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# Free-Living Tube Worm Endosymbionts Found at Deep-Sea Vents<sup>∇†</sup>

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**Recent evidence suggests that deep-sea vestimentiferan tube worms acquire their endosymbiotic bacteria from the environment each generation; thus, free-living symbionts should exist. Here, free-living tube worm symbiont phylotypes were detected in vent seawater and in biofilms at multiple deep-sea vent habitats by PCR amplification, DNA sequence analysis, and fluorescence in situ hybridization. These findings support environmental transmission as a means of symbiont acquisition for deep-sea tube worms.**

The mode by which symbionts are passed between successive host generations is a primary question in symbiosis research. Symbiont transmission typically occurs vertically via transfer from parent to offspring, horizontally between cooccurring host individuals, or environmentally via uptake from a free-living population (3). Determining which of these mechanisms operates within a symbiosis is critical, as the transmission mode impacts fundamental ecological and evolutionary processes, including genome evolution, symbiont-host specificity, and coevolution (for examples, see references 7, 19, and 30).

Deep-sea vestimentiferan tube worms, which dominate the fauna at hydrothermal vents and cold seeps, are hypothesized to acquire their bacterial symbionts environmentally from a free-living population. Attempts to detect tube worm symbionts in host eggs and larvae by the use of microscopy and PCR have been unsuccessful (4–6, 12), suggesting that transmission does not occur vertically. Furthermore, most vent vestimentiferan species host symbionts that share identical 16S rRNA sequences, which is also consistent with the hypothesis of environmental transmission (14, 22). Unlike adults, the larvae and small juveniles of vestimentiferan tube worms have a mouth and gut, suggesting environmental acquisition via the ingestion of symbionts during larval development (12, 28). However, Nussbaumer et al. (23) recently demonstrated that bacterial symbionts are found on the developing tubes of settled larvae, entering the host worm through the epidermis and body wall of both larvae and young juveniles (23). These studies strongly suggest that tube worms acquire their symbionts from the surrounding environment and, therefore, that these endosymbionts should be detectable in a free-living form.

**Sample collection.** A systematic search for the free-living counterpart to the gammaproteobacterial endosymbiont phylotype shared by three species of vestimentiferan tube worms, *Riftia pachyptila*, *Oasisia alvinae*, and *Tevnia jerichonana*, was conducted at the Tica hydrothermal vent site (~2,600-m depth) on the East Pacific Rise (EPR) (9°50.447'N, 104°17.493'W) during December 2002 and December 2003. Symbiont-containing tissue was dissected from all three vestimentiferan tube worm hosts (from the trophosome) and from *Calyptogena magnifica* clams (from the gills) at the Tica vent site for future use as positive and negative controls, respectively. Environmental samples were collected from two distinct habitats: surface-attached biofilms and seawater.

Symbionts in surface-attached biofilms were collected on bacterial settlement devices deployed in four hydrothermal vent environments at increasing distances from tube worm clusters: (i) among tube worms, (ii) adjacent to tube worms, (iii) away from tube worms (~10 m), and (iv) off-axis (~100 m outside the axial summit of the caldera) (see Fig. S1 in the supplemental material). Settlement devices were constructed of polyvinyl chloride holders containing three to five basalt pieces (8 by 1 by 1 cm) and 4 to 12 glass microscope slides that were washed, autoclaved, and kept sterile until deployment. Devices were collected within 1 month or after 1 year. Upon collection, the basalt pieces were examined under a dissecting microscope to detect any settled tube worm larvae or juveniles and then immediately stored at –80°C. Pieces with observable tube worms were excluded to eliminate the risk of detecting symbionts living within host tissue. Microscope slides were fixed for fluorescence in situ hybridization (FISH) analysis in 4% paraformaldehyde and stored in 70% ethanol at 4°C.

Seawater samples were collected 1 m away from an *R. pachyptila* tube worm cluster using a McLane large-volume water transfer system water pump attached to the deep submergence vehicle *Alvin*. Samples (200 liters each) were filtered in situ through a 1- $\mu$ m Petex prefilter (Sefar) and then through a 0.45- $\mu$ m mixed-cellulose ester filter (Millipore). Control seawater samples (80 liters each) were collected from the ocean surface above the EPR and from the Atlantic Ocean in Nahant, MA. All filters were stored at –80°C until DNA extraction.

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TABLE 1. Detection of free-living symbiont phylotype of vent vestimentiferan tube worms via PCR and sequence analyses of biofilms<sup>a</sup>

Time of collection and type of deployment	Total no. of blocks <sup>d</sup>	No. of blocks with positive result by:			Sequence verification <sup>f</sup>
		PCR amplification <sup>b</sup>			
		Bacteria positive control	Tube worm symbiont	Tube worm host	
<b>~1 mo</b>					
Among <i>R. pachyptila</i>	3	3	3	ND <sup>e</sup>	3
Near <i>R. pachyptila</i>	4	4	4	ND	3
Away from <i>R. pachyptila</i>	5	5	5	ND	3
Off-axis	3	2	2	ND	1
<b>1 yr</b>					
Among <i>R. pachyptila</i>	1	1	1	ND	1
Near <i>R. pachyptila</i>	2	2	2	ND	NA <sup>f</sup>
Away from <i>R. pachyptila</i>	3	3	3	ND	3
Off-axis	3	3	3	ND	3

<sup>a</sup> Basalt blocks, deployed at various distances from *R. pachyptila* tube worm clusters at the Tica vent site along the EPR, were collected and analyzed after ~1 month and after 1 year.

<sup>b</sup> PCR amplification was performed using primers for universal *Bacteria* (27f and 1492r), vestimentiferan host-specific primers (for RP43, RifTOExoF and RifTOExoR), and symbiont-specific primers (RifTO44 and RifTO445).

<sup>c</sup> A subset of basalt biofilm samples was analyzed to verify the presence of the free-living tube worm symbiont phylotype.

<sup>d</sup> Number of basalt blocks analyzed for each deployment location and time.

<sup>e</sup> Tube worm hosts were not detected (ND) on any analyzed samples; samples with tube worm hosts were discarded.

<sup>f</sup> Only one sample was not analyzed (NA) because sequencing reactions did not work (including positive controls), perhaps due to inhibitory substrates in the DNA.

**16S rRNA gene sequence analyses.** PCR amplification and DNA sequence analyses were used to test for the presence of the vestimentiferan symbiont in biofilm and seawater samples. DNA was extracted by standard methods (27). The vestimentiferan symbiont 16S rRNA gene (a 401-bp fragment) was PCR amplified using primers specific for the shared 16S phylotype: RifTO44 (5'-GGCCTAGATTGACGCTGCGGTA-3') (this study) and RifTO445 (23). To detect contamination by host tissue, primers specific for the genes encoding the vestimentiferan host exoskeleton protein RP43 (GenBank accession no. AF233595), RifTOExoF (5'-CTAAAGGCAGTGTCAGAGCGGGAC-3') and RifTOExoR (5'-TTCCTCGAAGTTGCCGTATGCCG-3'), were used. PCR products were cloned into a pCR2 cloning vector (Invitrogen) and sequenced by standard methods using BigDye Terminator cycle sequencing reaction kits (PE Biosystems) with M13 forward and reverse primers. Symbiont- and host-specific primers amplified their target genes in the control symbiont-containing tissue samples from *R. pachyptila*, *T. jerichonana*, and *O. alvinae* worms, while vestimentiferan symbionts were not amplified from *C. magnifica* gill tissue, the negative control.

The free-living vestimentiferan symbiont 16S rRNA phylotype was detected in both biofilm and seawater samples collected at the Tica vent site. The symbiont phylotype (GenBank accession no. U77478) (9) was amplified from all basalt pieces retrieved after 1 month and after 1 year, including those from the off-axis site, away from active venting, and those from vent seawater samples on both 0.45- and 1- $\mu$ m-pore-size water filters (Table 1). Host tissue was detected only on a single pre-filter (1  $\mu$ m) water sample. PCR amplifications from surface

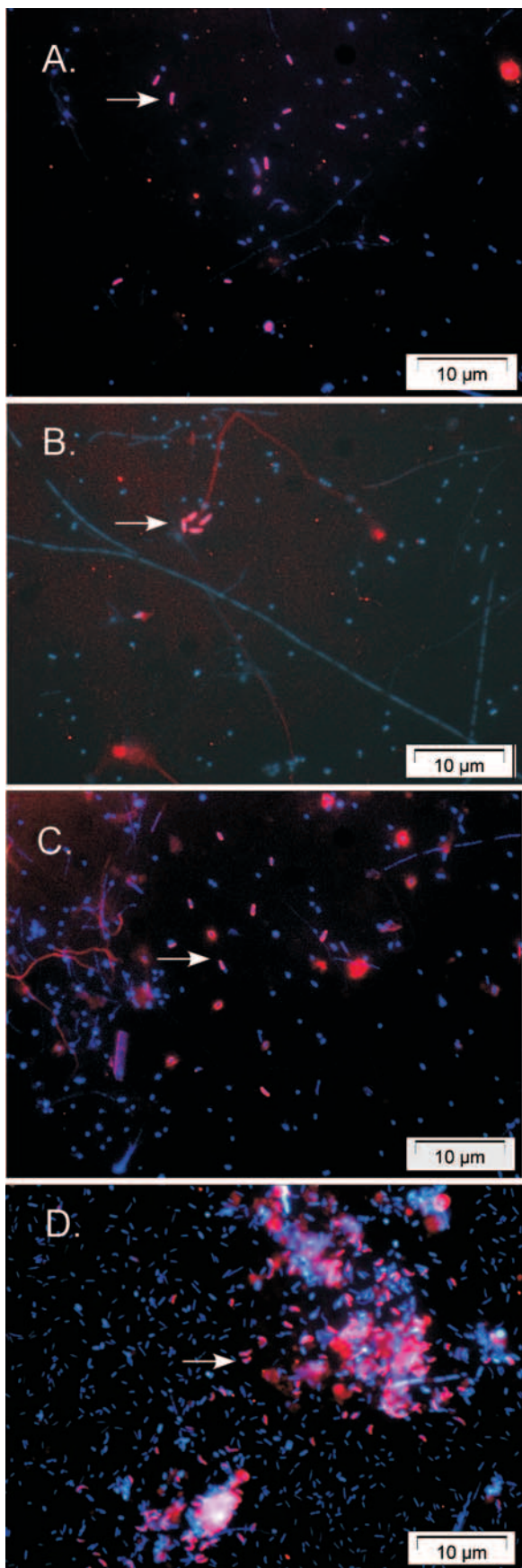
seawater control samples yielded positive PCR results with universal *Bacteria* primers (27f and 1492r) (13) but yielded negative results when either the vestimentiferan symbiont- or host-specific primers were used, suggesting that symbiont phylotypes were present only in deep-seawater samples. Little is yet known about the metabolic state or energy source for symbionts outside of their tube worm hosts, but it is possible that free-living symbionts may be cystic or quiescent while awaiting the inoculation of larval or juvenile tube worms.

**FISH.** FISH was used to provide direct visual evidence of the tube worm symbiont on glass slides recovered from bacterial settlement devices. For each slide, a universal *Bacteria* probe, Eub338 (1), either 5' end labeled with fluorescein or stained with the DNA-binding fluorescent dye 4',6'-diamidino-2-phenylindole (DAPI), was used as a positive control along with the symbiont-specific probe RifTO147, RifTO445, or RifTO830 that was 5' end labeled with Cy3 (23). The images from the control and symbiont-specific probes were then overlaid. The probe specificity was tested on *R. pachyptila* trophosome tissue, and the formamide concentration was increased until no probe remained hybridized (probe dependent, 20% [for Fig. S2 in the supplemental material] or 35% [for Fig. 1]). On each slide, either a nonsense probe, NON338 (23), or a 1-base-mismatch probe was used as a negative control. Hybridized slides were viewed and digitally photographed using a Leica model DMRB fluorescence microscope.

The tube worm symbiont phylotype was detected using FISH on all slides tested (Fig. 1; see Fig. S2 in the supplemental material) with the exception of the off-axis samples that were collected from devices deployed for less than 1 month. Although not directly quantified, the overall bacterial abundance appeared to be greatest on slides deployed for 1 year among, adjacent to, or away from the tube worms. The direct detection of the tube worm symbiont in biofilms supports the hypothesis that these bacteria exist in the free-living vent environment. Indeed, in a coastal marine endosymbiosis, the 16S phylotype of bacterial symbionts of *Codakia orbicularis* clams is readily found in the sea grass sediment surrounding their hosts (11).

**Endosymbiont ITS diversity.** If vestimentiferan tube worms acquire their symbionts from a diverse environmental source population, it can be hypothesized that the symbiont population within a host may consist of multiple closely related phylotypes (8, 31). The symbiont internal transcribed spacer (ITS), which is under relaxed selection relative to the 16S and has been used extensively to assess strain-level variation in bacteria (29), was cloned and sequenced to test for the presence of multiple symbiont phylotypes within individual tube worms. The ITS, located between the 16S and 23S rRNA genes in the bacterial rRNA operon, occurs as a single copy in the vestimentiferan symbiont genome (16, 26). By using symbiont-specific primers embedded in the 16S and 23S rRNA genes (Sym-ITS-1322F and Sym-ITS-23SR) (31), the ITS was PCR amplified (30 cycles with *Taq* polymerase) from DNA extracted from the trophosomes of three adult *R. pachyptila* worms. PCR products were cloned and sequenced (96 clones per specimen; 288 in total).

Analysis of the ITS sequences from the three *R. pachyptila* symbiont clone libraries revealed high levels of genetic homogeneity in intracellular symbiont populations. Sequence analysis revealed one dominant symbiont phylotype within each of



the three host specimens (accounting for 65, 77, and 41% of the sequences, respectively), and the third specimen hosted a second phylotype (27%), which consistently differed by the same two nucleotides. The majority of the remaining ITS sequences were singletons that cannot be distinguished from errors resulting from PCR or *Taq* analyses. The detection of diverse ITS sequences in *R. pachyptila* worms further supports the acquisition of bacteria from the environment, but the diversity of free-living symbionts has not yet been investigated.

**Evidence for environmental symbiont acquisition.** Detection of the free-living tube worm symbiont phylotype supports the hypothesis that newly settled tube worms obtain their bacteria from the vent environment. Along a spatial gradient, free-living symbionts were present among, adjacent to, and away from (within 10 m) tube worms and were also detected 100 m outside the areas of hydrothermal activity. The presence of free-living symbiotic bacteria at multiple spatial scales within a vent site suggests a potentially large environmental pool of symbionts. During host larval development and the colonization of new vents (17, 20, 21), an abundant free-living bacterial population would facilitate the initiation of the symbiosis. The environmental transmission of symbionts seems to be a risky strategy for obligate tube worm symbioses, as the survival of the mouthless and gutless adult host requires that developing larvae or juveniles successfully acquire their symbionts from a potentially unstable free-living source population. However, this developmental mode might be beneficial if it provides the host with opportunities to acquire specific, locally adapted symbiont genotypes.

**Influence of symbiotic bacteria on free-living microbial diversity.** Symbioses, notably those that are facultative, clearly have an impact on and may be a driving force of local microbial diversity in varied ecosystems (2, 10). Indeed, the bacterial symbionts of the shrimp *Rimicaris exoculata* make up a major component of the surrounding microbial community at hydrothermal vents in the Atlantic Ocean (25). Likewise, a free-living counterpart to the bioluminescent symbiotic bacterium *Vibrio fischeri* of squid has been identified in coastal environments, revealing a connection between the symbiotic relationship and microbial abundance and distribution (15). The same situation appears to be true in legume-rhizobium symbioses; the host species is thought to be a major factor in determining the characteristics of the soil microbial community (18). Endosymbiont and free-living populations may affect each other via positive feedback cycles, whereby the host inoculates the free-living population, and the free-living population inoculates the host (24). This study serves as the basis for future investigations of the biodiversity and biogeography of free-living marine symbionts at multiple spatial scales.

FIG. 1. FISH detection of free-living vestimentiferan bacterial symbionts. Representative slides deployed at the Tica hydrothermal vent site on the EPR for ~1 month among *Riftia pachyptila* tube worms (A), near tube worms (B and C), and 10 m away from tube worms (D) are shown. The overlay of two images with symbiont-specific probes (red [Cy3]) and DAPI (blue) shows the free-living symbionts (arrows) labeled with the symbiont-specific probes RifTO830 (A and C), RifTO147 (B), and RifTO445 (D).

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