

Parasitism in species of *Bathymodiolus* (Bivalvia: Mytilidae) mussels from deep-sea seep and hydrothermal vents

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ABSTRACT: Bivalve species, especially mussels, are biomass dominants in many deep-sea chemosynthetic ecosystems. As in shallow-water environments, parasites are likely to be important factors in the population dynamics of bivalve communities in chemosynthetic ecosystems, but there has been little study of parasitism in deep-sea seep or vent molluscs. In this study, parasite types, diversity, prevalence, infection density and non-infectious indicators of stress or disease as related to host age, reproductive condition, and endosymbiont density were assessed in mussels (*Bathymodiolus heckeriae*) from 2 seep sites and mussels (*B. puteoserpentis*) from 2 vent sites. We identified 10 microbial or parasitic agents in histological sections. Parasite types included 3 viral-like gut inclusions, 2 rickettsia-like gill inclusions, a rickettsia-like mantle inclusion, a bacterial gill-rosette, a chlamydia-like gut inclusion, gill-dwelling ciliates, and an unidentified inclusion in gut tissues. Parasite species richness was greater in seep mussels than in vent mussels, with the seep mussels possessing 9 types of parasites compared to 2 in the vent mussels. One of the viral-like inclusions infecting the seep mussel *B. heckeriae* was pathogenic, causing lysis of the digestive tubules. The prevalence and intensity of infection by this pathogen were greater in hosts with shell lengths less than 100 mm. Mussels from all 4 sites also exhibited intense infiltration of tissues and blood spaces by enlarged hemocytes. Hemocytic infiltration (hemocytosis) showed variable degrees of severity that were not associated with other host factors examined.

KEY WORDS: Chemosynthetic ecosystems · Mollusc · Virus · Pathogenicity · Community ecology · Parasite ecology · Endosymbiotic bacteria · Commensal polychaetes

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INTRODUCTION

Parasitic infections can impair growth, reproduction, competitive ability, stress tolerance and survival of host species (Kautsky 1982, Price et al. 1986, Esch et al. 1990, Calvo-Ugarteburu & McQuaid 1998). In addition, parasites may regulate host population structure and influence coexisting species, such as the host's prey, predators or competitors (Anderson 1978, Anderson & May 1978, Price et al. 1986, Calvo-Ugarteburu & McQuaid 1998). Parasitism is thus an important factor to consider in studies of the ecology and dynamics of populations within any ecosystem (Esch et al. 1990, Powell et al. 1999, Montaudouin et al. 2000). The

paucity of studies of parasitism in organisms living in bathyal environments is a consequence of both the relative inaccessibility of the deep sea and the low biomass and abundance of the organisms living there. Chemosynthetic environments (hydrothermal vents and cold seeps) in the deep sea are celebrated for the high biomass and abundance of organisms adapted for life under extreme conditions, yet surprisingly little is known about parasitism in organisms living in these environments.

Hydrothermal vents are submarine hot springs located along mid-ocean ridges. They are dynamic, ephemeral ecosystems characterized by metal-rich fluids and temperatures elevated above those of the

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ambient deep sea (see Van Dover 2000). Deep-sea seeps are located primarily on continental margins where reduced compounds diffuse from the sea floor into the overlying water column. Seeps occur in a variety of settings (Sibuet & Olu 1998) and include petroleum seeps, which are common in the Gulf of Mexico (MacDonald et al. 1990), brine seeps found on the Florida Escarpment (Paull et al. 1984), and gas-hydrate seeps, including the methane-hydrate seep at Blake Ridge, off the coast of South Carolina (Van Dover et al. 2003). In contrast to hydrothermal vents, seeps are thought to be more stable, longer-lived ecosystems (Sibuet & Olu 1998). Ecological studies in these environments are still in their infancy and investigations of parasite burdens, pathology and disease in species living at seeps and vents are scarce.

Bathymodiolin mussels (Mytilidae) are one of the dominant megafaunal taxa found at seeps and vents. Bathymodiolin mussels at vents typically live in water between 5 and 15°C (Van Dover 2000); mussels at seeps are generally present at ambient seawater temperatures (~2°C; Sibuet & Olu 1998). Mussels play key roles in seep and vent ecosystems: they serve as physical habitats for other invertebrates, and they house chemoautotrophic, endosymbiotic bacteria within gill epithelial cells, called bacteriocytes (Cavanaugh 1983, Le Pennec & Hily 1984). The endosymbionts account for a large proportion of the primary production in seep and vent communities. The energy required for carbon dioxide fixation is obtained by oxidation of reduced compounds in the endosymbionts; fixed carbon is translocated to the host mussel (Cavanaugh 1983). Bathymodiolin mussels can obtain nutrition from single (e.g. thiotrophic bacteria) or dual (e.g. thiotrophic and methanotrophic bacteria) symbioses. Healthy mussels retain a functional gut and an ability to filter-feed on particulate organic matter (Le Pennec et al. 1990), but in the absence of reduced compounds, gill condition deteriorates (Raulfs et al. in press).

Although diseases of commercially valuable intertidal bivalves have been extensively studied, little is known about parasites in bivalves found at deep-sea seeps, and nothing is known of parasites in bivalves found at hydrothermal vents. Extreme conditions in chemosynthetic ecosystems include high concentrations of metals at vents, brines and hydrocarbons at some seeps, and sulfide-rich effluents or pore-waters. Organisms endemic to chemosynthetic environments have adapted to these environments and require what would normally be considered noxious conditions.

Powell et al. (1999) examined the relationship between elevated concentrations of polycyclic aromatic hydrocarbons (PAH) and parasite burdens in mussels *Bathymodiolus* sp., from 4 hydrocarbon seep sites in the Gulf of Mexico (depth 550 to 650 m). In coastal

mussels, PAHs disrupt membranes, damage lysozymes and inhibit hemocyte phagocytic ability (Pipe & Coles 1995, Fernley et al. 2000). Unlike shallow-water mussels, the mussels from hydrocarbon seeps were obligately associated with high PAH concentrations; 5 types of parasites were documented in the hydrocarbon-seep mussels, including extracellular gill ciliates, bacterial gill rosettes, *Bucephalus*-like trematodes, and chlamydia/rickettsia-like gill and gut inclusions (Powell et al. 1999). In comparison to mussels from intertidal sites, the mussels from hydrocarbon seeps had similar types of parasites, but were more heavily parasitized. Infections of *Bucephalus*-like trematodes caused severe castration of hydrocarbon-seep mussels, leaving 40% of the populations in the study reproductively compromised (Powell et al. 1999).

To explore further the relationship between parasite burdens in bathymodiolin mussels and chemosynthetic environments, we conducted a comparative study of parasite burdens in *Bathymodiolus heckeriae* from 2 deep-sea seeps and *B. puteoserpentis* from 2 deep-sea hydrothermal vents. Both species contain methanotrophic and thiotrophic endosymbiotic bacteria. Differences in types of parasites and abundance of parasites were documented between the 2 geologically and ecologically distinct habitat types. Prevalence, infection densities, and noninfectious indicators of stress or disease as they related to age, reproductive condition and an index of endosymbiont density were also assessed.

MATERIALS AND METHODS

Specimens of *Bathymodiolus heckeriae* were collected using the deep-sea submersible 'Alvin' from 2 seep sites: a cold-water, saline site at the Florida Escarpment (26° 2' N, 84° 55' W, 3300 m; sampled in October 2000) and a gas-hydrate site off the Carolina coast (Fig. 1; Blake Ridge 32° 31' N, 76° 12' W, 2155 m; sampled in September 2001). Species of *B. puteoserpentis* were collected from the Logatchev vent (15° 45' N, 44° 58' W, 3300 m, sampled in July 2001; Batuyev et al. 1994) and Snake Pit (23° 22' N, 44° 56' W, 3600 m, sampled in July 2001; Karson & Brown 1988).

Mussels were sorted by length, and 30 to 32 specimens were systematically selected to represent an even distribution across the range of sizes available. Shell lengths were recorded for each specimen. Within 3 h of collection, mussels were removed from the shell, tagged, fixed in Davidson's solution (Humason 1972) for 24 h, and stored in 70% ethanol. The number and lengths of commensal polychaetes *Branchipolynoe seepensis* in the mantle cavities of the mussels were determined.

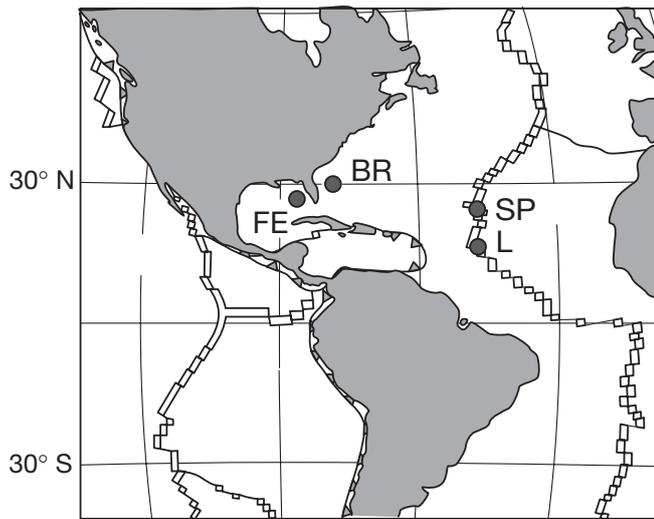


Fig. 1. Locations of Florida Escarpment (FE) and Blake Ridge (BR) seep sites and Logatchev (L) and Snake Pit (SP) vent sites

Wet weight was determined for whole, fixed mussels without the shell. An oblique, ~5 mm thick, transverse tissue slice, containing mantle, gonad, digestive tract and gill was dissected from each specimen. This tissue slice was dehydrated in a graded ethanol series, embedded in paraffin, and transverse sections were taken at 5 to 6 μm . Sections were stained with Gill's hematoxylin and eosin (H&E; Stevens 1990) or Ordway-Machiavello for rickettsia (Humason 1972) to identify parasite types and prevalence (no. of host individuals infected with any parasite; Margolis et al. 1982) using light microscopy. A Spot camera (Diagnostic Instruments) was used for capturing images for light microscopy. Contrasts were adjusted and photomosaics were generated using Adobe Photoshop (Adobe Systems).

Using an ocular grid, parasite densities (number of parasites per area standardized to 500 μm^2 of tissue; Margolis et al. 1982) were determined for infected individuals. We examined 5 grids per section for 3 sections that were separated by ~1 mm or more. Body burden (mean infection densities for all parasitic infections) of infected individuals was also determined. A quantitative measure of bacterial endosymbiont density was determined for *Bathymodiolus puteoserpentis* by measuring the area of the gill bacteriocytes occupied by the endosymbiotic bacteria to derive a ratio of the area occupied by the endosymbiotic bacteria to the area of the gill bacteriocytes. Cells used to measure bacterial endosymbiont density were located on sections of the transverse gill filaments in approximately the same region of each mussel. Host sex and gonad stage were measured. Semi-quantitative scales were assigned for intensity of hemocyte infiltration

Table 1. Index used to determine intensity of hemocyte infiltration in *Bathymodiolus* spp.

Intensity	Description
0	No areas of infiltration
1	Slight infiltration, small, focal, usually only around intestine
2	Several small foci of infiltration or a few larger areas; sometimes in more than 1 type of tissue
3	Disseminating foci ranging from small to large areas; infiltration extensive and usually in several or all types of tissue
4	A few blood vessels enlarged due to heavy infiltration of hemocytes
5	Most blood vessels enlarged due to massive infiltration

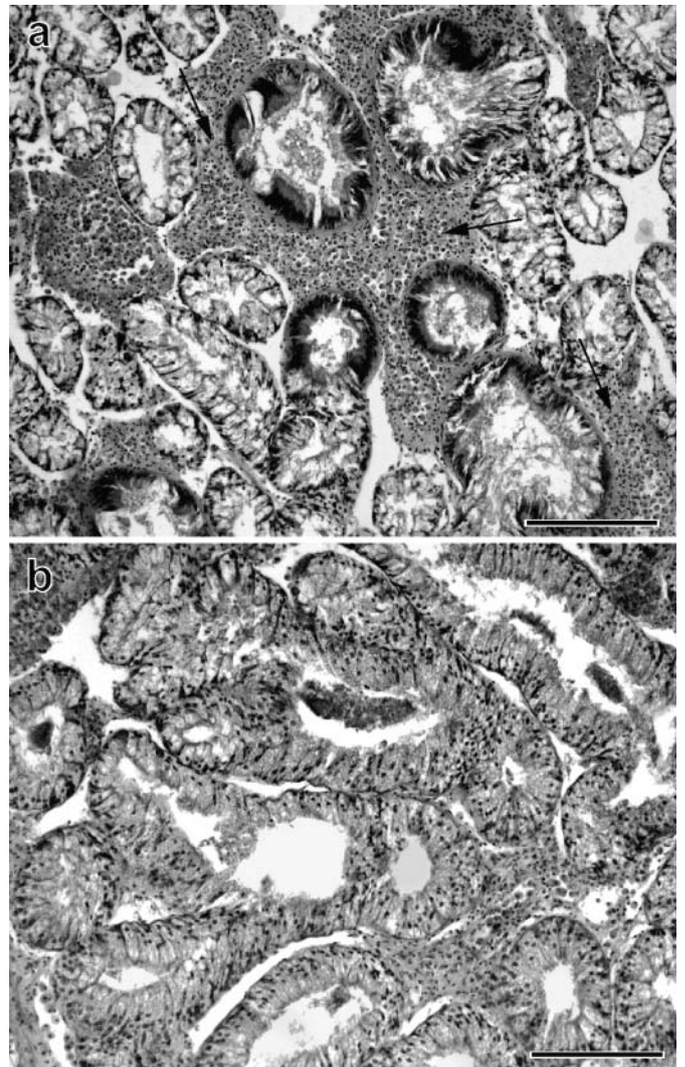


Fig. 2. *Bathymodiolus puteoserpentis*. Hemocyte infiltration. (a) Heavy infiltration of hemocytes (arrows) around digestive diverticula and intestine; (b) healthy visceral mass tissue without hemocyte infiltration. Scale bars = 200 μm

Table 2. Index used to determine gonad development in *Bathymodiolus* spp.

Gonad stage	Description
1	Undifferentiated; few areas in visceral mass with gonad
2	Early stages of developing gametes; several areas of visceral mass with small regions of gonadal development
3	Approximately equal proportions of developing and mature gametes; large areas of visceral mass taken up by gonads
4	Early stages of gametogenesis rare; follicles contain mature gametes; visceral mass densely packed with gonad

per cross-section (Table 1, Fig. 2) and density and stage of development of gonads (Table 2).

Statistical tests were performed using MINITAB software, Version 13.20, 2000. Comparisons of means among sites were determined using Kruskal-Wallis 1-way analysis of variance. If significant results ($p < 0.05$) were obtained, ANOVA and Tukey's pairwise comparisons were used to evaluate significant differences among sample pairs. Multi-dimensional scaling (MDS; PRIMER v5; Clarke & Gorley 2001) was used to examine similarities among individuals based on the parasite type–density matrix, using Bray-Curtis coefficients calculated from non-transformed data. The 16 individuals not infected by any parasite were excluded from the MDS analysis. MDS plots position individuals on a unitless 2-dimensional map; closely spaced points represent the most similar individuals in a multivariate space based on parasite types and density. Analysis of similarity (ANOSIM subroutine of PRIMER v5) was used to determine significant differences between groups of individuals evident in MDS plots. ANOSIM provides R statistics: when $R > 0.75$, groups are well-separated; when $0.75 > R > 0.5$, groups are overlapping but clearly different; when $R < 0.25$, groups are not separable (Clarke & Gorley 2001). Factors contributing to these differences of parasite infection densities were determined from similarity

percentages (SIMPER subroutine in PRIMER v5). The Shannon-Wiener diversity index ($H'_{\log(e)}$) was calculated using the DIVERSE subroutine in PRIMER v5. This diversity index takes into account both species richness (the number of parasite types) and evenness (the distribution of individuals among species; Hayek & Buzas 1997).

RESULTS

Parasite fauna

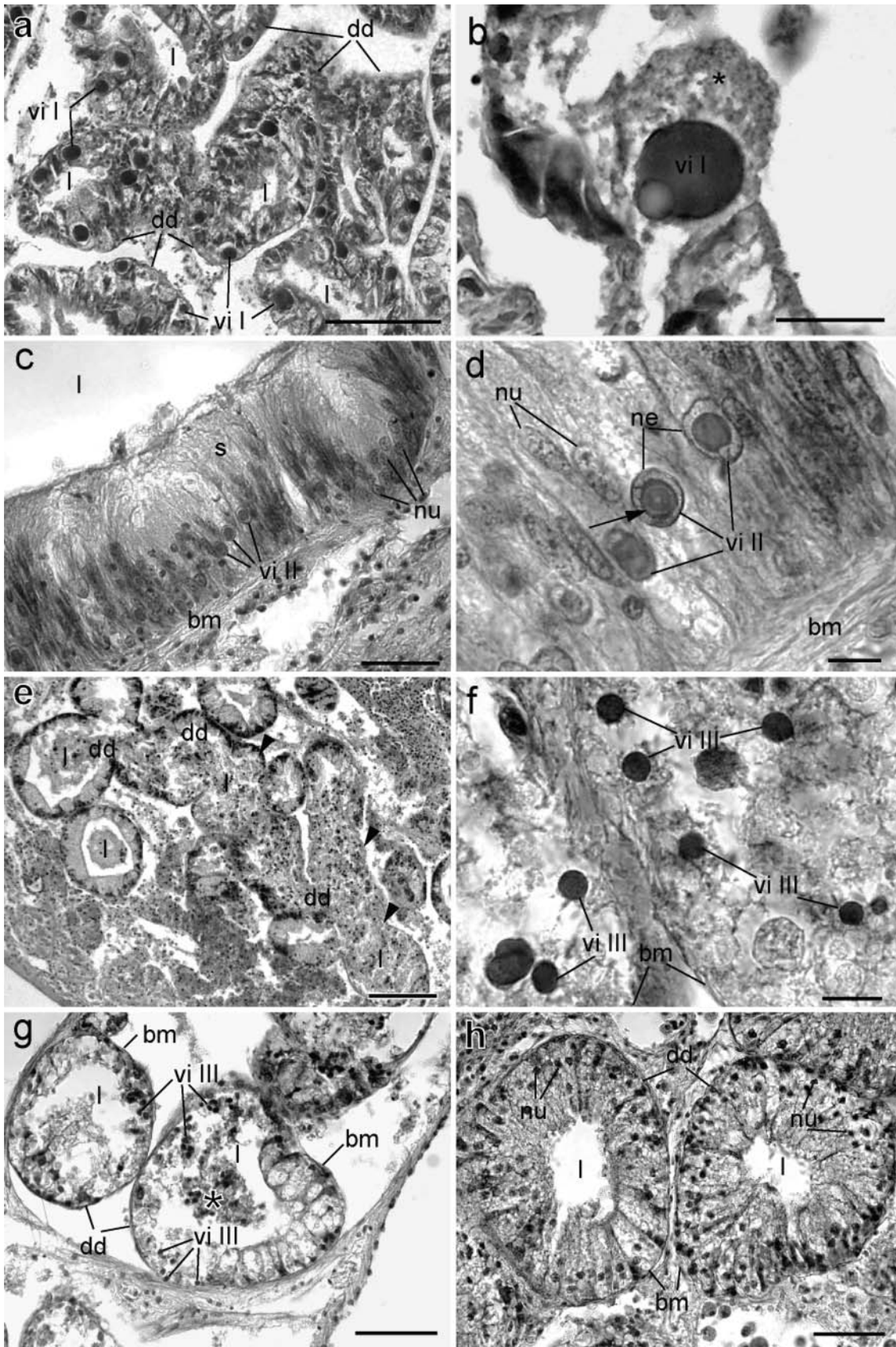
We identified 10 types of parasites (9 in *Bathymodiolus heckerae*, 2 in *B. puteoserpentis*) in histological sections: 3 viral-like inclusions in the gut, 2 rickettsia-like inclusions in the gill, rickettsia-like inclusions in the mantle, bacterial rosettes in the gill, chlamydia-like inclusions in the gut, gill ciliates and an unidentified inclusion in the gut.

One of the viral-like inclusions in the gut, referred to hereafter as Viral Inclusion I, was observed only in the vent mussel *Bathymodiolus puteoserpentis*, where it occurred in the digestive diverticula (Fig. 3a). These eosinophilic inclusions were generally spherical in shape with an average diameter of 20 μm ($n = 20$; Fig. 3b). Occasionally, the cell contents were peripherally located in the host cell or the cell was hypertrophied (Fig. 3b).

In the gut, 2 viral-like inclusions were associated with the seep mussel *Bathymodiolus heckerae*. Of these, one, hereafter referred to as Viral Inclusion II, occurred in the nuclei of stomach and intestinal epithelia of mussels from the Florida Escarpment (Fig. 3c). The inclusion had a waxy, eosinophilic appearance and was oval or spherical in shape with an average diameter of 11 μm ($n = 20$). The inclusion was surrounded by what we infer to be the hypertrophied nuclear envelope of the host cell. Lighter staining rings were occasionally visible within the inclusion (Fig. 3d). No obvious tissue pathology was associated with this infection.

A third viral-like inclusion in the gut (Viral Inclusion III), infected the epithelial nuclei of the digestive diver-

Fig. 3. *Bathymodiolus* spp. containing viral-like inclusions. (a) Individual heavily infected with Viral Inclusion I in digestive diverticula; scale bar = 200 μm . (b) Viral Inclusion I in digestive diverticula causing hypertrophy and necrosis of the host cell (*); scale bar = 20 μm . (c) Viral Inclusion II infecting stomach epithelial cells; scale bar = 50 μm . (d) Detail of Viral Inclusion II showing hypertrophied nuclear envelope (arrow); scale bar = 10 μm . (e) Viral Inclusion III infecting digestive diverticula and intestinal tissue; viral-like inclusion bodies visible in lumen and tissues with necrotic areas of digestive diverticula (arrowheads); scale bar = 200 μm . (f) Viral Inclusion III within 2 tubules of the digestive diverticula; scale bar = 10 μm . (g) Digestive diverticula infected with Viral Inclusion III with necrotic tissue caused by the virus; mass of viral-like inclusions and necrotic cells are visible in 1 tubule (asterisk); scale bar = 60 μm . (h) Uninfected digestive diverticula for comparison; scale bar = 50 μm . bm: basement membrane; dd: digestive diverticula; l: lumen; ne: nuclear envelope; nu: nucleus; s: stomach; vi I: Viral Inclusion I; vi II: Viral Inclusion II; vi III: Viral Inclusion III



ticula and intestines of *Bathymodiolus heckeræ* from both seep sites (Fig. 3e). Infections were characterized by small (4.8 µm diameter; n = 25) eosinophilic spherical bodies within host cells, free in the lumen of the digestive tract (Fig. 3e,f), or occasionally in the connective tissue of the visceral mass. The spherical bodies showed no internal structure at the level of light microscopy. Severe tissue pathology was often associated with infection by Viral Inclusion III (Fig. 3e,f,g). In moderately or heavily infected mussels, infected cells had either lysed or had sloughed into the lumen of the digestive tubules (Fig. 3g). Widespread necrosis of the digestive tissue was evident with large regions of the digestive tissue destroyed, leaving a mass of viral inclusions among the remnants of the cells (Fig. 3e,f). In adjacent healthy mussel tissues, nuclei were basally located and individual cells were easily distinguishable (Fig. 3h).

One of the rickettsia-like inclusions in the gill, referred to as Gill Rickettsia I, was found in both mussel species from all 4 sites. These basophilic, intracytoplasmic inclusions were spherical, with an average diameter of 20 µm (n = 20) and could be found in all cell types along the entire length of the gill filament (Fig. 4a,b). Many inclusions had a filamentous internal structure (Fig. 4b) while other inclusions appeared smooth.

A second type of rickettsia-like inclusion in the gill, referred to as Gill Rickettsia II was found only in *Bathymodiolus heckeræ* from the Florida Escarpment. The intracytoplasmic inclusions were basophilic and were found along the entire length of the gill filament within the host bacteriocytes (Fig. 4c). These inclusions were morphologically different from Gill Rickettsia I, being larger (average diameter 37 µm; n = 20) and more irregular in shape (Fig. 4c,d). These inclusions also had a filamentous internal structure (Fig. 4d). Both types of Gill Rickettsia were capable of displacing the endosymbiotic bacteria to the periphery of the host cell.

The third type of rickettsia-like inclusion, referred to as Mantle Rickettsia, was found in the cytoplasm of mantle epithelial cells in *Bathymodiolus heckeræ*. These inclusions were morphologically similar to Gill Rickettsia I, but due to their different location within

the host, they were considered a different parasite. No obvious tissue pathology was observed for this parasite.

Bacterial gill rosettes similar to those described by Powell et al. (1999) were observed in bacteriocytes. They were associated with the host bacteriocytes of *Bathymodiolus heckeræ*, averaged 9 µm (n = 15) in diameter, and were located within or penetrating the membrane of the host gill cell (Fig. 4e). The rosette structures were composed of groups of 30 or more basophilic, spherical bodies, each with an average diameter of 2.8 µm (n = 25; Fig. 4f). No tissue damage was observed except where rosettes penetrated the gill cell membranes.

Chlamydia-like intracytoplasmic inclusions occurred in epithelial cells of the stomach, intestine, and digestive diverticula of *Bathymodiolus heckeræ*. These parasites were basophilic, finely granular, and irregular in shape and size (average diameter 50 µm; n = 20). The finely granular inclusions appeared either tightly compacted (Fig. 4g) or diffuse (Fig. 4h). Host cells were hypertrophied or lysed by the inclusion (Fig. 4g,h).

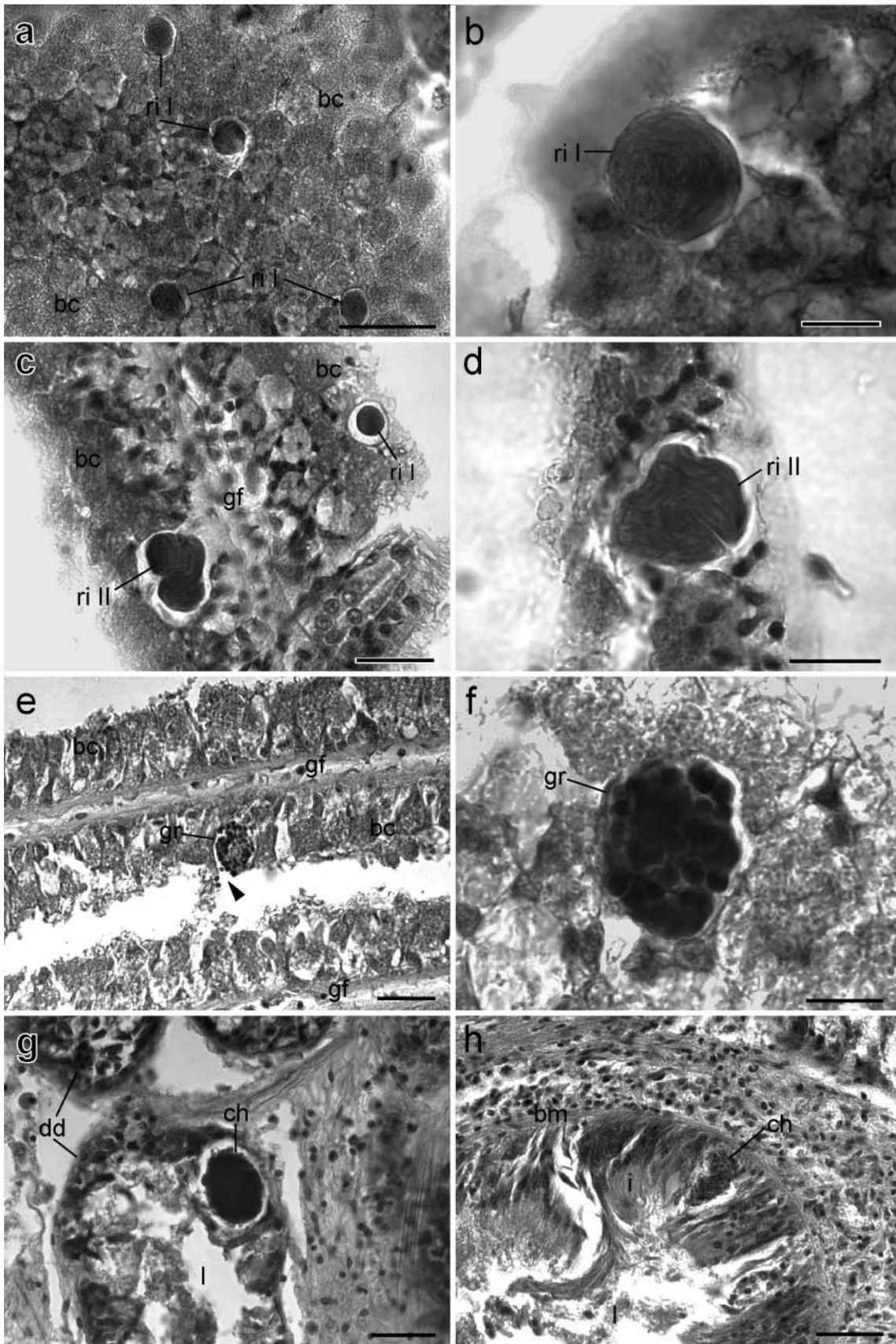
Ciliates were found among the gills of *Bathymodiolus heckeræ* (Fig. 5a). They had an average length of 30 µm (n = 15) and were associated with, but did not appear to be attached to, the gill filaments of the host mussel (Fig. 5a). No obvious tissue pathology was observed.

An unidentified parasite occurred only in *Bathymodiolus heckeræ*. It was characterized by a slightly refringent, intracytoplasmic, basophilic, spherical inclusion with an average diameter of 8 µm (n = 10; Fig. 5b). These inclusions were located in the epithelial cells of the intestine and digestive diverticula of the host. With the exception of a slight hypertrophy of the host cell, no obvious tissue pathology was observed.

Polychaetes

Commensal polynoid polychaetes *Branchipolynoe seepensis* were retained between the gill filaments of 2 vent mussels and subsequently sectioned (Fig. 6a). Displacement of gill filaments and some physical trauma of the filaments resulting in necrosis or poor

Fig. 4. *Bathymodiolus* spp. with bacteria-like infections. (a) Moderate infection with Gill Rickettsia I; rickettsial inclusions are located in gill bacteriocytes; scale bar = 50 µm. (b) Gill Rickettsia I with filamentous internal structure; scale bar = 10 µm. (c) Gill filament infected with both Gill Rickettsia I and Gill Rickettsia II; note difference in size and shape of the 2 inclusions; scale bar = 30 µm. (d) Gill Rickettsia II with filamentous internal structure and irregular shape; scale bar = 20 µm. (e) Gill rosettes located among bacteriocytes of host gill filament; gill rosette appears to be breaking through the host gill cell (arrowhead); scale bar = 30 µm. (f) Gill rosette composed of several spherical bodies with average diameter of 2.8 µm; scale bar = 10 µm. (g) Tightly compacted chlamydia-like inclusion in the digestive diverticula; scale bar = 40 µm; (h) More diffuse chlamydia-like inclusion in intestine; scale bar = 40 µm. bc: bacteriocytes; bm: basement membrane; ch: chlamydia-like inclusion; dd: digestive diverticula; gf: gill filament; gr: gill rosette; i: intestine; l: lumen; ri I: Gill Rickettsia I; ri II: Gill Rickettsia II



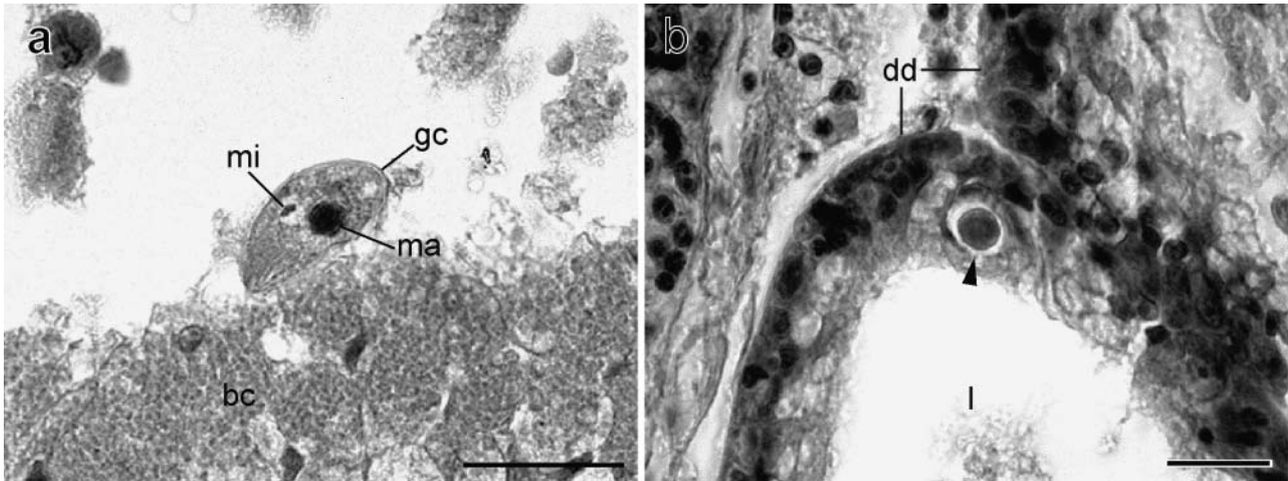


Fig. 5. *Bathymodiolus* spp. containing gill ciliates and an unidentified parasite. (a) Gill ciliate showing micro- and macro-nucleus; scale bar = 30 μ m. (b) Unidentified intracytoplasmic inclusion (arrowhead) in a digestive diverticulum; scale bar = 20 μ m. bc: bacteriocytes; dd: digestive diverticula; gc: gill ciliate; l: lumen; ma: macro-nucleus; mi: micro-nucleus

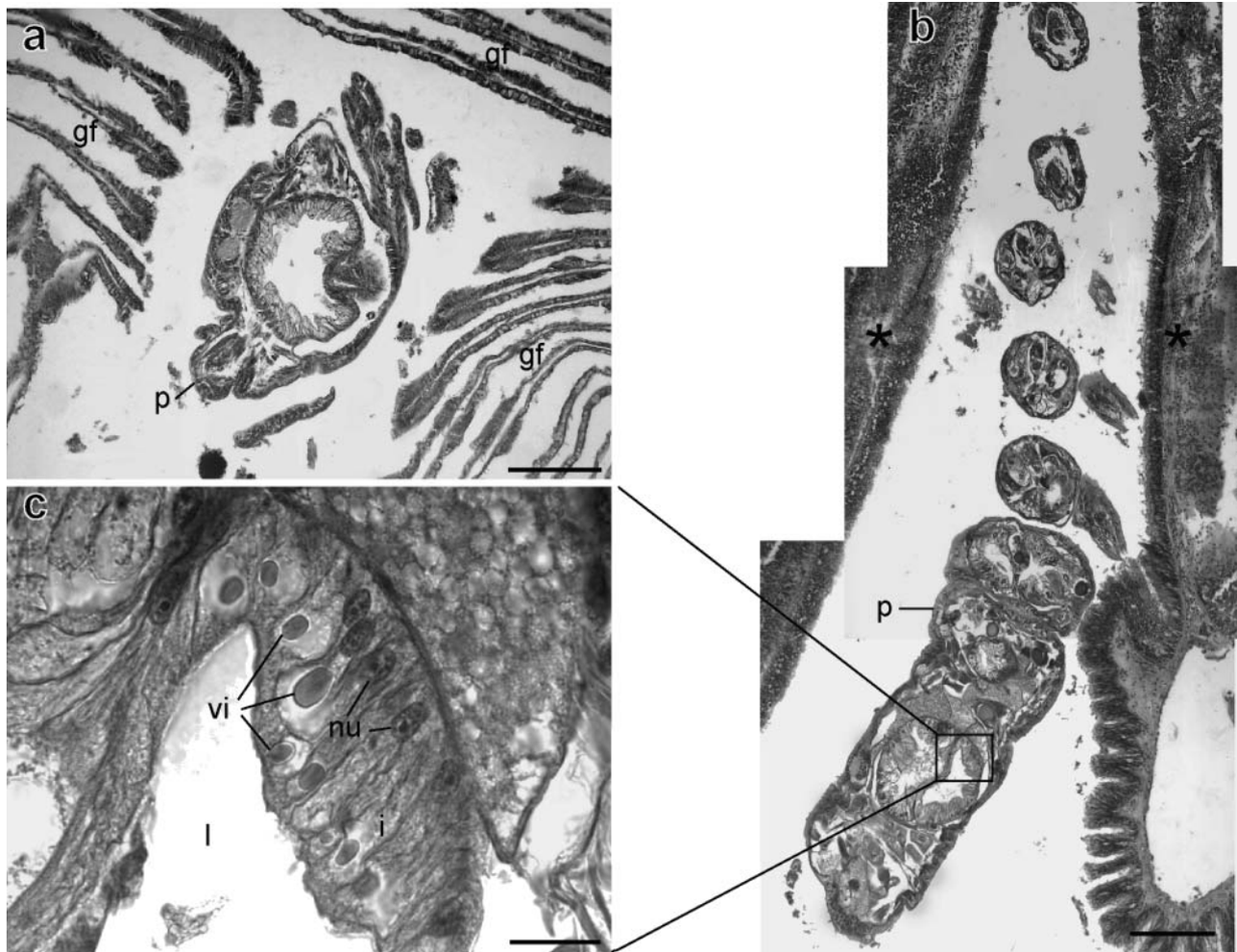


Fig. 6. *Bathymodiolus* spp. containing commensal polychaetes. (a) Polynoid polychaete *Branchipolynoe seepensis* between gill filaments of vent mussel *B. puteoserpentis*; scale bar = 200 μ m. (b) Nautiliniellid polychaete between demibranchs (*) of *B. heckerae* from Florida Escarpment; scale bar = 200 μ m. (c) Viral-like inclusions in intestine of the nautiliniellid; scale bar = 20 μ m. gf: gill filaments; i: intestine; l: lumen; nu: nucleus; p: polychaete; vi: viral-like inclusion

condition were attributed to the polychaetes. In *Bathymodiolus puteoserpentis*, the commensal polychaete had a prevalence of 30 to 40%. An even greater prevalence of the commensal polychaete was observed in *B. heckeræ* from the Florida Escarpment seep site (>60%). *Branchiopolynoe seepensis* was not observed in *B. heckeræ* from the Blake Ridge seep.

In 3 *Bathymodiolus heckeræ* from the Florida Escarpment, 4 nautiliniellid polychaetes (possibly *Laubierus mucronatus* based on characteristics of the setae) were observed in section (Fig. 6b). The polychaetes were located between the ascending and descending filaments of the demibranchs. Little evidence of damage on the surrounding filaments was observed, although cells in the adjacent gill tissue were detached from the filament or the tissue was in poor condition; 3 of the polychaetes were infected with viral-like inclusions in the digestive tract (Fig. 6c).

Parasite diversity and abundance

Parasite diversity (H') was highest ($H' = 0.92$) in *Bathymodiolus heckeræ* from the Florida Escarpment (Table 3). Parasite diversities at the other 3 sites were

considerably lower, with H' values of <0.20. Species richness (number of parasite types) was greater in *B. heckeræ* than in *B. puteoserpentis* (Table 3, Fig. 7a). Mussels from the Florida Escarpment were infected with 9 different types of parasites and *B. heckeræ* from the Blake Ridge were infected with a subset of at least 6 of these. Viral Inclusion I was unique to *B. puteoserpentis* at the vents. Gill Rickettsia I was observed in both mussel species at all 4 sites.

The number of parasite types in infected individuals (Fig. 7b) was species- and site-dependent. *Bathymodiolus heckeræ* from Florida Escarpment seeps had ~4 times more parasites per individual on average than did *B. heckeræ* from Blake Ridge (Kruskal-Wallis: $p < 0.001$). Infected individuals of *B. puteoserpentis* from the Logatchev vent had a slightly but significantly greater number of parasites per individuals on average (Fig. 7b) than did infected individuals from Snake Pit (1.1; Kruskal-Wallis: $p = 0.017$).

More than 70% of the mussels sampled from all 4 sites were parasitized by 1 or more types of parasites. At the Florida Escarpment site, 100% of the mussels were infected with at least 1 type of parasite (Table 3, Fig. 8a). Despite large numbers of infected individuals, total body burdens were relatively low except in

Table 3. *Bathymodiolus* spp. Prevalence (% of individuals infected) and densities of infection per 500 μm^2 of tissue for parasitic agents except viral-like infections and data on polychaete infestations. Number of individuals sampled is shown in parentheses next to site names

Infection	<i>Bathymodiolus heckeræ</i>		<i>Bathymodiolus puteoserpentis</i>	
	Seep sites		Hydrothermal vent sites	
	Florida Escarpment (32)	Blake Ridge (32)	Logatchev (32)	Snake Pit (30)
Gill Rickettsia I				
Prevalence	94 %	13 %	87 %	20 %
Density (\pm SD)	0.3 (\pm 0.34)	0.1 (\pm 0.09)	1.1 (\pm 1.20)	0.06 (\pm 0.04)
Gill Rickettsia II				
Prevalence	69 %	0	0	0
Density (\pm SD)	0.3 (\pm 0.34)	0	0	0
Mantle Rickettsia				
Prevalence	3 %	3 %	0	0
Density (\pm SD)	0.03	0.7	0	0
Bacterial gill rosette				
Prevalence	34 %	0	0	0
Density (\pm SD)	0.2 (\pm 0.35)	0	0	0
Chlamydia-like inclusion				
Prevalence	21 %	3 %	0	0
Density (\pm SD)	0.3 (\pm 0.56)	0.25	0	0
Gill ciliate				
Prevalence	40 %	3 %	0	0
Density (\pm SD)	0.04 (\pm 0.05)	0.02	0	0
Unidentified gut inclusion				
Prevalence	50 %	12 %	0	0
Density (\pm SD)	0.08 (\pm 0.05)	0.03 (\pm 0.02)	0	0
<i>Branchiopolynoe seepensis</i> polychaetes				
Prevalence	62 %	0	31 %	40 %
# polychaetes per individual (\pm SD)	1.5 (\pm 1.26)	0	1.1 (\pm 0.55)	1.7 (\pm 1.24)

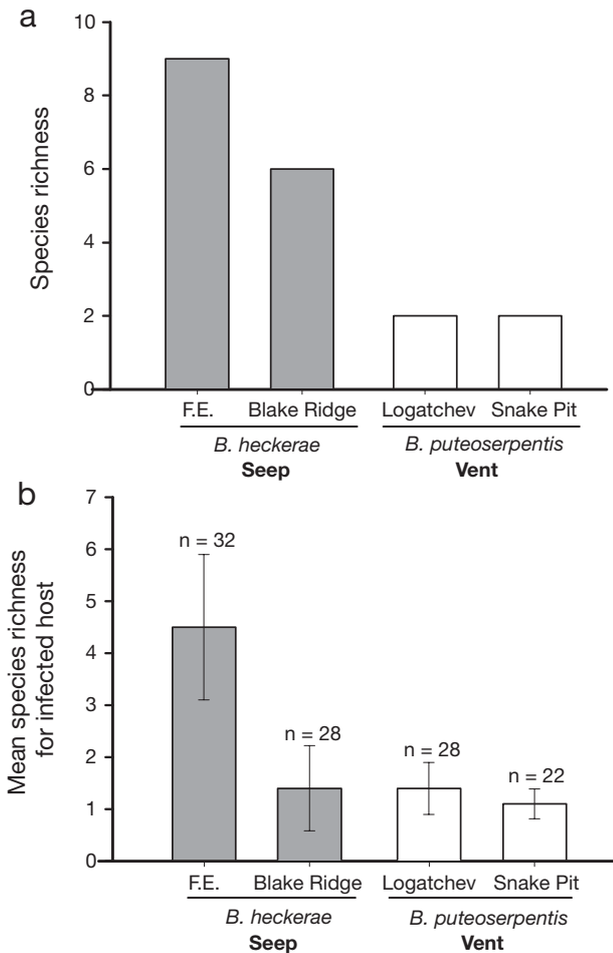


Fig. 7. *Bathymodiolus heckeriae* and *B. puteoserpentis* (a) Parasite species richness (no. of parasite species) in mussels; (b) mean species richness for infected mussels as a function of site. F.E.: Florida Escarpment; n: number of individuals

mussels from the Blake Ridge seep (Tukey's, $p < 0.001$; Fig. 8b). Individuals from the 4 sites were mapped as significantly different groups on MDS plots based on the parasite type–density matrix (Fig. 9; ANOSIM, $R > 0.40$; excluding uninfected hosts). Differences in parasite burdens between *Bathymodiolus heckeriae* and *B. puteoserpentis* ($R \geq 0.8$) and between *B. heckeriae* from Florida Escarpment and Blake Ridge ($R = 0.43$) were attributable to infection by Viral Inclusion III, which accounted for an average of 68% of the differences (SIMPER). *B. heckeriae* from the Florida Escarpment were mapped into 2 significantly different groups ($R = 0.58$; ANOSIM; Fig. 9). Infection with Viral Inclusion III accounted for 67% of the difference between these groups (SIMPER). *B. heckeriae* individuals infected with Viral Inclusion III mapped closest to the *B. heckeriae* individuals from Blake Ridge that were also infected with the virus (Fig. 9).

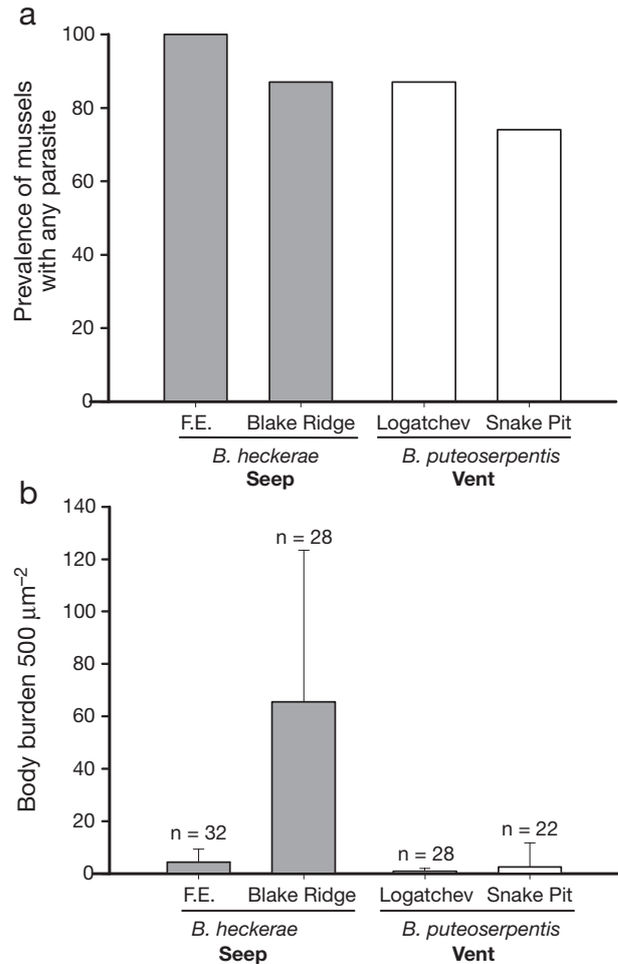


Fig. 8. *Bathymodiolus heckeriae* and *B. puteoserpentis*. (a) Prevalence of mussels infected with any parasite; (b) body burden (mean infection densities for all parasitic infections) of infected mussels ($500 \mu\text{m}^{-2}$ tissue). F.E.: Florida Escarpment; n: number of individuals

Viral-like gut inclusions occurred in mussels from all 4 sites, but *Bathymodiolus heckeriae* had a greater prevalence of viral inclusions than *B. puteoserpentis* (Fig. 10a). At Snake Pit, 60% of *B. puteoserpentis* were infected with Viral Inclusion I, but <30% of the mussels from the Logatchev site were infected with the same or similar Viral Inclusion I. More than 90% of the *B. heckeriae* from Florida Escarpment were infected with either Viral Inclusion II or Viral Inclusion III; of these, 53% were infected by both viruses. Almost 90% of *B. heckeriae* from Blake Ridge were infected with the pathogenic Viral Inclusion III. Infection density of Viral Inclusion III was significantly greater in *B. heckeriae* from Blake Ridge than from the Florida Escarpment (Tukey's: $p < 0.001$; Fig. 10b). Host size was an important factor in infection of *B. heckeriae*. Mean infection density for Viral Inclusion III was greater in individuals with shell lengths < 100 mm (Kruskal-

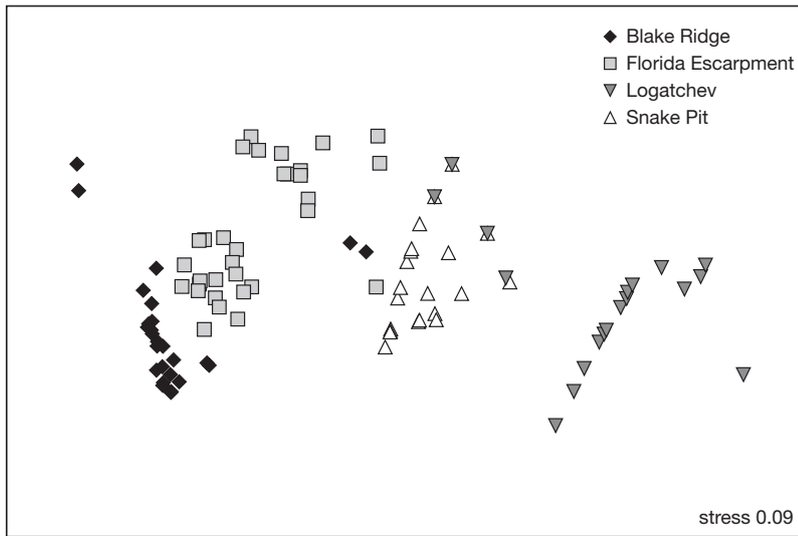


Fig. 9. *Bathymodiolus heckeriae* and *B. puteoserpentis*. Body burdens of all mussels infected with at least 1 parasite, analyzed by MDS. Samples were placed on a unitless map and distances between samples represent dissimilarity in parasite types and abundance infecting individual hosts (*B. heckeriae* = Florida Escarpment and Blake Ridge; *B. puteoserpentis* = Logatchev and Snake Pit). Note separation between sites

Wallis with outlier 0.005; Kruskal-Wallis without outlier: $p < 0.007$; Florida Escarpment, Fig. 11a; Blake Ridge, Fig. 11b). Mussels with shell lengths >100 mm were not infected or only slightly infected with Viral Inclusion III (Fig. 11).

Gill Rickettsia I was present in both species of mussels, but its prevalence varied between habitats. *Bathymodiolus puteoserpentis* from Logatchev vents had a significantly higher density of Gill Rickettsia I than *B. puteoserpentis* or *B. heckeriae* from the other sites (Tukey's: $p < 0.02$).

Non-infectious diseases

Heavy hemocyte infiltration (Fig. 2a) occurred in mussels from all 4 sites (Table 4). Hemocyte infiltration was observed in the majority of mussels

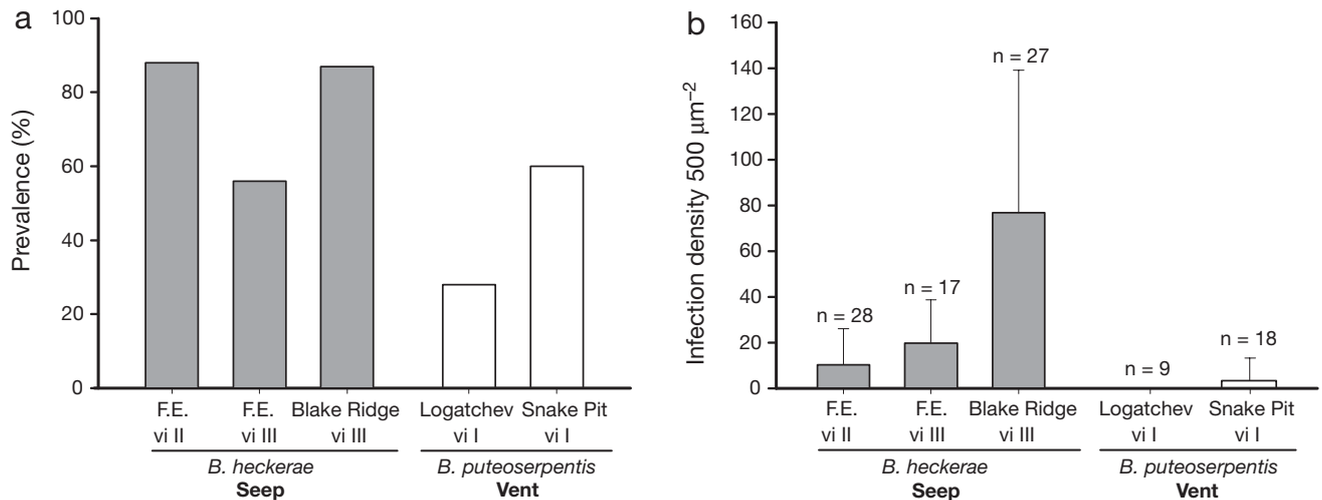


Fig. 10. *Bathymodiolus heckeriae* and *B. puteoserpentis*. (a) Prevalence of viral-like gut inclusions in mussels (b) density of infection of viral-like gut inclusions ($500 \mu\text{m}^{-2}$). F.E.: Florida Escarpment; n: number of individuals; vi: viral inclusion

Table 4. *Bathymodiolus* spp. Non-infectious indices of disease and index of gonad development. Prevalence of individuals with hemocyte infiltration and intensity of infiltration ($500 \mu\text{m}^{-2}$ tissue). Number of individuals sampled is shown in parentheses next to site names

Parameter	<i>Bathymodiolus heckeriae</i> Seep sites		<i>Bathymodiolus puteoserpentis</i> Hydrothermal vent sites	
	Florida Escarpment (32)	Blake Ridge (32)	Logatchev (32)	Snake Pit (30)
Prevalence of hemocyte infiltration	91%	97%	63%	90%
Mean hemocyte infiltration (\pm SD)	2.2 (\pm 1.4)	2.9 (\pm 2.0)	2.3 (\pm 1.5)	2.8 (\pm 1.6)
Mean gonad development (\pm SD)	2.3 (\pm 0.97)	1.3 (\pm 0.70)	3 (\pm 1.13)	2.5 (\pm 1.10)

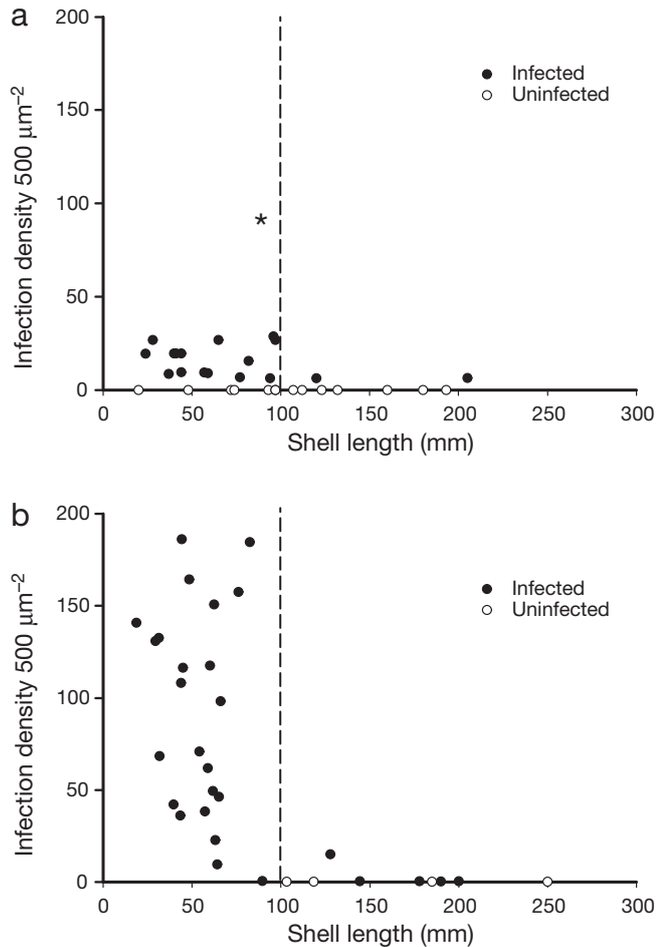


Fig. 11. *Bathymodiolus heckeræ*. Infection density of Viral Inclusion III infections ($500 \mu\text{m}^{-2}$ tissue) in mussels from (a) Florida Escarpment and (b) Blake Ridge in relation to mussel shell length (mm). *: outlier; dashed line: 100 mm shell length

(107 of 126) and, in 42 of these mussels, massive hemocyte infiltration resulted in abnormal expansion of blood vessels (Fig. 12). Most hemocytes appeared to be enlarged, with the nuclei displaced to the side of the cells (Fig. 12b). A positive relationship (Pearson's: $p < 0.002$) was observed between hemocyte infiltration and host length in mussels from all 4 sites.

Index of gonad development

Gonads of *Bathymodiolus heckeræ* from Florida Escarpment were more developed than those of *B. heckeræ* from the Blake Ridge (Kruskal-Wallis: $p < 0.001$). In *B. heckeræ* from the Florida Escarpment, individuals were more evenly distributed across the

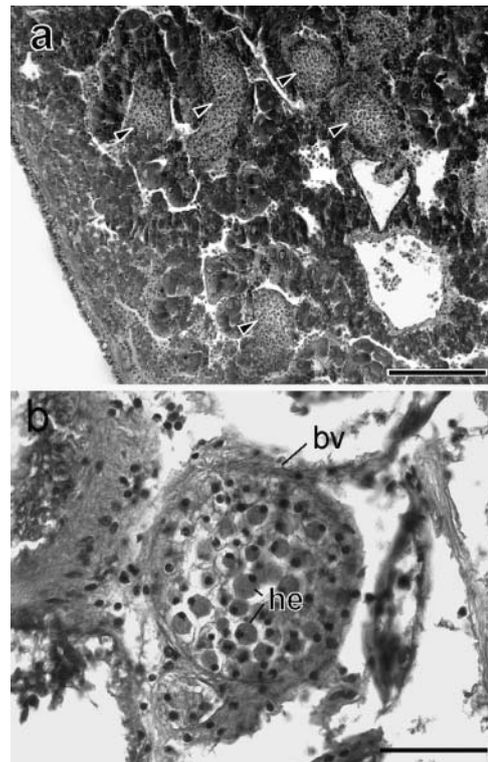


Fig. 12. *Bathymodiolus heckeræ*. (a) Hemocytic infiltration causing enlargement of blood vessels (intensity of 5; arrowheads) in visceral mass; scale bar = $200 \mu\text{m}$. (b) Enlarged blood vessel filled with hemocytes; hemocytes are enlarged, with nuclei displaced to side of cells; scale bar = $50 \mu\text{m}$. bv: blood vessel; he: hemocytes

4 categories of gonad development, with only 18% individuals possessing undifferentiated gonads. In *B. heckeræ* from Blake Ridge, 81% of individuals sampled had undifferentiated gonads. There was a negative correlation between total parasite body burden and degree of gonad development in *B. heckeræ* (Pearson's: $p < 0.030$); the density of Viral Inclusion III was negatively correlated with gonad development (Pearson's: $p < 0.040$). There was no difference in gonadal development in *B. puteoserpentis* from Snake Pit and Logatchev.

Index of endosymbiont density

In *Bathymodiolus puteoserpentis*, endosymbiont density was positively correlated with infection density of Gill Rickettsia I (Pearson's: $p < 0.020$; Fig. 13a). In *B. puteoserpentis* from Snake Pit, endosymbiont density also decreased with greater total body burdens of parasites in individual mussels and with an increase in infection density of Viral Inclusion I (Pearson's: $p < 0.020$; Fig. 13b).

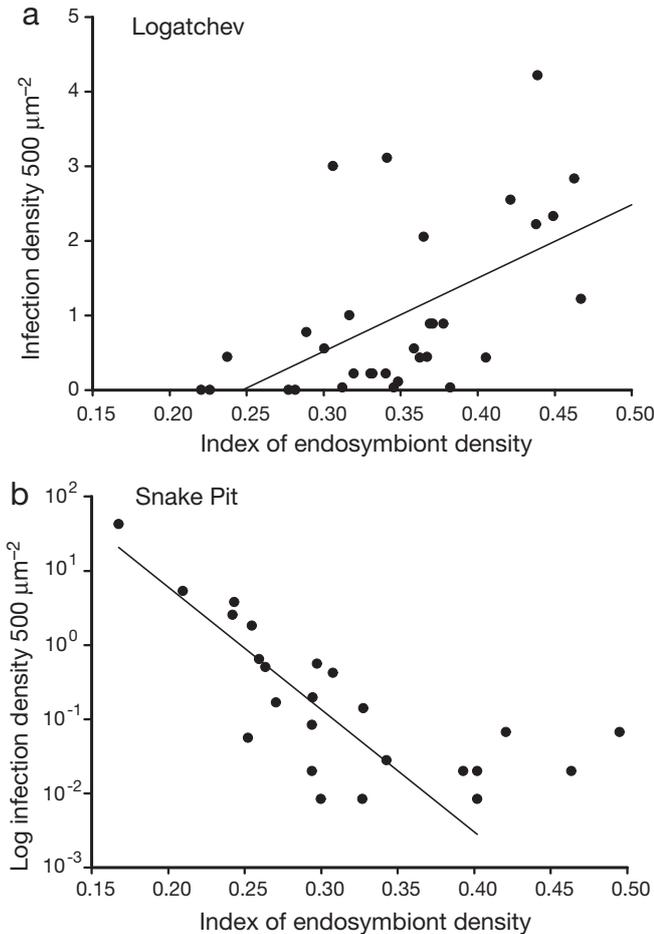


Fig. 13. *Bathymodiolus puteoserpentis*. Relationships between infection densities and index of endosymbiont density in vent mussels. (a) Infection density of Gill Rickettsia I ($500 \mu\text{m}^{-2}$ tissue) in mussels from Logatchev in relation to index of endosymbiont density; similar relationship was observed in mussels from Snake Pit. (b) Infection density of Viral Inclusion I ($500 \mu\text{m}^{-2}$ tissue) in mussels from Snake Pit in relation to index of endosymbiont density (Pearson's: $p = 0.013$); this relationship was not observed in mussels from Logatchev

DISCUSSION

Parasitism in seep and vent mussels

We observed 9 different types of parasites in *Bathymodiolus heckeriae* from the seep sites, and only 2 types of parasites in *B. puteoserpentis* from the vent sites. Only 1 type of parasite, Gill Rickettsia I, was common to mussels from both seep and vent habitats. We speculate that the ephemeral and extreme nature of vents may limit colonization and transmission of parasites, resulting in low parasite diversity in vent organisms (viz. Esch et al. 1990, Price 1990). Low diversity of invertebrates at vents compared to seeps has been attributed to the more ephemeral nature of vents

(Craddock et al. 1995, Sibuet & Olu 1998) and to the potential for greater barriers to invasion of vent habitats due to their more extreme conditions (e.g. elevated metal concentrations, greater sulfide flux; Turnipseed et al. 2003). Other factors might also be implicated in determining the lower parasite diversity at hydrothermal vents. For example, invasion of deep-sea habitats by ancestral, shallow-water host species and their parasites may be a function of proximity to shallow-water sites. The Florida Escarpment and Blake Ridge seep sites, located along the continental margins of eastern North America, are closer to shallow-water habitats than are the Logatchev and Snake Pit vent sites. Seep habitats, with a higher diversity of invertebrate taxa, are more likely to have higher parasite diversity than vent habitats, due to their greater number of potential hosts (Price 1990).

In shallow-water systems, species richness generally decreases as environmental stressors increase (Warwick & Clarke 1995, Menge & Branch 2001). Environmental stress, such as pollutants, may affect the physiology of the host organism, thus negatively affecting a host response to disease-causing agents or restricting available habitat within the host for parasitic agents (Laird 1961, Lauckner 1983, Cheng 1988, Winstead & Couch 1988). Parasites themselves can also be affected by environmental extremes. Deep-sea vent and seep sites contain noxious compounds such as sulfides, brines and metals. These compounds may act as barriers to invasion by parasites and limit infections to microbial agents, as few protozoan and no metazoan parasites other than commensal polychaetes were found in these deep sea mussels.

Pathogenicity

Viral Inclusion III

Viral-like infections were the most significant potential disease-causing agents found in the seep and vent mussels. Several lines of evidence, including relationships between mussel size and infection intensities of Viral Inclusion III, tissue destruction associated with Viral Inclusion III, undeveloped gonads in mussels infected by Viral Inclusion III, and *in situ* observations of extensive mussel mortality, suggest that Viral Inclusion III may be a significant pathogen with a potentially adverse affect on mussel populations at the Blake Ridge seep. Infection by Viral Inclusion III was pathogenic, causing significant damage to the digestive diverticula and connective tissues in the visceral mass. Although bacterial endosymbionts provide most of the nutrition to seep and vent mussels (Felbeck et al. 1981, Cavanaugh 1983, Childress et al. 1986), bathy-

modiolin mussels retain the ability to filter-feed, which presumably provides some nutrition to the mussels (Le Pennec & Hily 1984, Page et al. 1991). Viruses that form intranuclear inclusion bodies, such as those observed for Viral Inclusion III, can result in reduced food uptake in bivalves due to necrosis within the digestive tract (see Lauckner 1983). The high prevalence of mussels with undifferentiated gonads from Blake Ridge may have been a result of castration by Viral Infection III or resorption of gonads by an infected host, which is a common response of molluscs to disease (Lauckner 1983).

Several patches of empty valves have been observed in mussel populations from Blake Ridge, suggesting that mass mortalities have occurred at this site (Van Dover et al. 2003). While mortality could not be attributed to a specific cause, the valves were relatively uniform in size, suggesting a common cause of death. Mortality was attributed to shifting foci of seepage (Van Dover et al. 2003), but the pathogenicity of Viral Inclusion III suggests an alternative hypothesis, namely that viral infection may have contributed to mortality. Mussels with shell lengths >100 mm were either lightly infected or uninfected with Viral Inclusion III, and may represent a subset of the population that was immune to infection or strong enough to overcome the infection. Alternatively, Viral Inclusion III may be an opportunistic parasite infecting stressed hosts.

Hemocytic infiltration

Heavy infiltration by enlarged hemocytes was observed in the majority of mussels sampled. The predominant mechanism of molluscan internal defense involves phagocytosis by circulating hemocytes (Pipe & Coles 1995); and infiltration by hemocytes is part of the immune response to parasitic infection in molluscs (Quayle 1969, Lauckner 1983, Pipe & Coles 1995). The enlarged hemocytes seen in *Bathymodiolus heckeriae* and *B. puteoserpentis* may be abnormal and may impair host immune functions. The extreme infiltration by hemocytes into tissues and blood vessels was comparable to that observed in diseased, shallow-water bivalves (reviewed in Lauckner 1983). Although hemocyte infiltration can increase with parasitic infection, no correlation was observed between parasite infection densities and hemocyte infiltration in *B. heckeriae* or *B. puteoserpentis*. While hemocytes are important in molluscan internal defense, heavy hemocyte infiltration can hinder the normal flow of hemolymph, give rise to a reduction of host tissues needed for vital functions, and has been suggested as a possible cause of mass mortalities (Lauckner 1983, Villalba et al. 1997, Lee et al. 2001).

Prevalences of parasites

Of the parasite types observed in *Bathymodiolus heckeriae* (Gill Rickettsia I & II, gill ciliates, gill 'rosettes', and gut chlamydia) 50% were similar to parasites present in mussels (*Bathymodiolus* sp.) of relatively shallow (550 to 650 m) hydrocarbon seeps (Powell et al. 1999), and all but the gill rosettes are similar to common parasites found in coastal-zone mussels (Kim et al. 1998). Parasites resembling gill rosettes occur in the shallow-water lucinid clam *Loripes lucinalis*, which also hosts chemoautotrophic, endosymbiotic bacteria in its gills (Johnson & Pennec 1995, Powell et al. 1999). Trematodes (*Bucephalus*-like), which were the most pathogenic parasite observed by Powell et al. (1999) and which can be common in intertidal mussels (Lauckner 1983, Kim et al. 1998, Montaudouin et al. 2000), were entirely absent from *Bathymodiolus heckeriae* and *B. puteoserpentis* from the deeper vents and seeps. Trematodes were also absent from limpets from deep-sea hydrothermal vents and a deep-sea seep site (Terlizzi et al. 2004, this issue). Larval trematodes have a high host-specificity and use molluscs as their first intermediate hosts (see Lauckner 1983, Mouritsen et al. 1997). Transmission and infection by trematodes are affected by locality, season and host (Lauckner 1983, Montaudouin et al. 2000). The absence of trematodes in this study could be the result of factors such as depth, susceptibility to heavy-metal toxicity in trematode larval stages, or the lack of suitable intermediate and definitive hosts.

Polychaetes

Polychaetes are commonly found within the mantle cavities and between the gills of bathymodiolin mussels (Van Dover et al. 1999), but their true relationship with the host mussel is unknown. Gill flatworms (e.g. *Urastoma cyprinae*) are also common inhabitants of mantle cavities in coastal mussels (Villalba et al. 1997). Flatworms can disrupt the structure of gill filaments, cause necrosis of gill tissue, and thereby reduce the feeding capacity of the host, resulting in loss of condition. From these observations, Villalba et al. (1997) concluded that the flatworms were parasitic. Both the polynoid *Branchipolynoe seepensis* and the nautilielid *Laubierus mucronatus* polychaetes found in vent and seep mussels cause minor tissue damage to the gills and displacement of gill filaments, but there is no strong evidence for a detrimental effect on host condition. This is consistent with observations of Fisher et al. (1988), who found that a species of commensal polynoid polychaete in mussels at the Rose Garden hydrothermal vent (Galapagos Spreading Center;

eastern Pacific Ocean) did not cause gross lesions in soft tissues of the infected mussels. The polychaetes probably consume bacterial-laden mucus or pseudofeces, or free-living bacteria rather than tissues of the host mussel.

In summary, parasite burdens were greater in mussels at seeps than at vents. In some instances, parasites may affect reproductive output and overall health of the mussel populations, as observed in shallow-water bivalve populations. We suggest that infection by Viral Inclusion III influences population dynamics of *Bathymodiolus heckerae* at the Blake Ridge seep site, possibly causing widespread mortalities in juvenile mussels. Mussels from both seep and vent sites suffered from severe hemocytic infiltration by enlarged hemocytes. Hemocytic infiltration may be a symptom of disease or it may be an independent agent of mortality. How parasitic infections and hemocytic conditions affect individual mussels and the invertebrate communities associated with deep-sea mussel beds remains to be determined. Therefore, parasitological studies are a necessary addition to the more traditional ecological approaches used, in order to obtain an accurate representation of the factors affecting community structure.

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