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1 **Epsilonproteobacteria as gill epibionts of the hydrothermal vent gastropod *Cyathermia***  
2 ***naticoides* (North East-Pacific Rise)**

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13

14 **Abstract**

15

16 Molluscs, and particularly gastropods, are one of the major taxonomic groups at vents. In  
17 these ecosystems, devoid of light, chemoautotrophic bacteria are at the base of the food web,  
18 and symbiotic association between metazoa and these bacteria are numerous. Nevertheless,  
19 apart few "large size" well known species, the "small size" gastropods (shell < 5mm),  
20 although very abundant, remain poorly studied regarding symbioses. We investigated here  
21 *Cyathermia naticoides* (Warén and Bouchet 1989), a small coiled gastropod found in  
22 abundance on the East Pacific Rise among *Riftia pachyptila* tubes and usually inferred to  
23 graze on tubeworms bacterial cover, and/or filter feeding. Among molluscs, symbioses are  
24 well known in large species and almost exclusively rely on sulfide or methane-oxidizing  
25 Proteobacterial endosymbionts, occurring within the host tissues in gill epithelial  
26 bacteriocytes. Combining several approaches (molecular biology, microscopy, stable isotopes  
27 analyses), we described here an unusual symbiosis, where autotrophic filamentous  
28 Epsilonproteobacteria are located extracellularly, at the base of hosts gill filaments. Numerous  
29 endocytotic lysosome-like structures were observed in the gill epithelium of the animal  
30 suggesting bacteria may contribute to its nutrition through intracellular digestion by gill cells.  
31 Additional food source by non-symbiotic Proteobacteria grazed on *R. pachyptila* tubes could  
32 complete the diet. The possible role of temperature in the selection of Epsilon vs Gamma  
33 proteobacterial partners is discussed.

34

35

## 36 **Introduction**

37 To date, about 600 metazoan species have been reported at hydrothermal vents, belonging to  
38 12 phyla. Among those, 150 species of Mollusca and more than 100 species of Gastropoda  
39 have been described (Desbruyères et al. 2006), making them one of the major taxonomic  
40 groups at vents. Gastropod feeding habits are extremely diverse, although most species make  
41 use of a radula in some aspect of their feeding behavior (see review in Kohn 1983). Grazers  
42 can be herbivorous, rasping either micro- or macroalgae, or predators, rasping on encrusting  
43 invertebrates such as hydroids, sponges, cnidarians or ascidians. Herbivorous may also  
44 swallow sand containing algae. And carnivores may also hunt their prey and use their radula  
45 to drill mollusc shells or calcareous echinoids test, or perforate prey soft tissues (polychaetes,  
46 fishes,...). Some predators have lost the radula and engulf animal prey whole. Various  
47 feeding modes, using no radula, are also encountered in gastropods. In filter feeders,  
48 hypertrophy of the ctenidium as a cilary-mucous food collecting device is used as a trap to  
49 capture and sort particles suspended in seawater. Other feeding strategies include parasitic  
50 species, devoid of radula, that feed on body fluids thank to a sucker, and establishment of  
51 nutritional symbioses. Some herbivorous species can suck algal cell content and establish  
52 symbioses with chloroplastes (family Elysiidae) or zooxanthellae (family Aeolidae).  
53 Chemoautotrophic symbioses also occur in a wide range of habitats, including cold seeps,  
54 whale and wood falls, shallow-water coastal sediments and continental margins (Dubilier et  
55 al. 2008).

56 In the hydrothermal vent environment, chemosynthetic production by bacteria is the main  
57 food source of primary consumers (Felbeck and Somero 1982). The majority of hydrothermal  
58 gastropods are thus grazers or filter feeders that appear to feed on free-living bacteria (Bates  
59 2007a). Another widespread strategy at vents is symbiotic association with chemoautotrophic  
60 bacteria. Up to now, such symbioses have been demonstrated in at least 7 different phyla  
61 (Dubilier et al. 2008), including molluscs, the most famous examples being described in gills  
62 of bivalves (Bathymodiolinae and Vesicomidae) and involve sulfur-oxidizing  
63 Gammaproteobacteria endosymbionts. But it also exists among gastropods. The best known  
64 examples are *Alviniconcha hessleri* and *Ifremeria nautilei*, found in the western Pacific  
65 (Windoffer and Giere 1997; Borowski et al. 2002; Suzuki et al. 2005a, 2005b, 2006; Urakawa  
66 et al. 2005; Saito and Hashimoto 2010). Most of the examples described for these two species  
67 also rely on sulfur-oxidizing Gammaproteobacteria gill endosymbionts. But recently,  
68 Epsilonproteobacteria were described as gill endosymbionts in some species of Provannidae  
69 (Urakawa et al. 2005; Suzuki et al. 2006, Beinart et al. 2013). Symbioses in smaller gastropod  
70 species remain poorly studied and the presence of bacteria as symbionts has only been

71 documented in *Lepetodrilus fucensis* from the Juan de Fuca Ridge (Bates 2007a ), in which a  
72 nutritional role has been suggested.  
73 In this study, we investigate a coiled gastropod, the Neomphalina *Cyathermia naticoides*  
74 (Warén and Bouchet 1989). Not much is known about it, despite it is a common species,  
75 found in abundance among *Riftia pachyptila* clumps (Mills et al. 2007), and in lower  
76 abundances among *Alvinella pompejana* and *Bathymodiolus thermophilus* (Warén et al.  
77 2006). Up to now, *C. naticoides* was inferred to graze on tubeworms bacterial cover, but also  
78 to use filter feeding, based on its very large bipectinate gill (Warén and Bouchet 1989). A  
79 distinct labial notch described by Warén and Bouchet (1989) in the shell morphology is  
80 interpreted as an adaptation to allow the gill to be extended outside the shell even when the  
81 snail is resting on the substrate, partially retracted into the shell (Warén and Bouchet 1989;  
82 Sasaki et al. 2010). Here we investigate an additional hypothesis as feeding strategy in *C.*  
83 *naticoides*. Our study describes a unusual symbiosis, where epibiotic autotrophic  
84 Epsilonproteobacteria are endocytosed within the gill filaments of the animal. The large size  
85 of the gill, the recurrent observation of endocytosis and lysis of bacteria, and the stable  
86 isotopes results advocate for a nutritional symbiosis.

87

88

89

## 90 **Material and methods**

91 **Animal collection and conditioning.** *Cyathermia naticoides* specimens were collected,  
92 among *Riftia pachyptila* tubes, using the DSV Nautille during the Mescal 2010 cruise (East  
93 Pacific Rise, 2,500 m depth), on two different sites : 9°50'N (Bio9 site) and 12°50'N (Genesis  
94 site). Once on board, the entire specimens were fixed (operculum removed) in: 1) 2.5%  
95 glutaraldehyde (for light and electron microscopies), 2) ethanol (for DNA extraction), 3) 2-  
96 4% formaldehyde (for Fluorescent In Situ Hybridization, FISH) and 4) liquid nitrogen (for  
97 stable isotope analyses and chitinase activity assays).

98

99 **Light and Electron Microscopies.** Gut and gill tissues of 3 specimens of *C. naticoides* from  
100 9°50'N were dissected. Two other specimens were embedded whole. Samples were post-fixed  
101 in osmium tetroxide 1%, dehydrated in increasing ethanol series (50, 70, 95 and 100%) and  
102 embedded in Epon resin (48 h, 60°C). Sections were cut using a Reichert–Jung  
103 ultramicrotome. Semi-thin (600 nm) sections were stained with toluidine blue and observed  
104 with an Olympus BX 61. Thin (60 nm) sections were mounted on copper grids, contrasted  
105 using uranyl acetate and observed using a HITACHI H-7100 transmission electron

106 microscope, operated at 80 kV.

107

### 108 **Fluorescence In situ hybridization (FISH)**

109 Four specimens from 9°50'N and 2 from 12°50'N were used. Specimens were pulled out their  
110 shells, and after 2-4 hours in 4% formaldehyde, were rinsed and dehydrated in 50, 70 and  
111 96% ethanol. They were then embedded whole in polyethylene glycol (PEG) distearate: 1-  
112 hexadecanol (9:1). Sections of 7-10 µm were cut using a Jung microtome and deposited on  
113 Superfrost Plus slides. Wax was removed and tissue rehydrated in decreasing ethanol series.  
114 Sections were hybridized as described in Zbinden et al. (2010), using 30% formamide for 3  
115 hours at 46°C, rinsed (15 min, 48°C), covered with SlowFade containing DAPI, and  
116 examined under an Olympus BX-61 epifluorescence microscope (Olympus, Japan).  
117 Following probes, labeled with Cy-3 and Cy-5, were used: Eub-338 (5'-  
118 GCTGCCTCCCGTAGGAGT-3', Amann et al. 1990), Gam-42 (5'-  
119 GCCTTCCCACATCGTTT-3', Manz et al. 1992), Del-495a (5'- AGTTAGCCGGTGCTTST  
120 - 3', Loy et al. 2002), Epsy-549 (5'-CAGTGATTCCGAGTAACG-3', Manz et al. 1992) and  
121 Arc-94 (5'- TGCGCCACTTAGCTGACA - 3', Moreno et al. 2003). Phylotypes targeted by  
122 the different probes are indicated in Table 1.

123

### 124 **DNA extraction and 16S rRNA amplification**

125 DNA was extracted from 3 specimens from 9°50'N and 3 from 12°50'N. Specimens were  
126 dissected as follows: at 9°50'N, gill was extracted from two of the specimens (9-1-Gi and 9-2-  
127 Gi), while the visceral mass (i.e. the rest of the animal containing the digestive tract and the  
128 heart, gonad, digestive gland, liver, and excretory organs), was treated separately (9-1-VM  
129 and 9-2-VM). A third specimen (9-3) was treated as a whole. DNA was also extracted from  
130 the empty shell of specimen 2 (9-2-Sh). At 12°50' N, DNA was extracted separately from the  
131 gill and visceral mass of three specimens (12-1, 12-2, 12-3). Extractions were performed  
132 using the DNA Tissue Kit (Qiagen). A ~1500bp fragment of the 16S rRNA-encoding gene  
133 was amplified by PCR using primers 27F and 1492R, over 32 cycles. Three PCR products  
134 were pooled together for each sample, to reduce PCR bias. Fragments were cloned using a  
135 TOPO TA Kit (Invitrogen, CA). 11 to 32 clones were successfully sequenced from each  
136 sample by GATC Biotech (Table 1).

137 Genes encoding for key enzymes of sulphur oxidation (*aprA*) and autotrophic carbon fixation  
138 (*aclB*) were sought. Fragments of the *aprA* gene encoding APS (adenosine 5'-phosphosulfate)  
139 reductase and the *aclB* gene encoding ATP Citrate Lyase (the key enzyme in reverse  
140 tricarboxylic acide (rTCA) cycle) were amplified using primer sets *aps1F/aps4R* and

141 892F/1204R, respectively as described previously and using 32 to 35 PCR cycles (Campbell  
142 et al. 2003, Meyer and Kuever 2007). Obtained PCR products were cloned and inserts were  
143 sequenced (Table 2).

144

### 145 **Gene sequence analyses**

146 Chromatograms were checked for quality. For each sample, sequences were aligned, and  
147 grouped in Operational Taxonomic Units (OTUs) when >97% of the nucleotide positions  
148 were identical. For the 16S rRNA-encoding genes, overall 11 OTUs were present in more  
149 than a single sample. Sequences were compared with databases using BLAST (Altschul et al.  
150 1990; Cole et al. 2009), and the 8 OTUs for which full sequences were available were  
151 included in a dataset with their best hits and reference sequences, and aligned using SINA  
152 Web Aligner (Pruesse et al. 2007). Alignment was manually checked, and phylogenetic  
153 relationships were inferred by using the Maximum Likelihood (ML) method. For the  
154 reconstruction, a General Time Reversible model, with a Gamma distribution of evolutionary  
155 rates among sites was used (5 categories and invariant sites). For *aprA* and *aclB* genes,  
156 recovered sequences were compared to the database using BlastX (Table 2). For *aclB*, best  
157 hits and representative sequences were included in a dataset, and phylogenetic reconstruction  
158 was based on a 100 aa-long fragment using a ML approach, a JTT model of amino acid  
159 substitution and a Gamma distribution of evolutionary rates among sites (tree in Fig. S1). All  
160 analyses were conducted using MEGA6 (Tamura et al. 2013).

161

### 162 **Stable isotope analysis**

163 Frozen *Cyathernia naticoides* ( $n = 4$  for  $\delta^{13}\text{C}$  and  $\delta^{15}\text{N}$ ,  $n = 10$  for  $\delta^{34}\text{S}$  for  $9^{\circ}50'\text{N}$  and  $n = 5$ ,  
164 only  $\delta^{13}\text{C}$  and  $\delta^{15}\text{N}$ , for  $12^{\circ}50'\text{N}$ ) were dissected under a dissecting microscope to remove the  
165 shell. Specimen tissues were rinsed in distilled water, dried (3 days,  $60^{\circ}\text{C}$ ) and then reduced  
166 into powder. To avoid significant changes in  $\delta^{15}\text{N}$  isotopic composition, no HCl was used to  
167 remove carbonates (Kaehler and Pakhomov 2001). About 1 mg ( $\pm 0.1$  mg) of dried tissues  
168 (except 8 mg for sulfur stable isotopes; pool of 10 specimens) were analyzed by a GV  
169 IsoPrime (UK) stable isotope mass spectrometer (Iso-Analytical, Crewe, UK). Values of  $\delta^{13}\text{C}$ ,  
170  $\delta^{15}\text{N}$  and  $\delta^{34}\text{S}$  were determined and expressed as relative per-mil (‰) differences between  
171 samples and Pee Dee Belemnite (PDB) for carbon, air  $\text{N}_2$  for nitrogen and Canyon Diablo  
172 Troilite for sulfur according to the following equation:

$$173 \quad \delta(X) = \left[ \left( \frac{R_{\text{sample}}}{R_{\text{standard}}} \right) - 1 \right] * 1000$$

174 where  $X$  (‰) is  $^{13}\text{C}$ ,  $^{15}\text{N}$  or  $^{34}\text{S}$  abundance and  $R$  is the  $^{13}\text{C}/^{12}\text{C}$ ,  $^{15}\text{N}/^{14}\text{N}$  or  $^{34}\text{S}/^{32}\text{S}$  ratios.

175

176 **Chitinolytic activity assays.**

177 Chitinolytic activity was determined using a modification by Gutowska et al. (2004) of the  
178 standard procedure of Jeuniaux (1966), which measures the production of N-Acetyl  
179 glucosamine (NAG). Five specimens from 12°50'N were removed from their shell and ground  
180 together in liquid nitrogen. The powder was homogenized in 0.15 M citric acid, 0.3 M  
181 Na<sub>2</sub>HPO<sub>4</sub> buffer (pH = 5 and pH = 7). The homogenates were then centrifuged at 2000 g for  
182 10 min at 4°C, the supernatants were recovered and assayed for their chitinolytic activity. The  
183 standard mixture consisted of 2 vol. of tissues extract, 1 vol. of chitinase and 1 vol. of chitin  
184 suspension (5 mg.ml<sup>-1</sup>). Two control assays were added, with either the tissues extract or the  
185 chitin solution replaced by distilled water. For comparison, assays with commercial  
186 *Streptomyces griseus* chitinase (Sigma C6137) were also conducted in parallel. They were all  
187 incubated at 37°C, and aliquots were taken after t = 0 min, t = 90 min and t = 180 min.  
188 Chitinolytic reaction was stopped by mixing 1 vol. of the reaction medium with 1 vol. of  
189 boiling water. The mix was placed 10 min at 100 ° C, then centrifuged at 2000 g for 10 min.  
190 The supernatants were used for NAG measurements by adding K<sub>2</sub>B<sub>4</sub>O<sub>7</sub> 0.8M, and further  
191 incubating for 3 min at 100°C. p-dimethylaminobenzaldehyde (DMAB) was added and after  
192 20 min at 37°C, the concentration of NAG released was determined by comparing each  
193 sample's absorbance at 585 nm to NAG standard curves. The activity was expressed as µg of  
194 NAG released per gram protein per hour.

195

196 **Results**

197 **Morphology and ultrastructure of gut and gill**

198 *Cyathernia naticoides* is a gastropod with a regularly coiled shell (diameter up to 7 mm),  
199 with a deep notch at basal side of the outer lip. This gastropod possesses a very large  
200 bipectinate gill, which occupies most of the anterior part of the animal (Fig. 1a, b). Semi-thin  
201 sections revealed an abundant bacterial community associated with the gill (Fig. 1c, d). These  
202 gram-negative filamentous bacteria (0.5 to 0.6 µm in diameter and up to 5.8 µm in length,  
203 Fig. 2a, b) were located extracellularly, between the gill filaments, mainly at their base. Some  
204 of the bacteria were free in the inter-lamellar space, but many have also been observed  
205 trapped in lysosome-like structures in the gill epithelium (see Fig. 1d and 2a), and appeared to  
206 undergo different stages of degradation (Fig. 2b). Bacterial colonization and endocytosis  
207 occurred on each animal and all sections observed.

208 The digestive tract contained pieces of *Riftia pachyptila* tubes (not shown), recognizable by  
209 chitin microfibrils organized in parallel bundles with various orientations (Gaill and Shillito

210 1995). No bacteria similar to those present on the gill were observed in the gut contents.

211

### 212 **Fluorescence microscopy observations**

213 Probes Eub-338 and Arc-94 yielded strong signals in regions of the gill filaments of  
214 *Cyathernia naticoides* from both 9°50'N and 12°50'N (Fig. 1e, f). Cy-3 labeled probe Epsy-  
215 549 yielded weaker signal, but signal-to-noise ratio was greatly improved when letting tissue  
216 autofluorescence decrease under the laser for 30 seconds. Signals from the three probes fully  
217 overlapped in gills (not shown). Hybridized objects corresponded to thin filamentous bacteria.  
218 Parts of the gill filaments were free of bacteria and did not display any signal, suggesting that  
219 the distribution of bacteria was not homogeneous (Fig. 1f). Probes Gam-42 and Del-495a did  
220 not display any signal in the gills. No FISH signal was observed from the gut epithelium or  
221 content.

222

### 223 **Chitinolytic activity**

224 Chitinase activity was assayed on crude extracts from whole specimens. The measured  
225 activity (20 µg NAG released g<sup>-1</sup> protein h<sup>-1</sup>) was very weak when compared to the reference  
226 sample (*Streptomyces griseus* chitinase, 6500 µg NAG released g<sup>-1</sup> protein h<sup>-1</sup>). Another  
227 gastropod (*Lepetodrilus elevatus*) found on *R. pachyptila* tubes was also analyzed for  
228 comparison, and showed a two-fold higher activity (42 µg NAG released g<sup>-1</sup> protein h<sup>-1</sup>),  
229 which was still very weak when compared to *S. griseus*.

230

### 231 **Bacterial communities associated with digestive tract and gill**

232 Out of 295 sequences obtained, 244 (83%) belonged to one of the 11 OTUs (defined as  
233 groups of sequences displaying above 97% identical positions) that were present in more than  
234 a single sample (see Table 1). The remaining sequences corresponded to single reads  
235 occurring in a single sample. Above 78.0% of total sequences, and 7 of the 11 OTUs (94.3%  
236 of the OTUs-assigned sequences) were assigned to the Epsilonproteobacteria. Five of the  
237 OTUs (1, 3, 8, 11, 16-p), including the 4 most abundant and representing 90% of the OTU-  
238 assigned sequences, were present at both sampling sites 9°50'N and 12°50'N.

239 OTUs 3, 1 and 8 dominated clone libraries, representing 23.1, 20.0 and 19.3% of the total  
240 sequences, respectively (Table 1). OTU 3 was present in gills and visceral mass at 9°50'N and  
241 gill at 12°50'N. This sequence displayed above 98% identity and was most closely related to  
242 sequences from an epibiont of the gastropod *Lepetodrilus fucensis*, and to various *Arcobacter*  
243 from the EPR (Fig. 3). OTU 1 was present in all samples from 9°50'N, and in visceral mass  
244 samples from two specimens at 12°50'N. The sequence was closely related and highly similar



245 (>98%) to sequences from bacteria associated with the tube of *Ridgeia piscesae* on the EPR.  
246 The third, OTU 8, was present in gill, visceral mass and shell at all sites and displayed above  
247 98% identity with several sequences related to the sulfur-oxidizing chemolithotroph  
248 *Sulfurovum* from the Brothers Seamount (Kermadec Arc) and vents.

249

### 250 **Functional gene analysis**

251 Overall 70 *aprA* sequences were obtained from the visceral mass of three specimens, and  
252 none from the gill tissue. Six distinct nucleotide sequences were obtained, 4 of which were  
253 related to Gammaproteobacteria and represented above 94% of total sequences (Table 2). The  
254 dominant sequence, clone 761, was 96% similar (amino acids) to a sequence from an epibiont  
255 of the Yeti crab *Kiwa hirsuta*. Other sequences were similar to a sequence from a bacterium  
256 associated with the oligochete *Tubificoides benedii* and from the tube of the siboglinid annelid  
257 *Lamellibrachia anaximandri* (Table 2).

258 PCRs on the *aclB* gene yielded faint bands from the visceral mass of specimen 9-1 and 9-2-,  
259 and from the gills of specimen 9-2. Out of 28 sequenced clones from each of these 3 samples,  
260 only 35 good quality sequences were obtained. The majority (24) were from the gill, of which  
261 19 corresponded to a single sequence, clone 765, and were related to various sequences of  
262 Epsilonproteobacteria from hydrothermal vents and to epibionts from the gill chamber of the  
263 vent shrimp *Rimicaris exoculata* (around 98% amino acid similarity, Table 2 and Fig S1).

264

### 265 **Stable isotope composition**

266 The  $\delta^{13}\text{C}$  values of *Cyathernia naticoides* varied following vent sites, with  $-9.0\text{‰}$  ( $\pm 0.3\text{‰}$ )  
267 for  $9^{\circ}50'\text{N}$  and  $-10.8\text{‰}$  ( $\pm 0.4\text{‰}$ ) for  $12^{\circ}50'\text{N}$  (Fig. 4). These stable isotopic ratios of carbon  
268 fall after correction of fractionation ( $1\text{‰}$  for  $\delta^{13}\text{C}$  and  $3.3\text{‰}$  for  $\delta^{15}\text{N}$ ) into the range of stable  
269 isotopic ratios of carbon of Epsilonproteobacteria (range between  $-8$  and  $-12\text{‰}$ ; Campbell et  
270 al. 2003) (Fig. 4). Regarding  $\delta^{15}\text{N}$ , value of both sites was  $6.8\text{‰}$  ( $\pm 0.5\text{‰}$ ). This nitrogen stable  
271 isotopic ratio is typical of a primary consumer at vents (between  $4$  and  $8\text{‰}$ ). Isotopic signatures  
272 of  $\delta^{34}\text{S}$  measured in a pool of *C. naticoides* ( $n = 10$ ) was  $5.5\text{‰}$ .

273

## 274 **Discussion**

### 275 **Grazing vs. filter-feeding**

276 Gastropods have two main feeding strategies: i.e. grazing, using their radula to rasp various  
277 kinds of substrates, or filter feeding, using their gills as a trap to capture and sort particles  
278 suspended in seawater. At hydrothermal vents, where the organic matter synthesis relies on  
279 chemoautotrophic bacteria, most gastropods appear to feed on free-living bacteria (Bates et al.

280 2007b). Up to now, *Cyathernia naticoides* was inferred to graze on tubeworms bacterial  
281 cover, but also to use filter feeding, based on its very large bipectinate gill (Warén and  
282 Bouchet 1989). Grazing on tubeworms bacterial cover is congruent with our observations, as  
283 we noted the occurrence of *Riftia pachyptila* tube pieces in the gut content. Conversely, filter  
284 feeding on bacteria is not well-supported by our data. The position of the filamentous  
285 bacteria, deep between the filaments, and the large number of endocytosed and lysed bacteria  
286 advocates for a stronger association than a classic trapping through filter feeding mechanism  
287 for transport to the gut. Nevertheless, filter feeding on particulate organic matter cannot be  
288 discarded.

### 289 290 **A diet based on bacteria**

291 Despite *R. pachyptila* tube pieces, which contain up to 25% of chitin (Ravaux et al. 1998), are  
292 rasped and ingested by *C. naticoides*, a weak chitinolytic activity was measured for this  
293 species, when compared to the reference chitinase of *Streptomyces griseus*, or to values  
294 obtained for chitin degrading animals, such as fishes feeding on crustaceans (Gutowska et al.  
295 2004). This suggest a minor nutritional input of chitin and that *C. naticoides* rather grazes on  
296 *R. pachyptila* tubes for feeding either on proteins contained therein (representing 37-41% of  
297 the tube, Ravaux et al. 1998) or on the bacterial biofilm. López-García and collaborators  
298 (2002) observed dense microbial populations on *R. pachyptila* tubes, with very diverse 16S  
299 rRNA phylotypes, belonging mostly to Epsilon-, but also to Delta-, Alpha- and Gamma-  
300 proteobacteria. Interestingly, among our recovered bacterial phylotypes, OTUs 1 and 11 are  
301 closely related to several sequences from the tubes of *Ridgeia piscesae* and *Riftia pachyptila*  
302 (Fig. 3; Forget and Juniper 2013; López-García et al. 2002). These could be bacteria ingested  
303 alongside with tube fragments. Indeed, of the 59 sequences of OTU 1 for example, 36 were  
304 from the visceral mass of *C. naticoides* (3.4 to 63% of the sequences depending on the  
305 sample). This sequence displays 5 and 1 mismatches with FISH probes Arc94 and Epsy549  
306 and thus does not hybridize with them. It is thus surely not the sequence from the bacteria  
307 located in the gills, which respond to both probes. Sequences encoding APS reductase were  
308 successfully amplified from the visceral mass of all tested specimens. Related sequences were  
309 typically associated with the tube or cuticle of protostomes. These might again correspond to  
310 sequences of bacteria ingested with scrapings of tubes. The dense colonies of filamentous  
311 Epsilonproteobacteria observed on *C. naticoides* gill surface could be another nutritional  
312 pathway. Many are indeed endocytosed within lysosomes, arguing for a internal digestion of  
313 bacteria in the gill epithelium.

314 Vent gastropods at hydrothermal vents are considered as primary consumers feeding both on  
315 free-living bacteria of different origins but also on particulate organic matter (Limen et al.

316 2007), resulting in a range of stable isotopes nitrogen ratio between 4 and 8 ‰ (Bergquist et  
317 al. 2007; Limén et al. 2007; Gaudron et al. 2012), which includes our values for *Cyathernia*  
318 *naticoides*. Several species of *Lepetodrilus* are known to be also primary consumers  
319 (heterotrophic) such as *L. elevatus*, *L. pustulosus* and *L. ovalis*, displaying the same range of  
320 stable isotopic nitrogen values (Fig. 4). However *L. fucensis* known to harbor symbiotic  
321 bacteria within its gills also has a similar stable isotopic nitrogen value (7‰, Bates et al.  
322 2011), as well as others larger symbiotic gastropods (*Alviniconcha* spp) harboring  
323 Epsilonproteobacteria (Fig. 4), meaning that heterotrophic and symbiotic diet may co-occur in *C.*  
324 *naticoides* and cannot be easily distinguished based on nitrogen isotopes.

325 In the previously studied symbioses involving Epsilonproteobacteria, a trophic role has been  
326 suggested based on carbon stable isotope signatures of hosts ( $\delta^{13}\text{C}$  values between -11 and -  
327 10.7‰ for *A. aff hessleri* ; -12.8 and -11.2‰ for *Alvinella pompejana* and -12 to -10‰ for  
328 *Rimicaris exoculata*, Suzuki et al. 2005b; Desbruyères et al. 1998; Polz et al. 1998). These  
329 values indeed fall within the range of typical values measured in Epsilonproteobacteria which  
330 use the reverse TCA cycle for autotrophic carbon fixation (-12 to -8‰, Campbell et al. 2006;  
331 Sievert and Vetriani 2012). Similar values are measured in *C. naticoides* (-10.84 ± 0.58‰;  
332 Fig. 4), suggesting that similar bacteria may significantly contribute to the hosts diet, either  
333 those grazed on tubes or those endocyted in the gill. López-García and collaborators (2002)  
334 identified that most bacteria (68%) present on *R. pachyptila* tube surface belonged to the  
335 Epsilonproteobacteria ( $\delta^{13}\text{C}$  value for scrappings from *R. pachyptila* tubes are -12,5‰). In  
336 *Cyathernia*, this is further supported by the identification of ATP Citrate Lyase-encoding  
337 genes from the visceral mass of specimens which confirm the presence of rTCA. On the other  
338 hand, a single dominant ATP Citrate Lyase sequence is also identified from gill-associated  
339 bacteria, which supports the hypothesis of a significant contribution of the gill bacteria to the  
340 host carbon nutrition. The next step will be to quantify the respective roles of gill-associated  
341 versus ingested bacteria.

342 The vast majority of symbiotic bacteria described at present in molluscs are chemoautotrophic  
343 sulfur-oxidizing bacteria. The  $\delta^{34}\text{S}$  value of an animal indicates the origin of the assimilated  
344 sulfur. Marine invertebrates for which the sulfur source comes from chemosynthetic sulfur-  
345 oxidation have values lower than 5 ‰.  $\delta^{34}\text{S}$  of *C. naticoides* (5‰) is at the upper end  
346 normally measured into thiotrophic metazoans, which is between -25 to 5‰ (Vetter and Fry  
347 1998), allowing to suppose that some sulfur absorbed by the animal comes from  
348 chemosynthetic sulfur-oxidizers. If evidence for thiotrophic metabolism has been shown  
349 through sequencing of APS reductase in the visceral mass (possibly coming from the bacteria

350 in the gut, rasped on *R. pachyptila* tubes), no positive PCR result was obtained from gills. We  
351 cannot thus confirm the thiotrophic metabolism of gill-associated bacteria.

352

### 353 **Symbiosis**

354 A widespread feeding strategy at hydrothermal vents is to obtain organic carbon through  
355 symbiotic associations. Among molluscs, symbioses are well known and described in large  
356 species, such as the mussels and clams (Mytilidae and Vesicomidae, Dubilier et al. 2008) or  
357 the large gastropods *Ifremeria nautilei* and *Alviniconcha hessleri* (Provannidae, Borowski et  
358 al. 2002; Suzuki et al. 2005a, b). In all these symbioses, sulfide-oxidizing

359 Gammaproteobacterial symbionts are endosymbionts, occurring within the host tissues in gill  
360 epithelial bacteriocytes. The hosts are fuelled by by-products of bacterial metabolism  
361 (ultimately relying on sulfide oxidation) or intracellular bacterial digestion (Bates 2007b).

362 Here we described the occurrence of a dense population of filamentous bacterial located  
363 extracellularly at the base of the gill filaments. Some are free in the inter-lamellar space, but  
364 many have also been observed trapped in lysosome-like structures, in the gill epithelium. At  
365 hydrothermal vents, epibiotic symbioses have been described in only a few groups:

366 Ciliophora (Kouris et al. 2007), annelids (*Alvinella pompejana*, Haddad et al. 1995; Cary et  
367 al. 1997; Bright and Giere 2005) and crustaceans (*Rimicaris exoculata*, Segonzac et al. 1993;  
368 Zbinden et al. 2008; Petersen et al. 2010; the galatheid crabs *Kiwa hirsuta* (Macpherson et al.  
369 2005; Goffredi et al. 2008), *Kiwa puravida* (Thurber et al. 2011), and *Shinkaia crosnieri*  
370 (Miyake et al. 2007). In Molluscs, only very few examples are known : in Aplacophora (Katz  
371 et al. 2006) and in Gastropoda (*Lepetodrilus fucensis*, Bates 2007a, b). The kind of symbiosis  
372 described in *L. fucensis* is the closest to what we observed in *Cyathermia naticoides*, with a  
373 few exceptions. *L. fucensis* hosts dense colonies of filamentous bacteria on its gill surface,  
374 where bacteria are found partially embedded in the host's gill epithelium and extend into the  
375 fluid circulating between the lamellae (de Burgh and Singla 1894 ; Bates et al. 2007a, b).

376 Frequent endocytosis was observed in the epithelium (de Burgh and Singla 1984). Observed  
377 residual bodies of lysosome-like organelles, with concentric membrane stacks, mirror our  
378 observations. The main difference between *L. fucensis* and *C. naticoides* is that most  
379 abundant *L. fucensis* epibionts are Gammaproteobacteria (Bates et al. 2011), and those of *C.*  
380 *naticoides* belong to Epsilonproteobacteria. Furthermore, stable isotopes analyses ( $\delta^{13}\text{C} = -$   
381  $19.5$  to  $-14.8\text{‰}$  and  $\delta^{15}\text{N} = 2.5$  to  $5\text{‰}$ ) situate *L. fucensis* within a group of known deposit  
382 feeding invertebrates at the Juan de Fuca Ridge vents (Fox et al. 2002), whereas *Cyathermia*  
383 values fall within the range of typical values measured in Epsilonproteobacteria, and in

384 organisms living in symbiosis with these bacteria (-12 to -8‰, Campbell et al. 2006; Sievert  
385 and Vetriani 2012).

386 As suggested by de Burgh and Singla (1984) and Bates (2007a), there are 3 ways in which the  
387 gill bacteria may contribute to the organic carbon of the host : 1) the bacteria may be farmed  
388 and ingested; 2) dissolved organic molecules, byproduct of the bacterial metabolism, may  
389 pass from the bacteria to the host through the epithelium, as it was suggested for *Alvinella*  
390 *pompejana* and evidenced for the shrimp *Rimicaris exoculata* and its epibionts (Ponsard et al.  
391 2013); 3) bacteria may be endocytosed in the gill epithelium and digested within lysosomes.  
392 For *L. fucensis*, Bates (2007b) argues that endocytosis of bacteria by the gill epithelium  
393 followed by lysosomal digestion (de Burgh and Singla 1984) may not be an important feeding  
394 mechanism. In our case, the huge number of lysosome-like structures observed, with bacteria  
395 at different stages of degradation conversely rather advocates for the third hypothesis.  
396 Nevertheless, additional contribution by the two other ways cannot be discarded.

397 These gill bacteria likely correspond to our OTU 3, which is the predominantly associated  
398 with gill samples. Indeed, 43 of the 68 sequences were from gill samples, representing  
399 between 22 to 100% of sequences in the various gill samples. Besides, OTU 3 was present in  
400 gills of all specimens at both sites, and far less abundant in visceral mass samples,  
401 representing only from 0 to 22% of the sequences. Furthermore, it responds to both Arc-94 and  
402 Epsy-549 probes, as do gill bacteria observed using FISH. Finally, it is closely related to one  
403 of the documented gill epibionts of *Lepetodrilus fucensis*. OTU 3 might be a widespread  
404 epibiont of gastropod gills.

405 The third most abundant OTU identified in our clone libraries, namely OTU 8, related to  
406 *Sulfurovum* also responds to both FISH probes. However, only 10 of the 57 recovered  
407 sequences were from gill tissue, representing 0 to 29% of sequences depending on gill  
408 sample, while 14 were found on the shell analyzed (45%) and 33 in the visceral mass. Closest  
409 relatives do not include any reported symbiont. This bacterium is thus most probably an  
410 environmental bacterium, although this cannot be ascertained using FISH probes from this  
411 study.

412 As seen above, the large majority of the mollusc-associated symbionts from chemosynthetic  
413 environments are Gammaproteobacteria (in the Thyasiridae, Lucinidae, Solemyidae,  
414 Vesicomomyidae, Mytilidae, and some Provanidae). But recently (Suzuki et al. 2005a, 2005b),  
415 Epsilonproteobacteria were described as symbionts (and as endosymbionts) in some species  
416 of Provannidae (*Alviniconcha* sp.). *A. hessleri* from the Mariana Trough, *Alviniconcha* sp.  
417 type 1 from Manus Basin and Fiji, and *Alviniconcha* sp. from Lau Basin harbors sulfur-  
418 oxidizing chemoautotrophic Gammaproteobacterial endosymbionts that mediate the Calvin-

419 Benson cycle to fix CO<sub>2</sub>. Whereas *Alviniconcha aff. hessleri* from the Central Indian Ridge  
420 and *Alviniconcha* sp. type 2 from Manus Basin and Fiji harbors chemoautotrophic  
421 Epsilonproteobacterial endosymbionts that mediate the reductive tricarboxylic acid (rTCA)  
422 cycle for CO<sub>2</sub> fixation (Urakawa et al. 2005; Suzuki et al. 2006). A fragment of the gene  
423 encoding ATP Citrate Lyase was identified in the gill and visceral mass of *C. naticoides*. In  
424 particular, the most abundant sequence in the gills was related to sequences from various vent  
425 bacteria including gill epibionts of the vent shrimp *R. exoculata*. This advocates for the  
426 presence of this pathway in gills of *C. naticoides*. This finding is congruent with the  
427 dominance of Epsilonproteobacteria in the gill, and it is possible that the dominant *aclB*  
428 sequence (clone 765) is indeed associated with the dominant 16S rRNA OTU 3, or one of the  
429 other dominant gill-associated phylotypes.

430 Symbiotic association with filamentous Epsilonproteobacteria have been described, but  
431 mostly as ectosymbioses, as in the crustaceans *R. exoculata*, *K. hirsuta*, *K. puravida* or *S.*  
432 *crossnieri* and in the annelid *A. pompejana* (see references above). The *C. naticoides*  
433 symbiosis described here thus represents a unusual type of association in the long list of  
434 symbiosis within Molluscs, with Epsilonproteobacteria as ectosymbionts being the first  
435 exemple of this combination in molluscs, to our knowledge.

436

#### 437 **Epsilon- versus Gammaproteobacteria : an issue with temperature ?**

438 Urakawa and collaborators (2005) suggest that thermal gradient may affect the acquisition  
439 and evolutionary selection of either Epsilon- or Gammaproteobacterial symbionts. Vent hosts  
440 harboring Epsilonproteobacterial symbionts such as shrimps or polychetes, usually live at  
441 higher temperatures than those harboring Gammaproteobacteria, such as clams or  
442 vestimentiferans. Indeed, the two Provannidae gastropods, *Alviniconcha* spp. and *Ifremeria*  
443 *nautiliei*, studied by Urakawa, co-occur at the same sites in the Manus Basin, the former  
444 harboring Epsilonproteobacterial symbionts living at higher temperatures than *I. nautiliei*  
445 which harbors Gammaproteobacteria. This could be congruent with our example, as *C.*  
446 *naticoides* that lives on tubes of *Riftia pachyptila* may in fact live in a warmer microhabitat  
447 that the tubeworm itself and its Gammaproteobacterial endosymbionts, the latter being  
448 protected by the chitinous tube. *C. naticoides* lives in sympatry with another small gastropod,  
449 *Lepetodrilus elevatus* on the tube of *R. pachyptila*, where a vertical microzonation has been  
450 observed. Individuals of *C. naticoides* cluster at the base of the tubes, where temperatures up  
451 to 25°C were measured (Sarradin et al. 1998), whereas *L. elevatus* rather graze higher up the  
452 tubes (P. Tyler, pers. obs. cited in Mills et al. 2007), where temperatures ranged between 1.6  
453 and 10°C (Sarradin et al. 1998). So *C. naticoides* is associated with the warmer part of *R.*

454 *pachyptila* tube, and is also sometimes found among *A. pompejana* tubes (Desbruyères et al.  
455 2006; Mills et al. 2007), which live on the chimney walls at even higher temperatures (up to  
456 50°C was measured at 2-5 cm within the tube assemblages, Le Bris et al. 2005). Temperature  
457 can also be put forward to explain the different bacterial partners in *C. naticoides*  
458 (Epsilonproteobacteria ) et *L. fucensis* (Gammaproteobacteria ) ectosymbioses. Indeed  
459 *Lepetodrilus fucensis* was reported (Bates et al. 2005) to be abundant in fluids with  
460 temperature between 4 and 10°C, and to be absent where maximum fluid temperature reached  
461 18°C. Although precise temperatures have not been reported in literature, *C. naticoides* is  
462 probably exposed to temperatures exceeding 20°C in the habitats its occupies (base of *R.*  
463 *pachyptila* tubes or *A. pompejana* clumps).  
464 This selection of either Epsilon- or Gammaproteobacterial symbionts which seem to be  
465 affected by temperature, could also be linked to oxygen availability (both being negatively  
466 correlated). Sulfur metabolism pathways are indeed not the same in Epsilon- and  
467 Gammaproteobacteria. Both of the pathways used by deep-sea hydrothermal  
468 Gammaproteobacteria (the reverse sulfate reduction and the Sox multienzyme system )  
469 require O<sub>2</sub> as a terminal electron acceptor in most cases. This indicates that a relatively O<sub>2</sub>-  
470 depleted environment is less suitable for their growth (Yamamoto and Takai, 2011). Thus, it  
471 is predicted that the metabolically habitable niches for deep-sea chemoautotrophic  
472 Gammaproteobacteria strictly require co-existence of reduced sulfur compounds and O<sub>2</sub>.  
473 Besides oxygen, some Epsilonproteobacteria are also able to use sulfur compounds as  
474 electron acceptors (Yamamoto and Takai, 2011), which may allow them to tolerate and  
475 colonize O<sub>2</sub>-depleted and warmer niches within the mixing zone, closer to the reducing  
476 hydrothermal fluid.

477

478

## 479 **Conclusion**

480 *Cyathernia naticoides* harbors dense populations of filamentous Epsilonproteobacteria in its  
481 gill which may contribute to their nutrition through intracellular digestion by gill cells. OTU 3  
482 was identified as a probable candidate dominant gill bacterium. Yet, the diet could be  
483 mixotrophic, an additional food source being the bacteria grazed on *R. pachyptila* tubes.  
484 OTUs 1 and 11 identified here are likely siboglinid tube-associated Epsilonproteobacteria that  
485 may be significant food sources on this route. Novel for molluscs by the combination of the  
486 location (ectosymbionts) and bacterial phylotype (Epsilonproteobacteria) encountered and the  
487 feeding mechanism, the symbiosis of *C. naticoides* represents an unusual type of association  
488 in the already long list of molluscan symbioses, of which more await characterization in  
489 particular in smaller-sized species.

490

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497

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671 Ethical standards  
672 The authors declare that the experiments comply with the current laws of the country they  
673 were performed (France).  
674

675 The authors declare that they have no conflict of interest.

676

677

## 678 **Figure Legends**

679

680 **Fig. 1** *Cyathernia naticoides* ctenidium. **a)** Specimen outside its shell showing the extension  
681 of the ctenidium (ct). **b)** Semi-thin section of the ctenidium. **c)** Basal part of the gill filaments  
682 showing the accumulation of bacteria (b) between the gill filaments (gf) and in lysosome-like  
683 structure (ly). **d)** Close-up of bacteria in the inter-filament space (ifs) and in lysosome-like  
684 structure (ly). **e)** Transverse section through the gill, displaying bacterial filaments in white,  
685 hybridized with the FISH probe Arc-94. **f)** Transverse section through the gill displaying  
686 animal cells (DAPI-labelled nuclei in blue) and bacteria labelled with probes Eur-338 (Cy3,  
687 red) and Arc-94 (Cy5, green), overlying signals resulting in a yellow color. Scale bars : a = 1  
688 mm, b = 100  $\mu$ m, c = 20  $\mu$ m, d = 10  $\mu$ m, e = 50  $\mu$ m, f = 100  $\mu$ m

689 **Fig. 2** Transmission electron micrographs of the gill (A, B) of *C. naticoides*. **a)** Bacteria are  
690 endocytosed in the gill epithelium and **b)** progressively degraded. Scale bars : a, b = 2,5  $\mu$ m

691 **Fig 3** Phylogenetic tree based on the analysis of 16S rRNA-encoding gene sequences. The  
692 tree with the highest log likelihood (-13800) is shown. The percentage of trees in which the  
693 associated taxa clustered together is shown next to the branches. Scale bar represents the  
694 number of substitutions per site. Full (>1400 bp) sequences from this study appear in bold and  
695 an asterisk indicates an OTU that is present at both 9°50'N and 12°50'N. Sequences  
696 corresponding to confirmed metazoan symbionts or epibionts are underlined. All positions  
697 with less than 95% site coverage were eliminated. There were a total of 1302 positions in the  
698 final dataset, and partial sequences were excluded

699 **Fig. 4**  $\delta^{13}\text{C}$  and  $\delta^{15}\text{N}$  values of *Cyathernia naticoides* and other heterotrophic and symbiotic  
700 vent gastropods. Triangle up are isotopic ratios of *Cyathernia naticoides*, with dark blue  
701 symbols, those sampled at EPR 9°50'N (9°) and light blue symbols, those sampled at EPR  
702 12°50'N (13°). Triangle down, displayed previous isotopic values respectively, after correction  
703 using trophic step fraction of 1.0‰ for  $\delta^{13}\text{C}$  and 3.3‰ for  $\delta^{15}\text{N}$ . The rectangle with dashed  
704 lines represents the stable isotopes ratios of  $\epsilon$ -proteobacteria from Campbell et al. (2003).  
705  $\delta^{13}\text{C}$  and  $\delta^{15}\text{N}$  values of other gastropods are from: Levesque et al. (2006) for *Lepetodrilus*  
706 *fucencis*; Gaudron et al. (in revision) for *L. elevatus* (9° and 13°), *L. pustulosus*, *L. ovalis* and  
707 *Eulepetopsis vitrea*; Henry et al. (2008) for *Ifremeria nautilei*; Goffredi et al. (2004) for the  
708 Scaly snail; Beinart et al. (2013) for  $\delta^{13}\text{C}$  for *Alviniconcha hessleri* dominated by  
709  $\epsilon$ -proteobacteria (*Alviniconcha* (Epsilon)) and for *Alviniconcha hessleri* dominated by  
710  $\gamma$ -proteobacteria (*Alviniconcha* (Gamma)).  $\delta^{15}\text{N}$  of *Alviniconcha hessleri* is not documented,  
711 except for *Alviniconcha* sp. from Henry et al. (2008), which was used in this study

712 **Table 1** Number of sequences representing each identified OTU in each sample investigated  
713 in this study. OTU names in bold correspond to full length sequences included in the  
714 phylogeny, the suffix '-p' indicates a partial sequence. Affiliation based on best BLAST hits:  
715 Epsilonproteobacteria (E), Deltaproteobacteria (D), Mollicutes (M) and Bacteroides (B).  
716 Names of samples are indicated by site (9 or 12), specimen ID (1 to 3) and tissue type (Gi-  
717 gill, VM-visceral mass, Sh-shell) as follows: site-specimen-tissue. Sum of sequences per  
718 OTU and percentage of total sequence counts are indicated. Finally, the two bottom rows  
719 indicate whether the OTU has no mismatch (+), a single mismatch (1mis) or more (-) to FISH  
720 probes Arc-94 and Epsy-549

721 **Table 2:** analysis of fragments of functional genes encoding APS reductase (*aprA*) and ATP  
722 citrate Lyase (*ACL*), their length, the identity of the representative sequence, GENBANK  
723 accession number, percentage out of 70 (*aprA*) and 35 sequences (*ACL*), number of  
724 specimens in which the sequence occurred out of 3 (*aprA*) and 2 (*AclB*), tissue occurrence  
725 (G: gill, R: visceral mass), and best hit according to BLASTX translated nucleotide sequence  
726 analysis

727

728 **Supplementary material Fig S1:** Phylogenetic reconstruction based on a 100 aa-long  
729 fragment of the *aclB* gene. A Maximum Likelihood approach using the JTT matrix-based  
730 model and a discrete Gamma distribution of evolutionary rates with a proportion of invariant  
731 sites was used. All bacteria from the ingroup are Epsilonproteobacteria, and the tree is rooted  
732 on two Aquificales. Bootstrap percentage values based on 500 replicates are displayed. Scale  
733 bar corresponds to 10% estimated sequence divergence



Figure 1  
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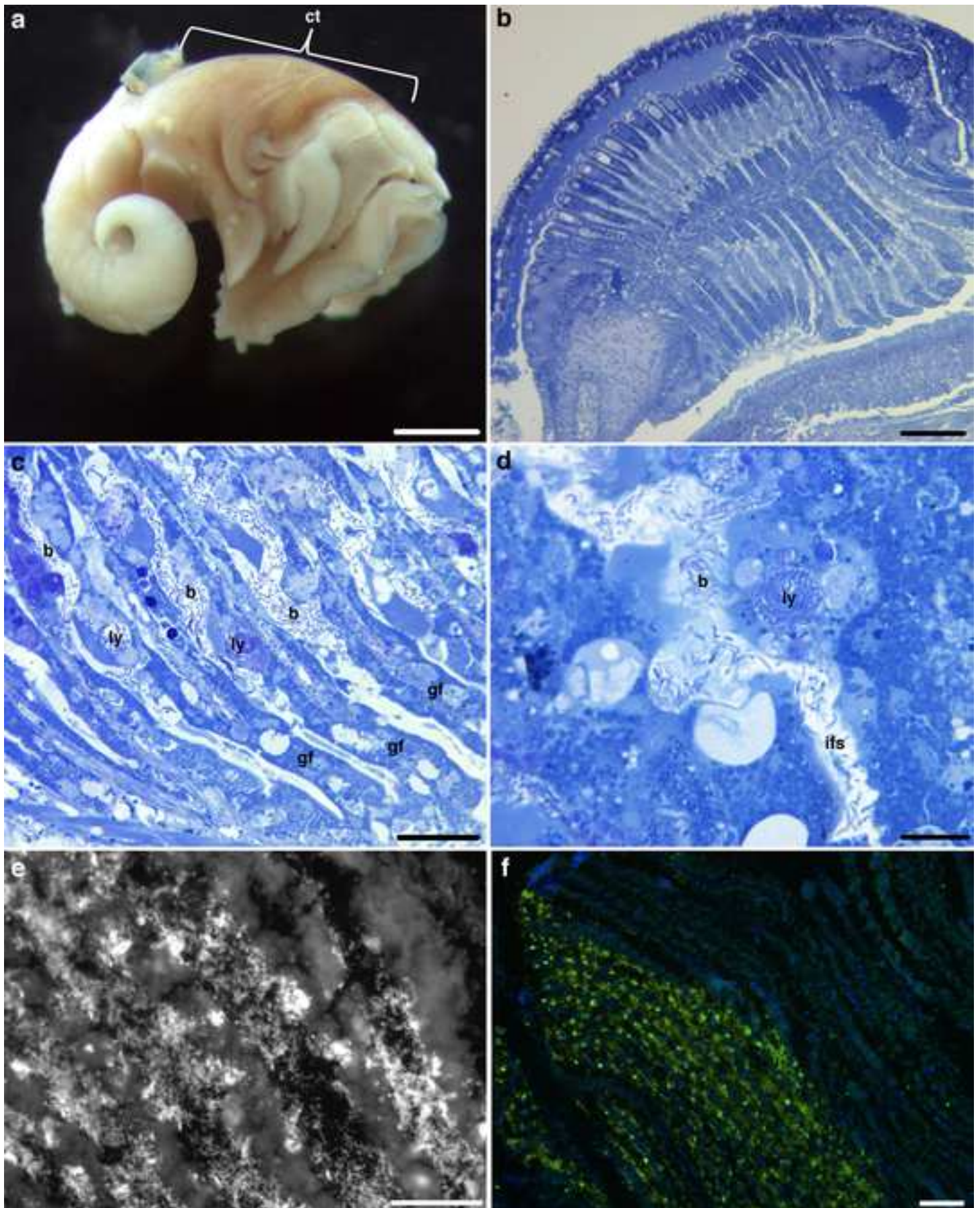


Figure 2  
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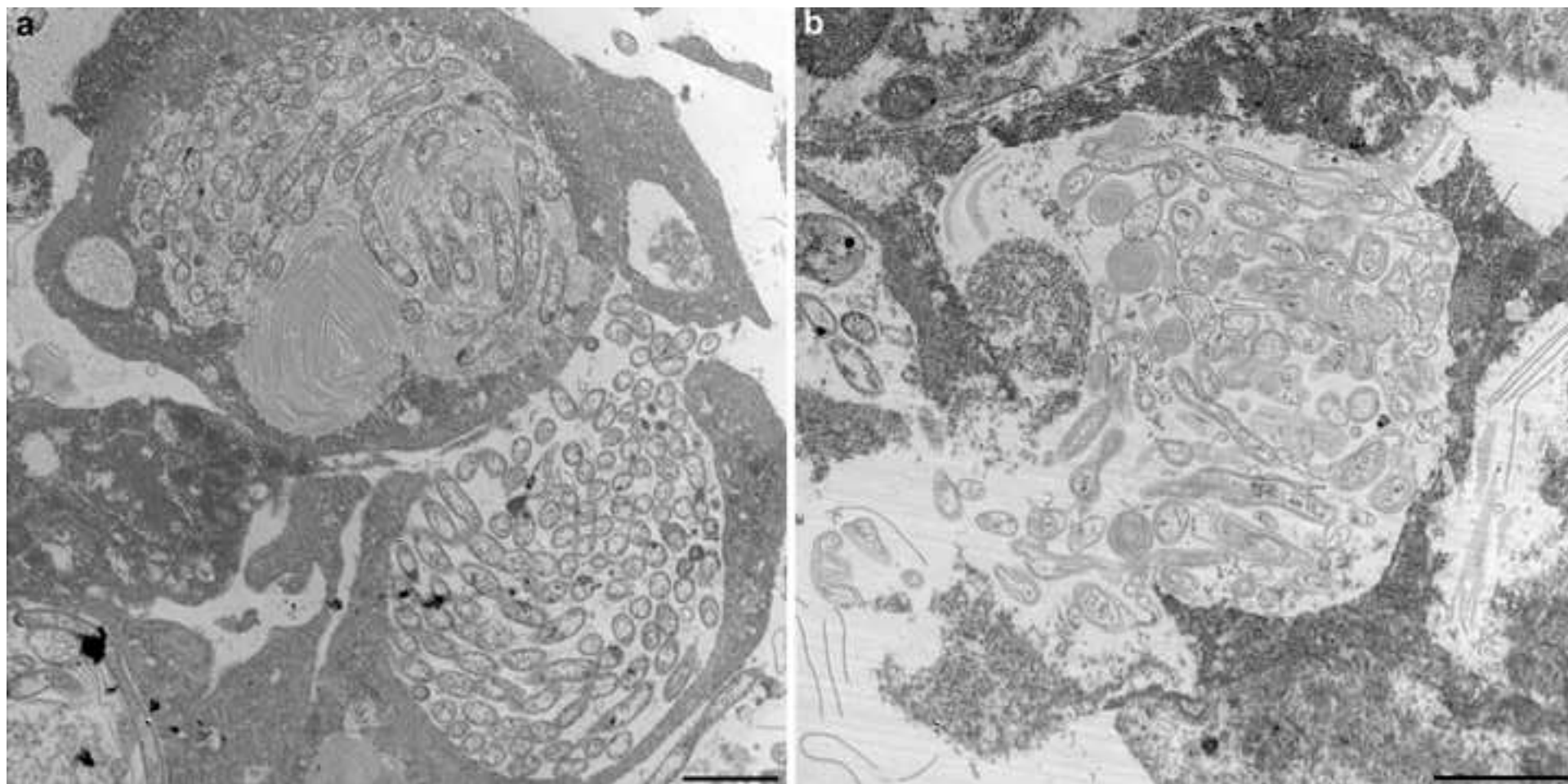




Figure 3

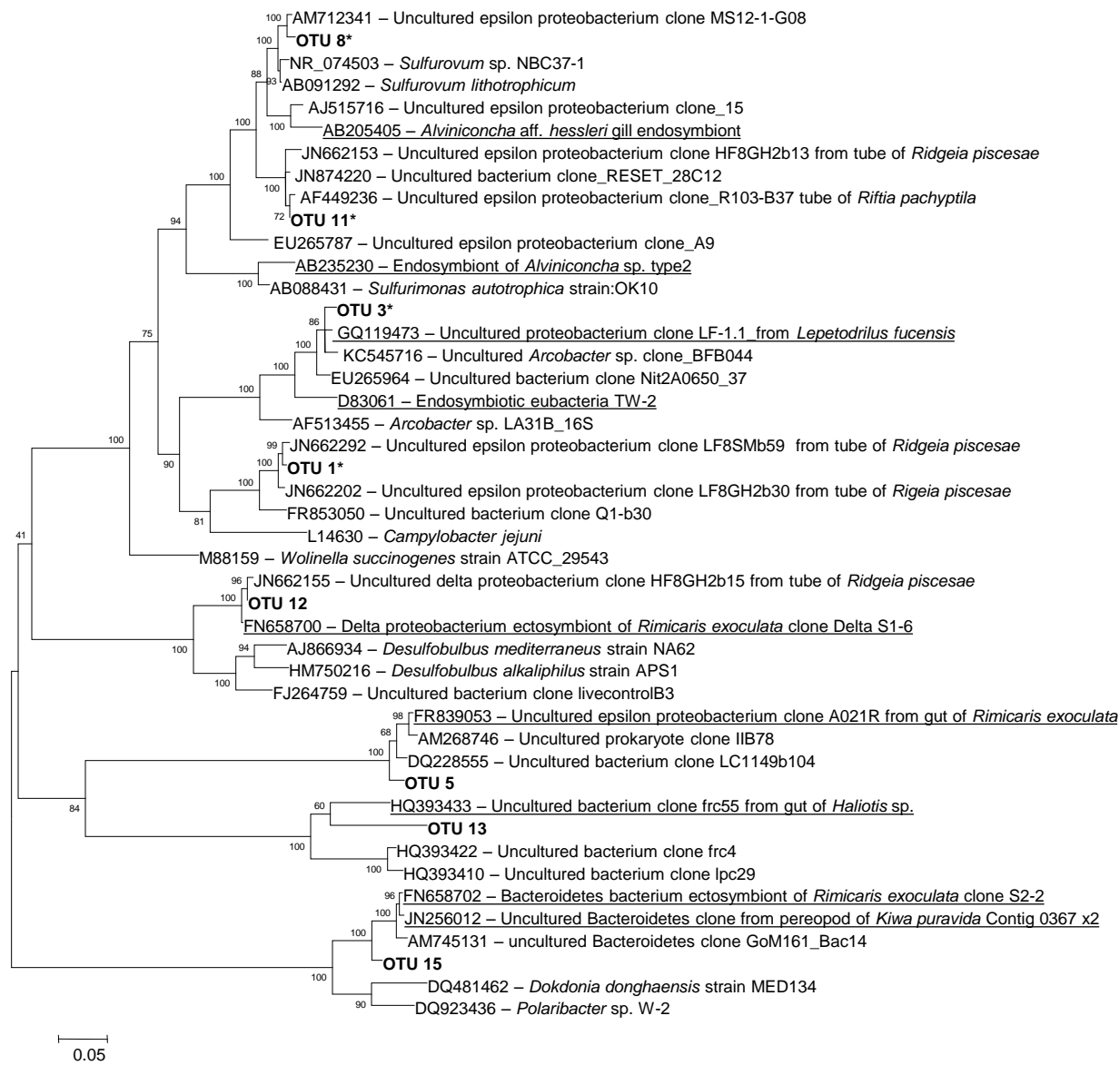






Table 1

<b>OTU ID</b>	<b>3</b>	<b>1</b>	<b>8</b>	<b>11</b>	<b>5</b>	<b>2-p</b>	<b>16-p</b>	<b>12</b>	<b>13</b>	<b>4-p</b>	<b>15</b>	<b>Sum</b>
<b>Accession</b>	KM213004	KM213002	KM213007	KM213008	KM213006	KM213003	KM213012	KM213009	KM213010	KM213005	KM213011	
<b>Affiliation</b>	E	E	E	E	E	E	E	D	M	M	B	
<b>9-1-Gi</b>	7	7			1	4				1		30
<b>9-1-VM</b>	3	20	3		1	3				1		32
<b>9-2-Gi</b>	14	7	3	1	5							32
<b>9-2-VM</b>	7	7	3		5			4	1		1	32
<b>9-2-Sh</b>		9	14	2				1			1	31
<b>9-3</b>	15	6	2		1		1	1	2	1		32
<b>12-1-Gi</b>	11											11
<b>12-1-VM</b>			11	7								19
<b>12-2-Gi</b>	5		5	2								17
<b>12-2-VM</b>		2	3	3								13
<b>12-3-Gi</b>	6		2	1			3					17
<b>12-3-VM</b>		1	11	5			1					29
<b>Sum per OTU</b>	68	59	57	21	13	7	5	6	3	3	2	244
<b>Percentage</b>	23,05	20	19,32	7,12	4,41	2,37	1,69	2,03	1,02	1,02	0,68	82,71
<b>Arc-94</b>	+	-	+	+	-	1 mis	+	-	-	-	-	
<b>Epsy-549</b>	+	1 mis	+	+	-	+	+	-	-	-	-	

Table 2

Fragment	Approx length	Clone ID	Accession number	%/total	Specimen occurrence	Tissue occurrence	Best BLAST hit (BlastX)
aprA	365 nt	761	KP115589	40.0	3	R	96% EU265804 Epibiont of the vent crab <i>Kiwa hirsuta</i> (Gammaproteobacteria)
		843	KP115590	24.3	2	R	91% GU197406 Bacterium associated with the Oligochete <i>Tubificoides benedii</i> (Gammaproteobacteria)
		144	KP115591	10.0	2	R	90% GU197406 Bacterium associated with the Oligochete <i>Tubificoides benedii</i> (Gammaproteobacteria)
		820	KP115592	18.6	2	R	100% FM165456 Bacterium associated with the tube of <i>Lamellibrachia anaximandri</i> (Gammaproteobacteria)
		786	KP115593	4.3	1	R	96% EF633097 Bacterium associated with <i>Echinocardium cordatum</i> (Deltaproteobacteria)
		827	KP115594	1.4	1	R	97% AM234053 <i>Olavius algarvensis</i> Delta-4 endosymbiont (Deltaproteobacteria)
		765	KP115581	54.3	2	G	98% FN659794 bacterium from branchial chamber of <i>Rimicaris exoculata</i> (Epsilonproteobacteria)
AclB	305nt	782	KP115582	11.4		G, R	98% FN908920 bacterium from hydrothermal fluid, Clueless (Epsilonproteobacteria)
		847	KP115583	2.9	1	R	99% FR670537 bacterium from Lucky Strike (Epsilonproteobacteria)
		766	KP115584	2.9	1	R	98% FN659786 branchial chamber of <i>Rimicaris exoculata</i> (Epsilonproteobacteria)
		805	KP115585	2.9	1	R	98% FN908920 bacterium from hydrothermal fluid, Clueless (Epsilonproteobacteria)
		808	KP115586	17.1		R, G	99% FN562694 bacterium from the Irina II vent, Logatchev (Epsilonproteobacteria)
		840	KP115587	2.9	1	R	99% FN908925 Bacterium from the Logatchev vent field (Epsilonproteobacteria)
		163	KP115588	5.8	1	R	97% FN562694 bacterium from the Irina II vent, Logatchev (Epsilonproteobacteria)