INTRODUCTION

Research at deep-sea hydrothermal vents (discovered only 25 yr ago) has resulted in a radically altered perception of life in the aphotic zone of the world’s oceans. Unrivaled biomass, coupled with extreme geological and chemical stochasticity, has redefined ideas regarding not only deep-sea biology, but the nature and function of ecosystems. Completely removed from sunlight, primary productivity at hydrothermal vents is driven by chemical energy. Life in these novel habitats is characterized by an astounding array of physiological, morphological and behavioral adaptations. The hallmark of hydrothermal vent biology is the recurring theme of chemoautotrophic bacterial symbioses, which have been described for the majority of the dominant vent macrofauna (Fisher 1990, Cavanaugh 1994, Nelson & Fisher 1995). The bacterial symbionts harness energy from reduced inorganic sulfur compounds to drive carbon fixation and, ultimately, either directly or indirectly provide their host with organic carbon. The host organism is able to use a combination of traits to sequester and transport necessary substrates to the aerobic chemoautotrophic bacteria.

As hydrothermal vent exploration has progressed, and additional locales have been identified and sampled, many key evolutionary questions regarding the widespread nature of chemoautotrophic symbioses have arisen. In particular, biogeographic ‘provinces’ are under continuous evaluation as more sites are explored (e.g. Tunnicliffe et al. 1998). The recent discoveries of the Kairei and Edmond hydrothermal vent fields on the Central Indian Ridge (CIR) were of profound importance in the context of further resolving...
biogeographic patterns in the world’s hydrothermal vent ecosystems (Hashimoto et al. 2001, Van Dover et al. 2002). The biology of CIR vents was a great mystery, given the disparate communities inhabiting the Atlantic and Pacific vents. Forming a key physical link between the Mid-Atlantic Ridge (MAR) hydrothermal vents and the western Pacific back arc basins, the CIR hydrothermal vents were found to be characterized by a blend of Atlantic and Pacific faunas (Van Dover et al. 2001). Massive swarms of shrimp dominated the vent fields, visually reminiscent of MAR vents. However, much like western Pacific vent sites, sea anemones, interspersed with clumps of mussels and snails, dominated the benthic environment.

The CIR mussels, designated *Bathymodiolus* aff. *brevior*, are more closely related to the western Pacific mussel *B. brevior* than to the MAR mussels *B. azoricus* and *B. puteoserpentis*, as revealed by molecular studies based on NADH dehydrogenase subunit 4 (ND4) (Van Dover et al. 2001, Won et al. 2002). Yamanaka et al. (2003) reported transmission electron microscope (TEM) and sulfur-isotope data that indicate the presence of bacterial symbionts in this species. This mussel has also been described as *B. marisindicus* based on morphological data (Hashimoto 2001); however, because morphological characters may reflect phenotypic plasticity, a new species designation may not be warranted. Pending additional data indicating that the CIR mussel is a separate species from *B. brevior*, this mussel will be referred to as *B. aff. brevior* herein, since the molecular results indicate high genetic identity with *B. brevior*.

While the CIR vents lack the well-studied vesicomyid clam and vestimentiferan tubeworm symbioses, the presence of mytilid mussels at this locale is significant given their ubiquity in the Atlantic and Pacific Oceans. Among symbiont-hosting vent fauna, only mussels have been described from all known hydrothermal vent biogeographic provinces (Table 1): Juan de Fuca (unnamed new species), East Pacific Rise, EPR, (*Bathymodiolus thermophilus*), MAR (*B. azoricus* and *B. puteoserpentis*), western Pacific (*B. brevior* and *B. elongatus*), and Okinawa Trough (*B. japonicus*, *B. platifrons*, and *B. septemdierum*). Thus, finding mytilid mussels at the CIR vents, although not unexpected, further establishes their global distribution. In addition to their ubiquity, *Bathymodiolus* spp. mussels are unique in the range of symbiont composition observed in the gills of different species (Table 1). To date, mussels are known from hydrothermal vents and cold seeps that host either sulfur-oxidizing chemosynthetic symbionts (Cavanaugh 1983, Nelson et al. 1995, Dubilier et al. 1998) or methane-oxidizing (methanotrophic) symbionts (Childress et al. 1986). Furthermore, mussels that host ‘dual symbioses’ involving both types of bacteria are known from the Gulf of Mexico cold seeps (Cavanaugh et al. 1987, Cavanaugh 1993, Fisher et al.

Table 1. *Bathymodiolus* spp. Symbiont composition and geographic distribution of mussels from deep-sea hydrothermal vents.

<table>
<thead>
<tr>
<th>Species</th>
<th>Symbiont type</th>
<th>Location</th>
<th>Depth (m)</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indian Ocean</td>
<td><em>Bathymodiolus aff. brevior</em></td>
<td>C</td>
<td>Central Indian Ridge</td>
<td>2450–3300</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Lau Basin</td>
<td>1850</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Galapagos Rift</td>
<td>2500</td>
</tr>
<tr>
<td></td>
<td><em>Bathymodiolus n. sp.</em></td>
<td>C</td>
<td>Juan de Fuca</td>
<td>2500</td>
</tr>
</tbody>
</table>

*aSingle specimen was collected from Juan de Fuca in August 1999; host taxonomy is unresolved*
1993) and the MAR hydrothermal vents (Cavanaugh et al. 1987, 1992, Cavanaugh 1993, Fisher et al. 1993, Distel et al. 1995, Robinson et al. 1998, Fiala-Médioni et al. 2002). Remarkably, the dual symbioses in these mytilids represent the only known example of the stable coexistence of 2 physiologically and phylogenetically distinct bacteria within the same eukaryotic host cell. Thus, the discovery of a new species of mussel provides potential evolutionary clues to the evolution of the neighboring MAR dual symbiosis species.

Here, the presence and nature of symbionts in the CIR mussel, *Bathymodiolus* aff. *brevior*, is evaluated using ultrastructural, enzymatic and molecular methods. The phylogenetic affinities of the putative symbiont are assessed using comparative sequence analyses including other known bivalve symbionts. We present evidence supporting the presence of a single, chemoautotrophic symbiont population in *B. aff. brevior*. Furthermore, the 16S rRNA phylotype of the *B. aff. brevior* symbiont is more closely related to the vesicomyid clam symbionts than the mytilid mussel chemoautotrophic symbionts.

**MATERIALS AND METHODS**

**Organisms.** *Bathymodiolus* aff. *brevior* specimens were collected in April 2001 by the remotely operated vehicle, ROV ‘Jason’ at the Kairei (25° 19′ S, 70° 02′ E; 2415 to 2460 m) and Edmond (23° 52′ S, 69° 35′ E; 3290 to 3320 m) vent fields on the CIR. The mussels were transported to the surface in insulated containers within 24 h of collection, and immediately transferred to chilled seawater (4°C). Specimens were dissected and tissues were preserved for microscopy or frozen in liquid nitrogen for enzymatic and molecular studies. The latter samples were stored at −70°C on board ship, then transported to the laboratory on dry ice and stored at −80°C.

**Microscopy.** Pieces of *Bathymodiolus* aff. *brevior* tissue were dissected from the middle portion of the gill on board the ship (RV ‘Knorr’) and preserved in 3% glutaraldehyde in 0.1 M sodium cacodylate, 0.4 M sodium chloride buffer, pH 7.4. Upon return to the laboratory, gill filaments were dehydrated through a graded ethanol series, post-fixed in osmium tetroxide, stained en bloc with uranyl acetate, and embedded in Spurr’s resin. Thick sections of gill filaments were examined from 2 individuals from Kairei and 1 from Edmond. Thin sections of gill filaments were examined from 2 individuals from Kairei mussel with a Zeiss 10CA transmission electron microscope.

**Enzyme assays.** Cell free extracts of *Bathymodiolus* aff. *brevior* gill tissue (n = 3) were analyzed for activities of enzymes diagnostic of carbon fixation (ribulose 1,5-bisphosphate carboxylase/oxygenase, RubisCO) and methylootrophy (methanol dehydrogenase). RubisCO activity was assayed using the 14C incorporation method as described in Robinson et al. (1998), with fresh spinach used as the positive control. Methanol dehydrogenase activity was tested using the spectrophotometric method of Anthony & Zatzman (1964). Gill tissue of the MAR mussel *B. azoricus*, which is known to host methanotrophic symbionts (Fiala-Médioni et al. 2002), was used as the positive control. For both assays, boiled cell free extracts and samples lacking substrate, ribulose 1,5-bisphosphate and methanol, respectively, were used as negative controls.

**PCR, cloning, and sequencing of 16S rRNA.** DNA was extracted from mussel gill tissue (n = 4) using the DNeasy tissue kit (Qiagen). The universal bacterial primers 27f and 1492r (Weisburg et al. 1991) were used to amplify 16S rRNA from gill DNA for *Bathymodiolus* aff. *brevior* from both Kairei (n = 2) and Edmond (n = 2) fields. Reactions included primers (1 µM each), 0.25 mM deoxynucleoside triphosphates, 2.5 mM MgCl₂, 1 U *Taq* polymerase, 1× PCR buffer and 0.01 to 1 ng DNA. PCR was performed using an initial 2 min denaturation at 94°C, followed by 35 cycles at 94°C for 1 min, 50°C for 1 min, and 74°C for 2 min, and concluding with 74°C for 10 min. PCR products were cloned using the TOPO TA cloning kit (Invitrogen). DNA was extracted from positive clones using the QIAprep Spin Miniprep kit (Qiagen) and screened using *RsaI* digests to discriminate between inserts. Partial (~700 bp) single strand sequences were obtained for all positive candidates. For a subset of these positive clones, the entire 16S rRNA insert was sequenced in both directions and was re-sequenced in cases of ambiguity. All sequencing reactions were carried out on an ABI 3100 sequencer (Applied Biosystems) and reactions used either M13 vector primers or universal bacteria 16S rRNA primers (27f, 530f, 1492r; Weisburg et al. 1991).

**Confirmation of symbiont 16S rRNA sequence.** RT-PCR was used to confirm that the bacterial 16S rRNA sequence obtained was not a contaminant and, in fact, was from the symbiont of *Bathymodiolus aff. brevior*. Since the presence of RNA is indicative of active cells, a sequence generated from RNA, via cDNA, supports the presence of a viable population of that phylotype in the RNA source. Total RNA, extracted from the symbiont-containing gill tissue of 1 mussel from each CIR vent field using the RNeasy Mini Kit (Qiagen), was reverse transcribed using the Thermoscript™ RT-PCR System (Invitrogen). The 16S rRNA gene was then amplified as described above, using the 27f and 1492r universal primers, and directly sequenced, using the following internal primers: 530f (universal), 42f GATTGAACGCTGGCGG, 515r CCGCGGCTGCTGGCAC, 1473r TTACCCCAGACCATCGACACACC (all numbering based on *Escherichia coli*).
Phylogenetic analyses. Sequence analyses were performed using the Genetics Computer Group (GCG) software package (Wisconsin Package) and PAUP 4.0 b10 (Swofford 2002). In GCG, sequences were aligned by Pileup and then manually edited based on the 16S rRNA secondary structure of *Escherichia coli* (Boros et al. 1979). Phylogenetic analyses were conducted using PAUP 4.0 b10. Maximum parsimony trees were generated with heuristic searches using random-sequence addition with 100 replicates and TBR branch-swapping methods. The same search conditions were used for a 500 replicate bootstrap analysis. Analyses were based on approximately 1.5 kb section of 16S rRNA; gaps were treated as missing data. The organisms included in the phylogenetic analyses are listed in Table 2. The 16S rRNA sequence of the *Bathymodiolus aff. brevior* symbiont has been deposited in GenBank under Accession No. DQ077891.

**RESULTS**

**Microscopy**

Transmission electron micrographs revealed Gram negative bacteria enclosed in vacuoles within *Bathymodiolus aff. brevior* gill epithelial cells (Fig. 1). The symbiont-containing bacteriocytes occurred between symbiont-free intercalary cells. Symbionts appeared in rod and coccoid shapes and lacked the intracytoplasmic membranes typical of methanotrophic bacteria. Multiple symbionts occurred within a single vacuole and each bacteriocyte included multiple vacuoles. These vacuoles tend to be positioned at the apical end of host cells, which bear microvilli and are exposed to seawater.

**Enzyme assays**

RubisCO activity was detected in cell free extracts of *Bathymodiolus aff. brevior* gill tissue from 3 specimens (0.007 ± 0.003 nmol min⁻¹ mg⁻¹ protein), while methanol dehydrogenase activity was not detected. For both assays, activity was detected in the respective positive controls, spinach and *B. azoricus*, and was not detected in boiled cell free extracts and substrate-free reactions.

<table>
<thead>
<tr>
<th>Host group</th>
<th>Species</th>
<th>GenBank Accession No.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Chemoautotrophic symbionts</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bivalvia</td>
<td><em>Bathymodiolus</em> aff. <em>brevior</em> symbiont</td>
<td>DQ077891</td>
</tr>
<tr>
<td></td>
<td><em>Bathymodiolus puteoserpentis</em> symbiont 1</td>
<td>U29163</td>
</tr>
<tr>
<td></td>
<td><em>Bathymodiolus thermophilus</em> symbiont</td>
<td>M99445</td>
</tr>
<tr>
<td>Solemyidae</td>
<td><em>Solemya velum</em> symbiont</td>
<td>M90415</td>
</tr>
<tr>
<td>Lucinidae</td>
<td><em>Lucinoma aequizonata</em> symbiont</td>
<td>M99448</td>
</tr>
<tr>
<td>Vesicomidae</td>
<td><em>Calyptogena elongata</em> symbiont</td>
<td>AF035719</td>
</tr>
<tr>
<td></td>
<td><em>Calyptogena magnifica</em> symbiont</td>
<td>AF035721</td>
</tr>
<tr>
<td></td>
<td><em>Calyptogena pacifica</em> symbiont</td>
<td>AF035723</td>
</tr>
<tr>
<td></td>
<td><em>Vesicomya gigas</em> symbiont</td>
<td>AF035726</td>
</tr>
<tr>
<td>Vestimentifera</td>
<td><em>Riftia pachyptila</em> symbiont</td>
<td>M99451</td>
</tr>
<tr>
<td><strong>Methanotrophic symbionts</strong></td>
<td></td>
<td></td>
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<tr>
<td>Bivalvia</td>
<td><em>Bathymodiolus</em> childressi symbiont</td>
<td>U05595</td>
</tr>
<tr>
<td></td>
<td><em>Bathymodiolus puteoserpentis</em> symbiont 2</td>
<td>U29164</td>
</tr>
<tr>
<td>Free-living bacteria</td>
<td><em>Escherichia coli</em></td>
<td>AF233451</td>
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<tr>
<td></td>
<td><em>Thiomicrospira crunogena</em></td>
<td>L40810</td>
</tr>
<tr>
<td></td>
<td><em>Thiomicrospira thyasirae</em></td>
<td>AF016046</td>
</tr>
</tbody>
</table>

Fig. 1. *Bathymodiolus aff. brevior*. Transmission electron micrograph of transverse section through gill filament showing bacterial symbionts, showing bacteriocyte (B) next to symbiont-free intercalary cell (IC). Symbionts (S) are located within vacuoles (V) in bacteriocyte. Putative lysosomal bodies (L) are located in distal portion of bacteriocyte. Scale bar = 5 µm.
Sequence analyses

PCR from the 4 samples using the 27f and 1492r Bacteria primers yielded a band that was the expected 1.5 kb in length. Each PCR was then cloned and 10 positive clones from each mussel were assessed. Restriction endonuclease analyses showed a dominant band pattern for 36 of 40 positive clones. The 36 positive clones were then screened using single-strand sequencing, and 30 of those clones yielded the same 16S rRNA sequence (<0.1% variation). For 2 mussels from the Kairei site, 2 clones each were fully sequenced in both directions, resulting in nearly identical 1473 bp 16S rRNA sequences; there were 2 bp substitutions in 1 clone as compared to the other 3, which were identical.

Subsequent sequence confirmation of the symbiont phylotype in host tissue was accomplished via RT-PCR and cDNA direct sequencing of 1 mussel from Kairei and 1 mussel from Edmond. The fully sequenced 16S rRNA RT-PCR products for both mussels were 100% identical to the three indistinguishable fully sequenced clones. This ‘consensus’ sequence was designated as the Bathymodiolus aff. brevier symbiont phylotype.

Phylogenetic analyses yielded 7 most parsimonious trees that all placed the symbiont of Bathymodiolus aff. brevier with the mytilid mussel and vesicomyid clam chemoautotrophic symbionts (Fig. 2). This clade, containing both mytilid mussel and vesicomyid clam symbionts, had 100% bootstrap support. The mytilid mussel methanotrophic symbionts fell in a separate clade, with 100% bootstrap support (as shown previously: Distel & Cavanaugh 1994); this was distinct from the clade containing the chemoautotrophic mussel symbionts, which included the B. aff. brevier symbiont. Interestingly, the B. aff. brevier symbiont groups with the vesicomyid clam symbionts (supported by a 97% bootstrap value) rather than clustering with the mytilid mussel chemoautotrophic symbiont group (82% bootstrap support). Additional phylogenetic analyses were conducted using different proportions of vesicomyid clam and mytilid mussel symbionts to evaluate the stability of this topology with respect to taxon sampling. All resulting phylogenies showed that the B. aff. brevier symbiont clustered with the vesicomyid clam symbionts, sister to the other mytilid mussel chemoautotrophic symbionts (results not shown).

DISCUSSION

Ultrastructural, physiological and molecular evidence all support the existence of a single population of chemoautotrophic symbiont in the mussel Bathymodiolus aff. brevier from the recently discovered CIR hydrothermal vents. Indeed, the western Pacific mussel B. brevier, which is most closely related to the CIR mussel, has been characterized as having only chemoautotrophic symbionts (Dubilier et al. 1998). In addition, a recent study of this mussel, referred to as B. marisindicus, reported TEM and stable sulfur isotope data that also are consistent with the presence of chemoautotrophic symbionts (Yamanaka et al. 2003).

In TEMs, a single endosymbiont morphotype was observed in Bathymodiolus aff. brevier gill epithelial cells; while both rod and coccolid shapes were observed, this may be an
artifact of symbiont orientation within the vacuole. This finding is in agreement with previously published TEMs for these mussels (Yamanaka et al. 2003). In addition, as seen in the chemosynthetic endosymbionts observed in *B. brevior* (Dubilier et al. 1998) and *B. thermophilus* (Le Pennec 1984, Fiala-Médioni et al. 1986), these bacteria lack the intracytoplasmic membranes typical of methanotrophic bacteria.

The Rubisco activity, albeit low in these mussels, is comparable to previously reported values for vent mytilids hosting chemosynthetic symbionts (Fisher et al. 1993, Nelson et al. 1995, Fiala-Médioni et al. 2002). Rubisco activity is notoriously difficult to measure in mytilid gill tissues, given their high levels of proteolytic activity (Nelson et al. 1995). In fact, for *Bathymodiolus puteoserpentis* (the MAR mussel which hosts both types of symbiont), Rubisco activity was undetectable (Cavanaugh et al. 1992), although immunological blots later confirmed its presence (Robinson et al. 1998). Consistent with the lack of observable methanotrophic endosymbionts in the TEMs and the single 16S rRNA sequence detected, methanol dehydrogenase activity was absent. Furthermore, this finding of a single population of chemosynthetic symbionts is consistent with the previously published stable carbon isotope value for *B. aff. brevior* mantle tissue of −29.9 ± 0.9‰ (Van Dover 2002).

Also in line with a single symbiont population, only one 16S rRNA phylotype was amplified and sequenced from mussels collected from both CIR vent fields. Sequence confirmation via RT-PCR indicated that this phylotype is the endosymbiont of *Bathymodiolus* aff. *brevior*. This additional step is critical in establishing the origin of the sequence in host tissue and ruling out the possibility that it is a contaminant.

The lack of variation in the 16S rRNA gene sequences obtained from multiple clones isolated from multiple individuals, is notable, though the implications in the context of symbiont transmission are unclear. Cary et al. (1993) reported preliminary work indicating maternal transmission in the chemosynthetic symbiont of the EPR mussel *Bathymodiolus thermophilus*. Conversely, recent evidence based on host and symbiont sequence variation in a hybrid zone, suggests environmental transmission of the chemosynthetic symbiont of the MAR mussel *B. azoricus*, which is known to host a dual symbiosis (Won et al. 2003). Won et al.’s (2003) report also featured ultrastructural evidence for environmental uptake of symbionts. However, these data may have been confounded by methodological artifacts, and maternal transmission cannot be eliminated as a possibility. A phylogenetic approach was recently used to address cospeciation (and indirectly symbiont transmission) in the bathymodioline mussel symbioses. Analyses of host and symbiont phylogenetic congruence provided weak support for cospeciation of chemosynthetic symbionts, but no evidence for cospeciation between the mussels and their methanotrophic symbionts, although small samples sizes may have biased these results (McKiness 2004). Future work aimed at evaluating mussel larvae and gametes for the presence of symbionts is necessary in order to establish the mode of symbiont transmission in these mussels. In addition, studies of genetic variation in both symbiont and host populations, as well as investigations of phylogenetic congruence of larger samples of host and symbiont, will allow a more robust assessment of cospeciation in bathymodioline mussel symbioses.

From an evolutionary perspective, the position of the *Bathymodiolus* aff. *brevior* symbiont within the vesicomyid clade is striking. This topology may reflect a shared evolutionary history for the mytilid mussel and vesicomyid clam symbioses. In a thoughtful discussion on the evolution of the mussel and vesicomyid clam symbioses, Distel (1998) notes that, based on the fossil record, vesicomyids evolved much more recently (95 to 135 million yr ago) than the mytilid mussel’s symbioses (150 million yr ago). This timing is coincident with the divergence of the mussel and vesicomyid clam symbionts, presumed to have occurred 125 to 300 million yr ago, based on a molecular clock estimate for 16S rRNA (Distel 1998). Thus, the question as to the nature of the origin of chemosynthetic symbioses characteristic of mytilid mussels and vesicomyid clams arises. Was the initial symbiosis in both mytilid mussels and vesicomyid clams with the same symbiont, which subsequently cospeciated with the respective host groups?

While the *Bathymodiolus* aff. *brevior* symbiont provides a potential link between these 2 host groups, there is a significant difference between the symbionts of the mussels and vesicomyid clams. Sequence data for the Rubisco gene, coupled with Western blots, show that the EPR vent mussel *B. thermophilus* has symbionts expressing a Form I Rubisco while the vent vesicomyid clam *Calyptogena magnifica* hosts symbionts with a Form II Rubisco (Cavanaugh & Robinson 1996). The Form I and Form II Rubisco enzymes are biochemically, physiologically and evolutionarily distinct (Cavanaugh & Robinson 1996, Tabita 1999), and the fact that bacteria which are closely related on the basis of 16S rRNA and which also inhabit similar environments express different forms of this enzyme is puzzling. While the Form I Rubisco is known to exhibit high CO₂ affinity, the Form II enzyme shows low CO₂ affinity, consistent with growth in high-CO₂ conditions (Jordan & Ogren 1981, Tabita 1988). Interestingly, there are examples of free-living bacteria that express both forms of Rubisco, such as the vent isolate *Thiomicospira* L-12 (C. Cavanaugh unpubl. data) and
Hydrogenovibrio marinus (Yaguchi et al. 1994). Thus, there is the potential for an ancestral symbiont with both Form I and Form II RubisCOs which gave rise to the mussel and clam symbionts with subsequent differential retention of the enzyme. Moreover, the endosymbionts probably experience very disparate conditions in the gills of their respective hosts, as mussels and vesicomyid clams have not only different CO₂ concentrations, but also different O₂ concentrations, given abundant hemoglobin present in vesicomyid clam blood (Arp et al. 1984). In addition, there are substantial differences in gill morphology and ciliation between filibranch mussels and eulamellibranch clams (Atkins 1937).

Evaluation of the Bathymodiolus aff. brevior symbiosis within the context of the chemical environment of the habitat is consistent with chemosymbiotic, as opposed to methanotrophic, symbionts. Preliminary analyses revealed high levels of sulfide dissolved in hydrothermal fluid sampled from both CIR vent fields (Van Dover et al. 2001). End member fluid methane concentrations reported for both Kairei and Edmond (0.2 and 0.4 mmol kg⁻¹, respectively) are very low compared to other hydrothermal vents (Van Dover et al. 2001), perhaps selecting against the presence of methanotrophic symbioses. Alternatively, methanotrophic symbionts may be present but escaped detection because of low abundance.

The prospect of a dual symbiosis is particularly compelling in the context of establishing the evolutionary origin of this phenomenon within the mytilid mussels. Future work characterizing the ancestral state of symbiosis in these hosts will be necessary in order to evaluate whether there was a single origin for dual symbioses. Coupled with forthcoming results of a host phylogeny, these data will provide critical insight into the evaluation of the evolution of mytilid dual symbioses.

This opportunity to study this mytilid–chemoautotrophic symbiosis was exciting given the key biogeographic location of the host Bathymodiolus aff. brevior as the previously unexplored CIR. These vents provide a physical link between the Atlantic and Pacific Oceans, evoking questions regarding the origin of the taxa at the Kairei and Edmond vent fields. Interestingly, this mussel appears to share evolutionary alliances with the western Pacific mussel B. brevior and not the mussels of the MAR, which hosts dual symbioses. Furthermore, its symbiont shows phylogenetic affinities to the vesicomyid clam symbionts, evoking questions regarding the origin of chemoautotrophic symbioses in the vesicomyid clams and mytilid mussels. By increasing the taxonomic sampling, patterns in the evolution of these symbionts can be more accurately assessed. Evaluation of the degree of genetic heterogeneity within the Kairei and Edmond vent mussels will provide a context within which to address the apparent lack of variation within and between the symbiont populations as well as, ultimately, the role of cospeciation in the evolution of this interaction. This finding opens the door for further questions regarding the evolutionary origin of symbioses in deep-sea mussels, as well as the nature of symbiont transmission and the role of cospeciation in the development of this interaction.

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