Fig. 20, panel a3) and inside the juvenile and adult LO crypt
2707 epithelia (Fig. 20, panel b3).

2708 **The Fundamental Role of Hemocytes in the Establishment** 2709 **of Symbiosis**

2710 Hemocytes play a major role in the establishment and maintenance of this interac-
2711 tion. As detailed in the previous section, these are motile cells that circulate through
2712 the squid vasculature and can reach sites where the bacteria are, and interact with
2713 them. For a review on the role of hemocytes on the squid–*Vibrio* symbiosis, the
2714 reader is directed to a recent publication by McAnulty and Nyholm (2017). The
2715 squid hemocytes play a pivotal role right from the initial stages of colonization.
2716 First, the presence of the symbiont causes the proliferation of hemocytes, the num-
2717 ber of which peaks about 36 h postcolonization (Koropatnick et al. 2007).
2718 Furthermore, these cells play an active role during the apoptotic regression of the
2719 LO epithelia, a behavior that is accredited to the presence of *V. fischeri* products

released in the LO crypts. Specifically, and in response to *V. fischeri* outer membrane vesicles (OMV) (Aschtgen et al. 2016) and PGN-tracheal cytotoxin (TCT) (Koropatnick et al. 2004), squid hemocytes move from the circulation and migrate to the sinus space in the ciliated appendages. This migration is also accompanied by upregulation of transcripts involved in protein degradation, suggesting that these cells are involved in facilitating the apoptosis and restructuring of epithelial cells during the LO metamorphosis (Koropatnick et al. 2007). This process is aided by the activity of a matrix metalloproteinase (Koropatnick et al. 2014), as suggested by the upregulation of this enzyme in hemocytes and the LO tissues of symbiotic squids (Chun et al. 2006; Collins et al. 2012b; Schleicher et al. 2014).

In vitro studies have also shown that *E. scolopes* hemocytes can selectively recognize, bind, and engulf bacteria, while showing a degree of tolerance of *V. fischeri* in comparison with other marine bacteria (Nyholm and McFall-Ngai 1998; Nyholm et al. 2009). This recognition is modulated by unknown factors secreted by the symbionts (Nyholm et al. 2009). In addition, to discriminate between bacterial species, hemocytes of adult squid also appear to be “trained” to tolerate the symbiont, as hemocytes from antibiotic-treated squids lose their symbiont recognition capacity and bind *V. fischeri* cells more readily (Nyholm et al. 2009).

Several transcriptome and proteomic studies comparing hemocytes from colonized and noncolonized animals have been performed, which enabled the sequence identification of a number of soluble immune factors (Collins et al. 2012b). Among these, a matrix metalloprotein, a cephalotoxin, a galectin, and a soluble peptidoglycan recognition protein (EsPGRP5) were found to be downregulated in cured hemocytes, while EsC3 transcripts could not be detected in symbiotic animals. These results suggested that the presence of the symbiont modulates the host immune system to avoid its removal (Collins et al. 2012b). The complement component C3 and other complement-like molecules—including CD109 antigen (Yazzie et al. 2015), other thioester-containing proteins, and alpha-2-macroglobulin (Collins et al. 2012b, personal observations)—have also been identified in hemocytes, but their specific role in symbiosis have not been described yet. Like C3, some of these transcripts appear to be modulated in symbiotic squid compared with those not exposed to bacteria, as was the case for CD109 antigen (Yazzie et al. 2015). Furthermore, several transcripts with homology to known PRRs have been identified in hemocytes, including PGRPs and TLRs (Collins et al. 2012b). Hemocyte–proteomics studies have also revealed at least 37 differentially expressed proteins in the adult symbiotic animals compared with cured squid. Some of these are known to be involved in immune-related functions, most notably cathepsins, lysosomal proteins, and various proteases (see section “Proteases and Protease Inhibitors”) (Schleicher et al. 2014). It is also worth noting that—as mentioned in section “A Short Journey in the ‘Immune System’ of Cephalopods,”—like all other cephalopods, squid appear to possess a well conserved immune signaling machinery. It is, however, still unclear how these immune sensors and effector molecules modulate or are modulated by the presence of the bacterial symbiont.

Hemocytes are not only important during the squid colonization process; they are also central to the homeostatic maintenance of the symbiosis. Recent studies

2765 have found that hemocytes have cytoplasmic vesicles that contain chitin (Heath-
2766 Heckman and McFall-Ngai 2011). Chitin is an abundant carbohydrate polymer in
2767 marine environments and a food source for many planktonic organisms, including
2768 bacteria. It has been suggested that hemocytes deliver this nutrient into the LO
2769 crypts during the evening and night hours, when the bacteria population is at its
2770 higher densities, to provide nutrients to the symbionts. In return, the symbionts
2771 utilize this resource via fermentation and, as a consequence, acidify the crypt spaces
2772 to a pH of about 5.5 (Kremer et al. 2014). Hemocyanin, the squid's blood pigment
2773 and oxygen carrier (Markl 2013), releases oxygen under acidic conditions. Since
2774 bacteria need oxygen to produce light, as in the luciferase reaction, the hemocytes
2775 are providing a source of food to the bacteria that will in turn promote the formation
2776 of the proper environment for light production, which the host uses for its nocturnal
2777 activities (Kremer et al. 2014).

2778 The large number of putative immune molecules identified in the aforementioned
2779 sequencing studies confirm the involvement of hemocytes in the host response to
2780 *V. fischeri* colonization. It is also interesting to note that multiple genes associated
2781 with cytoskeletal and lysosomal activities are modulated, reflecting the develop-
2782 mental and morphological changes the host undergoes in response to its association
2783 with its bacterial partner. For more information, the reader is directed to the primary
2784 study sources (Goodson et al. 2005; Collins et al. 2012b; Schleicher et al. 2014;
2785 Salazar et al. 2015).

2786 **The Immune Role of the Light Organ**

2787 In addition to hemocytes, other squid tissues express immune-related molecules.
2788 Many of these were originally discovered during an extensive analysis of expressed
2789 sequence tags (ESTs) from the juvenile LO at different times after colonization
2790 (Chun et al. 2006), in the transcriptomes of adult LOs at different times during the
2791 diel rhythm (Wier et al. 2010), or in a data set of LO transcripts differentially
2792 expressed in animals for 3 h to the symbiont (Kremer et al. 2013). The following
2793 paragraphs will describe these molecules and their suggested role in the symbiosis.

2794 **Receptors and Sensor Molecules**

2795 Several receptors were identified in the juvenile LO, including four PGRPs
2796 (PGRP1–4) (Chun et al. 2006), whose general role in invertebrate immunity is sum-
2797 marized in section “[Other Membrane-Bound Immune Receptors](#).” PGRP1 was
2798 found to be localized in the cytoplasm of surface epithelial cells and translocated to
2799 the nucleus, a change associated with the apoptosis of the LO appendages (Troll
2800 et al. 2009). PGRP2 was secreted in mucus and found to have PGN-catalytic activ-
2801 ity, suggesting an antimicrobial purpose (Troll et al. 2010). Furthermore, PGRP2
2802 was also secreted inside the LO crypts but only after colonization, possibly to aid in
2803 removal of PGN products released by the symbionts. Finally, PRGP3 had a glyco-
2804 phosphatidylinositol (GPI)-anchoring site, and PRGP4 was a true transmembrane
2805 receptor (McFall-Ngai et al. 2010). Additional PRRs identified in *E. scolopes* are
2806 members of the LBP/BPIs family of proteins (see section “[Lysozymes, BPIs and](#)
2807 [Other Pore-Forming Molecules](#)”). Not much is known about the function of these

sensor/effector molecules in squid, other than the fact that a BPI transcript was upregulated during LO apoptosis in symbiotic squid. Because of its localization in the LO crypts, this BPI might play a similar antimicrobial role to the PRGPs (Krasity et al. 2011).

Complement System

As mentioned earlier, bivalve molluscs possess a prototypical complement system (see section 4.4). Furthermore, C3-like transcripts have been found in squid hemocytes (Collins et al. 2012b; Schleicher et al. 2014). Transcripts for this and other complement-like molecules were first identified in ESTs from juvenile LOs (Castillo et al. 2009; McFall-Ngai et al. 2010). Immunocytochemical analysis detected the expression of C3 in epithelial cells of several tissues of juvenile squid, including the LO, gills, and skin (Castillo et al. 2009). Other complement homologs have also been identified in *E. scolopes* and its sister species *E. tasmanica* ([name], [year], unpublished data), including C1qDC proteins, C1qBP, and an MBL-like transcript (McFall-Ngai et al. 2010). Preliminary data also point toward the presence of several serine proteases with similarity to MASPs and Factor C ([name], [year], unpublished data), although biological activity for these and the other complement-like proteins remains to be confirmed. Furthermore, TEPs similar to C3 have been identified in *E. scolopes*. Initially thought to be a representative of the insect TEPs (iTEPs) subgroup, Es-CD109 was found to be expressed in several squid tissues, and its transcript was downregulated in the LO of juveniles harboring *V. fischeri* (Collins et al. 2012b; Yazzie et al. 2015). This suggested that, similarly to C3, this microbial sensor is modulated in order to avoid the removal of symbiont cells (Collins et al. 2012b; Yazzie et al. 2015).

Soluble Effector Molecules

One of the first immune-related molecules identified in *E. scolopes* was a halide peroxidase (Tomarev et al. 1993). This enzyme, localized to vesicles in the epithelial cells, was secreted on the ciliated appendages of symbiotic juveniles, possibly as an antimicrobial factor (Weis et al. 1996). Transcripts of enzymes such as chitinase and lysozyme have also been described as upregulated in the first hours of exposure to *V. fischeri*, suggesting a possible involvement in the symbiont selection process (Kremer et al. 2013). The finding of NOS in the squid LO represented another possible antimicrobial source (Davidson et al. 2004). Immunocytochemical studies found NOS and NO in vesicles localized to the mucus on ciliated epithelial cells, where the bacteria aggregate and symbiont selection starts. In addition, NOS was expressed in the crypt ducts and antechambers (Davidson et al. 2004). Furthermore, it was shown that the presence of the symbiont or its products (LPS and TCT) downregulated the expression of NOS and the production of NO (Davidson et al. 2004; Altura et al. 2011). The authors proposed that in this case, the attenuation of NO production was a response by the host, enacted to modify the crypt environment to ease colonization upon symbiont recognition (Altura et al. 2011).

Although hemocyanin is mainly expressed in gills and the branchial heart, it was also detected in the symbiotic LO crypts, where it was suggested to release oxygen,

2851 thereby promoting bacterial growth and bioluminescence (Kremer et al. 2014).
2852 Moreover, the detection of a hemocyanin isomer in the mucus secretions of the
2853 juvenile LO suggests that this molecule may have a dual role and serve in the sym-
2854 biont selection process as an antimicrobial agent against nonsymbiotic marine bac-
2855 teria (Kremer et al. 2014). An additional antimicrobial and bacteriostatic molecule
2856 recently reported in *E. scolopes* is galaxin, one of the most highly upregulated tran-
2857 scripts in colonized LOs (Chun et al. 2008; Wier et al. 2010), whose encoded pro-
2858 tein is localized to the epithelial cells and mucus secretions of the LO (Heath-Heckman
2859 et al. 2014). In vitro assays showed that a peptide fragment of galaxin had inhibitory
2860 effects mainly against Gram-positive bacteria, although the growth of *V. fischeri*
2861 was also affected (Heath-Heckman et al. 2014). As mentioned earlier, the sensor
2862 molecule PGRP2, which binds and degrades bacterial peptidoglycan, is localized to
2863 epithelial surfaces exposed to the environment and secreted into the LO mucus, sug-
2864 gesting a role during the initial stages of colonization and selection of the symbiont
2865 (Troll et al. 2010). This protein is also detected in the crypt lumen, suggesting that
2866 it also assists in modulating host–bacteria interactions once the symbiosis is estab-
2867 lished (Troll et al. 2010). Another soluble protein with antimicrobial properties
2868 found in this squid species is alkaline phosphatase (ALP) (Rader et al. 2012), whose
2869 enzymatic activity was upregulated in symbiotic hosts possibly in response to bacte-
2870 rial MAMPs. Indeed, the addition bacterial lipid A and TCT induced the enzymatic
2871 activity, while the addition of an inhibitor reduced bacterial colonization by more
2872 than 80%. Overall, it was suggested that *esap1* has a supporting role in the coloniza-
2873 tion and maintenance of symbiosis (Rader et al. 2012).

2874 Signaling Molecules

2875 Following the preliminary annotation of the LO-EST database, several molecules
2876 pertaining to the canonical TLR signaling (see section “[Canonical TLR Signaling](#)”)
2877 were identified (Goodson et al. 2005). In a related study, three p-63-like (a member
2878 of the p-53 family of tumor suppressor proteins) transcripts were identified and
2879 localized to the nuclei of LO cells in symbiotic animals, suggesting a role in the
2880 apoptosis of appendages (Goodson et al. 2006).

2881 This is a topic that warrants further study, as the capacity of the host to rec-
2882 ognize the correct bacterial symbiont from the multitude of bacterial cells in the
2883 water may reside in the signaling cascades triggered by *V. fischeri*. One interesting
2884 aspect that has been learned since the early studies of this symbiosis is that at first
2885 glance, *V. fischeri* bacteria do not seem to contain any evident “symbiont marker”
2886 that could help the host to discern the symbionts from other bacteria. Surprisingly,
2887 the same molecules present in nonsymbiotic bacteria, including pathogens, are
2888 used to communicate with the animal host. These MAMPs, such as LPS and PGN,
2889 should be readily recognized by the innate immune system as foreign and usu-
2890 ally elicit a response resulting in microbial removal (see section “[Phagocytosis](#)”).
2891 Similarly, the host interacts with the symbionts using PRRs and signaling pathways
2892 known to be usually activated by pathogens. Nonetheless, there is still the potential
2893 of discovering novel markers on the symbionts and receptors on the host, especially
2894 considering the scarce genomic resources currently available and the unknown

function of most cephalopod genes (see section “A Short Journey in the ‘Immune System’ of Cephalopods”). The described studies suggest that attention needs to be paid to the context, timing, and very possibly the effector mechanisms elicited in response to the bacterial signals that can make the difference between removal and accommodation.

Accessory Nidamental Gland

E. scolopes is also used to study another very interesting case of symbiosis, in this case involving a consortium of symbionts that may be acquired in different ways. This particular interaction occurs in the accessory nidamental gland (ANG) (Collins et al. 2012a). The ANG is part of the reproductive organs in female squid. This structure is formed by a series of epithelial tubules containing a mixture of bacterial species dominated by Rhodobacteriaceae (Barbieri et al. 2001; Collins and Nyholm 2011; Collins et al. 2012a, 2015). It is thought that some of the components of the ANG bacterial community are added to the jelly coat of eggs during their formation, and that the function of these microorganisms is to protect the developing embryos from environmental infections (Barbieri et al. 1997; Collins et al. 2012a, 2015). In a recent publication, Gromek and colleagues (2016) isolated one of the ANG bacteria (*Leisingera* sp.) from the jelly coat of *E. scolopes* eggs, and in in vitro studies demonstrated that it had antimicrobial activity, producing a pigment that selectively inhibited the growth of several marine bacteria, including *Vibrio* species.

Altogether, the knowledge obtained from the study of these two types of symbiosis has the potential to provide an improved understanding of the complex bacterial associations between animals and microbes. In particular, this might bring new elements to interpret the mechanisms of regulation of bacterial symbiosis in various organs, such as the digestive, respiratory, and urogenital tracts of mammals, further serving as a productive research field for deciphering the multifaceted roles of the immune system in metazoans, which are still not well understood.

Conclusions

The application of -omic tools to the study of bivalve and cephalopod immunology has recently led to exciting discoveries about the extent of the diversity of immune genes in these groups of diverse species. Comparative functional studies using natural and selectively bred disease-resistant strains of bivalves, and in-depth analysis of the powerful model system of the bobtail squid–*Vibrio* symbiosis, as well as the application of gene-editing technologies, have the potential to provide exciting insights into the functional relevance of immune gene family expansion in molluscs and the potential role of this diversity in the specificity and plasticity of immune responses. Other areas of molluscan immunity that have not been understudied until now, because of the lack of tools and resources, include the elucidation of the process of hematopoiesis, the molecular characterization of hemocyte subpopulations, and a thorough characterization of mechanisms underlying maternal immunity and immune priming.

2936 Molluscan immunobiology is gaining renewed importance from the growing
2937 challenges posed by human activities, which have a significant impact in particular
2938 on anthropized coastal regions (for a detailed discussion, see Chap. 12, section
2939 “Challenges for Molluscs in the Anthropocene Epoch”). This, together with the cur-
2940 rent trends of global climate change, is currently leading significant shifts in the
2941 structure of benthic communities due to the introduction of alien species, more
2942 resistant to the presence of pollutants and therefore outcompeting native species.
2943 Continuous research will be certainly needed to improve our knowledge of the
2944 immune system of molluscs, both to preserve endangered endemic populations and
2945 to face the challenges posed by emerging diseases targeting commercially and eco-
2946 logically important species (see Chap. 12, section “Molluscan Conservation
2947 Immunology” for a detailed discussion on molluscan conservation immunology).

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Author Queries

Chapter No.: 11 0003596912

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AU15	By 'would allow, in line of principle, additional physiological functions' do you mean 'would, in principle, allow additional physiological functions'?	
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