Molecular Microbial Diversity Survey of Sponge Reproductive Stages and Mechanistic Insights into Vertical Transmission of Microbial Symbionts

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Many marine sponges, hereafter termed high-microbial-abundance (HMA) sponges, harbor large and complex microbial consortia, including bacteria and archaea, within their mesohyl matrices. To investigate vertical microbial transmission as a strategy to maintain these complex associations, an extensive phylogenetic analysis was carried out with the 16S rRNA gene sequences of reproductive (n = 136) and adult (n = 88) material from five different Caribbean species, as well as all published 16S rRNA gene sequences from sponge offspring (n = 116). The overall microbial diversity, including members of at least 13 bacterial phyla and one archaeal phylum, in sponge reproductive stages is high. In total, 28 vertical-transmission clusters, defined as clusters of phylotypes that are found both in adult sponges and their offspring, were identified. They are distributed among at least 10 bacterial phyla and one archaeal phylum, demonstrating that the complex adult microbial community is collectively transmitted through reproductive stages. Indications of host-species specificity and cospeciation were not observed. Mechanistic insights were provided using a combined electron microscopy and fluorescence in situ hybridization analysis, and an indirect mechanism of vertical transmission via nurse cells is proposed for the oviparous sponge *Ectyoplasia ferox*. Based on these phylogenetic and mechanistic results, we suggest the following symbiont transmission model: entire microbial consortia are vertically transmitted in sponges. While vertical transmission is clearly present, additional environmental transfer between adult individuals of the same and even different species might obscure possible signals of cospeciation. We propose that associations of HMA sponges with highly sponge-specific microbial communities are maintained by this combination of vertical and horizontal symbiont transmission.
which rely on their chemosynthetic symbionts (34), and the sepiolid squid-vibrio symbiosis (33).

The first evidence for vertical transmission in sponges came from electron microscopy studies (references 6 and 9 and references cited therein; 25, 44, 52). To date, microbes have been detected in oocytes, embryos, and larvae of more than 20 oviparous sponges (which release gametes) and viviparous sponges (which release fully developed larvae) (9, 42). Recently, a culture-based study successfully isolated the same alphaproteobacterial strain from adult and embryo samples of the Caribbean sponge Mycale laxissima, and vertical transmission of this bacterium was suggested by the authors (8). Additional evidence for vertical transmission comes from molecular studies that allow the phylogenetic identification of microbes within sponge reproductive stages and comparison with the microbial community of the adult. Using Cyanobacteria-specific primers, Oren et al. (35) found a Synechococcus-related sequence type in Diacarnus erythraenus adult sponges and their embryos. However, so far only three molecular studies investigated the total microbial diversity in sponge offspring. Complex bacterial consortia were described for each of the species M. laxissima, Ircinia felix, and Corticium sp. (8, 45, 46, respectively). Sharp et al. (46) also described archaea within Corticium sp. embryos and verified the presence of microbes by fluorescence in situ hybridization (FISH). Additionally, a complex microbial community was described for the first time in I. felix juvenile sponges which were obtained after larval settlement experiments (45).

To provide insights into the phylogenetic diversity of bacteria and archaea in sponge reproductive stages, a comprehensive 16S rRNA gene survey was performed with >250 sequences obtained from eight sponges representing six orders, two geographic regions, and both oviparous and viviparous modes of reproduction. Furthermore, the mechanistic aspects of vertical transmission were investigated in the oviparous Caribbean sponge Ectyoplasia ferox using a combination of electron and fluorescence microscopy. E. ferox was chosen because its early reproductive stages are obtainable, allowing mechanistic investigations from an early developmental stage onward. Based on the phylogenetic and mechanistic results, we propose a symbiotic transmission model for HMA sponge symbioses, including a combination of vertical and horizontal symbiotic transmission to maintain the association with sponge-specific, though not species-specific, microbial consortia.

### MATERIALS AND METHODS

**Sponge collection.** The sponges Agelas conifera, Agelas wiedenmayeri, Ectyoplasia ferox, Smenospongia aurea, and Xestospongia muta were collected by scuba diving off the coast of Key Largo, Florida, in June 2002, June and August 2004, and May 2005 (Table 1). Additional material from E. ferox was collected by scuba diving offshore of French Wells, Bahamas, in July 2003. The samples were taken at depths of 5 to 30 m, transferred in ziplock bags containing seawater to the surface, and kept on ice in a cooler until further processing (within 4 to 5 h). Offspring samples (oocytes, embryos, and larvae) were collected in larval traps using the methodology of Lindquist et al. (27). Additionally, tissue samples of the corresponding adult individuals were taken. After the adult samples were cut into small pieces, the adult and offspring materials were washed in sterile filtered seawater, fixed either in 96% ethanol or in 2.5% glutaraldehyde-double-distilled water (ddH₂O), and stored at either −20 or 4°C. The field settlement experiments were performed with a portion of the captured S. aurea larvae as described by Schmitt et al. (45). Briefly, larvae were enclosed in plastic containers (~75 ml in volume), which were placed at a 9-m-deep reef site. These containers consisted of a plastic frame supporting a covering of 200-μm nylon mesh to allow water exchange. Juveniles were recovered 1 to 3 days postsettlement and preserved as described above.

**Extraction of DNA and 16S rRNA gene library construction.** Adult samples that had been fixed in 96% ethanol were air dried, ground with a mortar and pestle in liquid nitrogen, and DNA extracted using the Fast DNA Spin kit for soil (QiBiogene, Heidelberg, Germany) in accordance with the manufacturer’s instructions. Reproductive material containing oocytes and embryos was also air dried and transferred into 150 μl ddH₂O, and DNA was extracted by heating the samples in a water bath for 10 min at 100°C. This same methodology was used to

### TABLE 1. Compilation of all available 16S rRNA gene sequences from sponge reproductive stages until December 2007

<table>
<thead>
<tr>
<th>Host sponge (order)</th>
<th>Collection site(s) (latitude/longitude)</th>
<th>Mode of reproduction</th>
<th>Sequence source(s)</th>
<th>No. of offspring sequences</th>
<th>Reference</th>
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</thead>
<tbody>
<tr>
<td><strong>HMA sponges</strong></td>
<td></td>
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<td></td>
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<tr>
<td>Agelas conifera</td>
<td>Key Largo, FL (25°02’N/80°24’W)</td>
<td>Oviparous</td>
<td>16S rRNA library</td>
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<td>This study</td>
</tr>
<tr>
<td>Agelas wiedenmayeri</td>
<td>Key Largo, FL (25°02’N/80°24’W)</td>
<td>Oviparous</td>
<td>DGGE</td>
<td>7</td>
<td>This study</td>
</tr>
<tr>
<td>Corticium sp. (Agelasida)</td>
<td>Palau islands (07°23’N/134°38’E)</td>
<td>Viviparous</td>
<td>16S rRNA library</td>
<td>18</td>
<td>46</td>
</tr>
<tr>
<td>Ectyoplasia ferox</td>
<td>Key Largo, FL (25°02’N/80°24’W)</td>
<td>Oviparous</td>
<td>DGGE</td>
<td>28</td>
<td>This study</td>
</tr>
<tr>
<td>Ircinia felix</td>
<td>Bahamas islands (24°32’N/75°55’W)</td>
<td>Viviparous</td>
<td>DGGE</td>
<td>71</td>
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<tr>
<td>Smenospongia aurea</td>
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<td>Viviparous</td>
<td>16S rRNA library</td>
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<td>This study</td>
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<td>16S rRNA library</td>
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<td>Mycale laxissima</td>
<td>Key Largo, FL (25°02’N/80°24’W)</td>
<td>Viviparous</td>
<td>Isolates, 16S rRNA library</td>
<td>27</td>
<td>8</td>
</tr>
</tbody>
</table>

* All sponges belong to the class Demospongiae (classification after Systema Porifera by Hooper and van Soest [19]).

*b* Reclassified after the paper by Schmitt et al. (43).

*c* Sequences were obtained using universal 16S rRNA primers either by the construction of 16S rRNA libraries or by the excision of DGGE bands.

*d* The 16S rRNA library was constructed using group-specific primers.
obtain DNA from collections of larvae and juveniles (three, five, or seven individuals pooled immediately after collection). The DNA solutions were then used directly as templates (0.01 to 1.0 µg) in the following PCR.

The universal primers 27f and 1492r (23) were used for the PCR amplification of bacterial 16S rRNA genes. The primer pairs POR389f/POR1130r (12), 11f/ Ntspa865r (47), and Arch21f/Arch958r (7) were used for the specific PCR amplification of the candidate phylum “Poribacteria,” Nitrospira, and Archaes 16S rRNA genes, respectively. Negative controls (PCRs without DNA templates) were included for each 16S rRNA gene amplification. Cycling conditions on a Mastercycler gradient (Eppendorf, Hamburg, Germany) were as follows: an initial denaturing step at 96°C for 5 min; 30 cycles of denaturing at 96°C for 1 min; primer annealing at 54, 64, and 56°C for 1 min; and elongation at 72°C for 90, 50, and 50 s for universal, Poribacteria, Nitrospira, and Archaes PCRs, respectively. The PCR program ended with a final extension step at 72°C for 5 min.

**DGGE.** DNA was extracted as described above. The universal 16S rRNA gene primers 341f with GC-clamp and 907r (32) were used for the PCR amplification of bacterial 16S rRNA genes. The cycling conditions were as follows: initial denaturing step at 95°C for 5 min, 30 cycles of denaturing at 95°C for 1 min, primer annealing at 54°C for 1 min and elongation at 72°C for 45 s, followed by a final extension step at 72°C for 10 min. Denaturing gradient gel electrophoresis (DGGE) was performed with a Bio-Rad DCode universal mutation detection system (Bio-Rad, München, Germany) on a 10% (wt/vol) polyacrylamide gel in 1× Tris-acetate-EDTA using a 0 to 90% denaturing gradient; 100% denaturant corresponded to 7 M urea and 40% (vol/vol) formamide. Electrophoresis was performed for 6 h at 150 V and 60°C. The gels were stained for 30 min in 50 g/l 1-Butanol (Molecular Probes) and scanned on a Typhoon 8600 scanner (Amersham Biosciences). Selected bands were excised with an ethanol-sterilized scalpel and incubated in 25 µl ddH₂O overnight at 4°C. A total of 4 µl of eluted DNA was subsequently used for reamplification with primers 341f and 907r using the PCR conditions described above.

**Cloning, sequencing, and phylogenetic analysis.** The PCR products from the excised DGGE bands and for 16S rRNA gene libraries were ligated into the pGEM-T Easy vector (Promega) and transformed by electroporation into competent *Escherichia coli* XL-1-Blue cells. Plasmid DNA was isolated by standard miniprep procedures (40), and the correct insert size was verified using agarose gel electrophoresis following restriction digestion.

The clones from the 16S rRNA gene libraries were characterized by single digestions with the restriction endonucleases HaeIII and Sau3AI (restriction fragment length polymorphism analysis). The clones with identical restriction patterns were grouped together, and one random clone from each group was chosen for sequence analysis. Sequencing was performed on an ABI 377XL automated sequencer (Applied Biosystems), and the sequences were edited with the ContigExpress tool in the Vector NTI suite 6.0 (InforMax, Inc.). The sequences were checked for chimeras with the program Pintail (2) and for other amplification and sequencing artifacts. Following the removal of chimeras from the data set, the percentage similarities (p-distances) between the sequences from the same sponge developmental stage (adult, oocyte, embryo, larva, and juvenile) within each species were determined with the software Align, and those with identities above 99% were grouped together into operational taxonomic units. Only one randomly chosen sequence per operational taxonomic unit was used for further analysis. The sequences obtained in this study, all available sequences from sponge reproductive material in GenBank (http://www.ncbi.nlm.nih.gov), and the appropriate reference sequences (all nearest BLAST matches including sequences from adult sponges and, moreover, representatives of the respective phylum) were aligned automatically with ClustalX (51), and the alignment was subsequently corrected manually in Align. Phylogenic trees were constructed with the ARB software package (28). Initially, neighbor-joining (Jukes-Cantor correction) and maximum parsimony trees were calculated with nearly full-length sequences (>1,250 bp) and 100 pseudoreplicates. Subsequently, partial sequences were added to the trees without changing the topology by the use of the parsimony-interactive method in ARB.

**TEM.** Adult samples that had been fixed in 2.5% glutaraldehyde-0.1M H₂O were cut into small cubes of about 1 mm³. All samples (adults, material containing oocytes and embryos, larvae, and juveniles) were then washed five times in cadaverine buffer (50 mM, pH 7.2), fixed in 2% osmium tetroxide for 90 min, washed again five times in ddH₂O, and incubated overnight in 0.5% uranyl acetate. After being dehydrated in an ethanol series (30, 50, 70, 90, 96, and 3 times at 100% for 30 min each), the samples were incubated three times for 30 min in 1× propylene oxide, maintained overnight in 1:1 (vol/vol) propylene oxide–Epon 812 (Serva), incubated twice for 2 h in Epon 812, and finally embedded in Epon 812 for 48 h at 60°C. The samples were then sectioned with an ultramicrotome (OM U3; C. Reichert, Austria) and examined by transmission electron microscopy (TEM) (Zeiss EM 10; Zeiss, Germany).

**FISH.** Small pieces (ca. 5 mm³) of the ethanol-preserved *E. ferox* offspring material (oocytes/embryos embedded in a mucous sheath) were incubated in 50 and 30% ethanol for 3 min each, transferred into 1× phosphate-buffered saline for 3 min, and frozen in Tissue-Tek cryo-embedding medium (Miles Laboratories, Inc., Naperville, IL) at −80°C for at least 2 days. Frozen *E. ferox* offspring material was directly placed in cryo-embedding medium and kept at −80°C. For hybridization, pieces were sectioned with a cryomicrotome (Mikrom, Walldorf, Germany), placed on microscope slides, and air dried. After fixation in 4% paraformaldehyde (final concentration) overnight at 4°C in a humid chamber, the samples were washed twice in 1× phosphate-buffered saline for 5 min each, heat fixed at 42°C, dehydrated in an ethanol series (50, 75, and 100% for 5 min each), and air dried. FISH was performed in an isotonically equilibrated humid chamber at 46°C in hybridization buffer (0.9 M NaCl, 20 mM Tris-HCl [pH 8.0], 0.01% sodium dodecyl sulfate) and probe-dependent amounts of formamide (25 to 50%) for 4 h. The Cy3-labeled probe mix EUB III (5) was used at concentrations of 2 to 7.5 ng/µl, whereas the fluorescein isothiocyanate-labeled EUK516 probes (1) were used at concentrations of 15 to 50 ng/µl. Additionally, 5 µl of a DAPI stock solution (100 µg/ml) was added to each reaction 30 min prior to stopping the hybridization. The hybridization buffer was gently rinsed off, and the slides were incubated at 48°C in washing buffer (0.7 M NaCl, 20 mM Tris-HCl [pH 8.0], 50 mM EDTA, 0.01% sodium dodecyl sulfate) for 30 min. The washing buffer was rinsed off with ice-cold ddH₂O, and the slides were air dried in the dark and mounted in Citifluor (Citifluor Ltd., London, United Kingdom). Visualization was achieved with a confocal laser scanning microscope (LSM 510; Zeiss MicroImaging).

**Nucleotide sequence accession numbers.** The 16S rRNA gene sequences were deposited in EMBL/GenBank/DDJB under accession numbers EF159734 to EF159941 and EU273899 to EU273914.

**RESULTS**

**Phylogenetic sequence analysis.** In total, 218 bacterial 16S rRNA gene sequences were obtained in this study: 6 from *A. conifera* (3 adult, 3 offspring), 8 from *A. wiedenmayeri* (1 adult, 7 offspring), 33 from *E. ferox* (7 adult, 26 offspring), 2 from *I. felix* (1 adult, 1 offspring), 61 from *S. aurea* (42 adult, 19 offspring), and 108 from *X. muta* (32 adult, 76 offspring). Additionally, six archaeal 16S rRNA gene sequences were recovered from the sponges *A. conifera* (2 adult, 2 embryo) and *E. ferox* (2 embryo). These sequences, together with 18 sequences from *Cor-ticium* sp. offspring (46), 71 from *I. felix* offspring (45), 27 from *M. laxisima* offspring (8), and 2 from *Luffariella variabilis* offspring (48), were used for phylogenetic tree construction.

The overall microbial diversity in sponge offspring is high (Fig. 1 to 7). 16S rRNA gene phytypes are affiliated with the bacterial phyla *Aquificae*, *Acidobacteria* and *Bacteroidetes*. Filled and open circles represent neighbor-joining bootstrap values above 95% and 80%, respectively. Neighbor-joining and maximum parsimony bootstrap values are given for each VT cluster. Sequences obtained in this study are given in bold. Short sequences that were added to the tree using the parsimony interactive tool in ARB are indicated by dashed lines. Arrow, to outgroup (*Pyrobaculum calidiformis* AB078332 and *Sulfolobus metallicus* D85519). Gray boxes depict VT clusters that contain sequences obtained from adults and offspring of the same species. Scale bar indicates 10% sequence divergence.
FIG. 2. Evolutionary distance tree calculated with 16S rRNA gene sequences affiliated with the bacterial phyla **Nitrospira**, **Gemmata monadetes**, and **Actinobacteria**. Details are the same as those provided for Fig. 1.
FIG. 3. Evolutionary distance tree calculated with 16S rRNA gene sequences affiliated with the bacterial phyla **Cyanobacteria** and **Chloroflexi**. Details are the same as those provided for Fig. 1.
FIG. 4. Evolutionary distance tree calculated with 16S rRNA gene sequences affiliated with the bacterial class Alphaproteobacteria. Details are the same as those provided for Fig. 1. The cluster containing the isolate from *M. laxissima* adults and embryos is indicated by an unshaded box.
FIG. 5. Evolutionary distance tree calculated with 16S rRNA gene sequences affiliated with the bacterial classes Beta- and Gammaproteobacteria. Details are the same as those provided for Fig. 1.
FIG. 6. Evolutionary distance tree calculated with 16S rRNA gene sequences affiliated with the bacterial phyla Planctomycetes, Verrucomicrobia, and Firmicutes; with the class Deltaproteobacteria; and with the candidate phyla TM7 and Poribacteria. One group of sequences could not be unambiguously affiliated within the Bacteria and is marked as “uncertain affiliation.” Details are the same as those provided for Fig. 1.
(n = 4), Planctomycetes (n = 6), Proteobacteria (Alpha- [n = 55], Beta- [n = 1], Gamma- [n = 40], and Deltaproteobacteria [n = 8]), and Verrucomicrobia (n = 2), as well as with the bacterial candidate phyla Poribacteria (n = 4) and TM7 (n = 1). An additional cluster of bacterial sequences (n = 8) could not be unambiguously affiliated within the Bacteria (Fig. 6). Archaeal sequences obtained from sponge offspring all fell within marine group I of the Crenarchaeota (n = 6) (Fig. 7).

VT clusters. Sequences from an adult sponge and its offspring that cluster together in the phylogenetic analysis are referred to as vertically transmitted sequence pairs. Together with other sponge-specific sequences, they build clusters which are referred to as vertical-transmission (VT) clusters. Twenty-five VT clusters were found, distributed among 10 different bacterial phyla (including two candidate phyla) (Fig. 1 to 6). One additional VT cluster could not be assigned to any known bacterial phylum and was therefore termed VT-Uncertain (Fig. 6). Within the Archaea, two VT clusters were obtained (Fig. 7). Six clusters are double VT clusters (VT-Acido-IV, VT-Actino-II, VT-Alpha-II, VT-Gamma-III, VT-Gamma-IV, VT-Gamma-V).
and VT-Nitrospira), and one is a triple VT cluster (VT-Chloroflexi-II), as they contain vertically transmitted sequence pairs of two or three sponge species, respectively. Twenty-nine percent of *A. wiedenmayeri*, 53% of *Corticium* sp., 23% of *E. ferox*, 75% of *I. felix*, 0% of *M. laxissima*, 68% of *S. aurea*, and 23% of *X. muta* offspring sequences belong to VT clusters.

**TEM.** *E. ferox* released orange-colored oocytes which were embedded in a mucous sheath. Electron micrographs of the oocytes revealed the presence of many inclusions equally scattered within the cytoplasm (Fig. 8A). These inclusions vary in electron density and in size and represent, e.g., lipid droplets (1 to 1.5 μm in diameter) and yolk granules (2 to 3 μm in diameter). Microorganisms could not be detected in the oocyte cytoplasm. The oocyte is surrounded by a layer of amoeboid-shaped nurse cells which are 5 to 8 μm in size (Fig. 8A and B). Each nurse cell contains one 1- to 2-μm large nucleus, several smaller vacuoles, and usually one large vesicle with 15 to 20 (sometimes more) microorganisms of different sizes of up to 2.5 μm and various shapes including rod-like and coccus-like cells (Fig. 8B). At this stage of development, there is a clear space between nurse cells and the oocyte. However, the oocyte already shows signs of invagination (Fig. 8B). After 2 days, the oocytes developed into early-stage embryos, which were still embedded in the gelatinous sheath. The embryo consists of many small, densely packed amoeboid-shaped cells that contain one 2- to 3-μm large nucleus each and again diverse inclusions, including yolk granules (Fig. 8C and D). A thin lamella of collagen fibrils surrounds the embryo (Fig. 8C), and nurse cells are no longer present at this stage of development. Clusters of morphologically diverse microorganisms can be found extracellularly within the embryo, mostly located in the periphery.

**FISH.** A cohybridization with the bacterial probe mix EUB I-III and the eukaryotic probe EUK516, in addition to the DNA dye DAPI, was performed on cross-sections of *E. ferox* oocytes (Fig. 9). The 150- by 100-μm-large oocyte showed no background autofluorescence (Fig. 9A). The interior appeared heterogeneous (Fig. 9B). Eukaryotic ribosomes are present in large numbers as indicated by many green signals. The 4- to 5-μm-large nucleolus is especially prominent in the center of the oocyte within the nucleus, which has a size of 20 μm. Additionally, dark blue, circular signals (about 3 μm in diameter), as well as red signals of various sizes, are present. The oocyte is surrounded by nurse cells which can be identified by their nuclei due to blue DAPI signals and contain clusters of microorganisms represented by red signals (Fig. 9C).
DISCUSSION

Molecular microbial diversity survey of sponge reproductive stages. The first evidence for the vertical transmission of microorganisms in sponges came from electron microscopy studies that documented the presence of microbes in reproductive stages (references 6 and 9 and references cited therein; 25, 44, 52). The application of molecular tools allowed the phylogenetic identification of microorganisms in sponges and their reproductive stages. Prior to this study, the total microbial diversity in sponge offspring was analyzed for only three species in separate studies (8, 45, 46). Here we provide new 16S rRNA gene sequences from five additional sponges and report the first comprehensive phylogenetic analysis including more than 250 16S rRNA gene sequences from sponge reproductive stages.

The observed total phylogenetic diversity of microbes in sponge reproductive stages is high and comprises phylotypes within one archaeal phylum and 13 bacterial phyla (two of them are candidate phyla) including the four classes Alpha-, Beta-, Gamma-, and Deltaproteobacteria (Fig. 1 to 7). These high numbers almost match the known microbial diversity of adult sponges, which comprises two archaeal and 16 bacterial phyla and is predicted to be close to the total microbial diversity in sponges at the phylum level as assessed by the current approaches used (49). Most sponge offspring 16S rRNA gene sequences were derived from X. muta and I. felix, representing 11 and 8 phyla, respectively. However, just 7 sequences from A. wiedenmayeri embryos are affiliated with 4 phyla and 19 sequences from Corticium sp. already fall into 8 phyla. These numbers suggest that the presented data largely reflect the overall microbial diversity in sponge offspring, at least at the phylum level, and they show that equally complex microbial consortia are present in adult sponges and reproductive stages.

Microbes which are vertically transmitted were determined by VT clusters which are defined as phylotypes that are present in adult and offspring samples of the same species. More than two-thirds of the offspring sequences from I. felix and S. aurea and more than half from Corticium sp. fall into VT clusters implying that a large part of the adult microbial community is vertically transmitted in these three species. In A. wiedenmayeri, E. ferox, and X. muta, about one-third of the offspring-derived sequences are within VT clusters. Phyotypes that do not fall within VT clusters may represent environmental contamination, e.g., seawater microbes. However, it is noteworthy that many of the offspring sequences outside of the VT clusters still cluster with sequences obtained from other sponge species. For example, within the Gemmatimonadetes phylum (Fig. 2), three embryo sequences from Corticium sp. (clones CC08, -18, and -19) and one from E. ferox (clone EFE5) cluster with clones from the adult sponges I. felix (clone B3-1), Aplysina aerophoba (clone TK19), and Agelas dilatata (clone AD004). This cluster was not defined as a VT cluster because sequences from adult Corticium sp. and E. ferox are missing. Additionally, all VT clusters except clusters VT-TM7 (Fig. 6) and VT-Archaea-II (Fig. 7) contain additional sequences derived from other sponge species. The general overlap between offspring and adult microbial communities is striking. It is well recognized that HMA sponges share a uniform, sponge-specific microbial community (references 16 and 49 and references cited therein), and the presented data now demonstrate that this uniform microbial community is collectively transmitted through the reproductive stages.

In contrast to the results for the HMA sponges discussed above, sequences from M. laxissima embryos tend to cluster much less with other sponge-derived sequences, and none fall within a VT cluster. A recent analysis of the adult microbial community of M. laxissima also showed little overlap with the microbial diversity of HMA sponges but revealed a specific community in M. laxissima that is different from seawater bacterioplankton (30). Interestingly, M. laxissima was recently classified as an LMA sponge based on microscopic observations (44). The phylogenetic microbial signature of both the adult sponge and embryos that differ from the signature of HMA sponges supports this classification. M. laxissima might contain its own specific microbial community that is vertically transmitted. Notable is a bacterial isolate that was cultivated from the adults and embryos of M. laxissima. The respective sequence was also recovered from a 16S rRNA gene library of M. laxissima embryos and clusters with the marine alphaproteobacterium Pseudovibrio denitrificans (Fig. 4). The same phylotype was previously detected by cultivation-dependent- and -independent approaches in many other sponges but also in corals and marine environments (41, 49). It cannot be discounted that the Pseudovibrio-like bacterium was actually taken up by sponges from the seawater. However, because it is widespread among sponges and appears to be vertically trans-
mitted in *M. laxissima*, it probably represents a true sponge symbiont.

**Mechanism of vertical transmission in *E. ferox***. The phylogenetic results demonstrated vertical transmission in eight species representing six different orders and two modes of reproduction (ovipary and vivipary) (Table 1). The passage of symbionts through reproductive stages appears to be common and widespread among sponges. However, differences in the mechanism of symbiont transmission, e.g., between oviparous and viviparous species, are almost unknown (but see references 9 and 29). In this study, we conducted a combined electron and fluorescence microscopic examination of microbes within oocytes and embryos of the Caribbean sponge *E. ferox*. This oviparous species was chosen because oocytes can be collected, allowing the examination of vertical transmission from the early reproductive stages onward.

The oocyte is large and represents a late stage of gamete development. As fertilization was not observed, this unicellular stage could possibly also be a zygote. Typically, the oocyte contains many inclusions, including yolk and lipids, which mostly serve as nutrient reserves during the gamete and embryo growth periods (Fig. 8). The oocyte is metabolically highly active, as demonstrated by the large number of eukaryotic ribosomes throughout the cytoplasm (Fig. 9B). The nucleus is located in the center of the oocyte and contains several nucleoli (data not shown) of which one is visible in Fig. 9B as a bright green signal due to the synthesis of new rRNA. The oocyte is encircled by a layer of nurse cells (Fig. 8A and B). Nurse cells play a major role in the nutrition, and consequently the growth, of oocytes. The transfer of substances from nurse cells to oocytes via cytoplasmic bridges and/or direct phagocytosis of the whole nurse cell by oocytes has been described for several viviparous species (e.g., see references 11 and 22) as well as oviparous species (e.g., see references 14 and 24). In this developmental stage of *E. ferox*, the nurse cells have not yet made contact with the oocyte, but the phagocytic activity of the gamete is already visible (Fig. 8B). The presence of microorganisms within nurse cells was shown in electron micrographs (Fig. 8A and B) and further confirmed by clusters of bacterial-specific red signals encircling the oocyte in fluorescence micrographs (Fig. 9B and C). Single red signals were also observed within the oocyte in FISH analyses but were absent in the negative control (Fig. 9A). These signals might correspond to microbes. However, as microbes within the oocyte were never detected in the electron micrographs, the identity of these red signals remains unclear.

The embryonic stage of development that is represented by Fig. 8C and D is characterized by many densely packed sponge cells that include lipids, yolk reserves, and other inclusions. The embryo is surrounded by a lamella of collagen fibrils. Such filamentous structures were previously described, e.g., the enveloping embryos of *Spongia* sp. and *Hippospongia lachne* (22) as well as *Lutrunculia magnifica* (21). Ilan (21) suggested that the fibrils are secreted by a layer of sponge cells. However, the origin as well as the function of this lamella in *E. ferox* remains unknown. Clusters of microorganisms are present extracellularly within the embryo (Fig. 8D).

*E. ferox*, like all other sponges, lacks reproductive organs, and gamete production occurs within the mesohyl of adult sponges. Based on the microscopic observations, the following
mechanism of microbial transmission in *E. ferox* is proposed: during the establishment of oocytes, nurse cells take up adult microorganisms from the mesohyl via phagocytosis. Presumably, this uptake occurs mainly at random. However, nurse cell phagocytosis of cyanobacteria in the cortex region and subsequent transport to the developing eggs deeper within the sponge were observed in *Chondrilla australiensis* (52), suggesting a possible control of symbiont uptake by these cells. Afterwards, the reproductive propagules, along with associated nurse cells that now form a cell layer around the oocytes, become embedded in a mucous sheath that is produced and then released by the sponge. The same mode of reproduction was observed to occur in other sponges, including *X. muta, A. conifera*, and *A. wiedenmayeri* (N. Lindquist, personal observation). Microorganisms are then either transferred from nurse cells to oocytes or, at a later stage, to the developing embryo. In the majority of oviparous species as well as in at least one viviparous sponge, the transfer of microbes appears to occur directly as the microbes are taken up by the oocyte itself from the adult mesohyl (reference 9 and references cited therein). In *E. ferox* and some other sponges, such as *Chondrosia reniformis* (24), the transfer takes place indirectly because the microbes are translocated through nurse cells to the progeny.

Transmission model for symbioses of HMA sponges and complex microbial consortia. Coprecipitation between sponges and microbes is often difficult to prove due to the complexity of the microbial consortia and/or the generally conservative characteristic of some phylogenetic markers such as the 16S rDNA gene. Based on largely congruent phylogenies, cospeciation was demonstrated for the filamentous cyanobacterium *Oscillatoria spongiae* and dictyoceratid sponges (39, 50) and for an alphaproteobacterial symbiont and halichondrid sponges (10). However, in both cases, at least one host-switching event occurred. Our data demonstrate the transmission of a highly sponge-specific microbial community via reproductive stages. However, the phylogenetic trees show no indication of host-species specificity or cospeciation.

Based on our phylogenetic results and the mechanistic details described for *E. ferox*, we therefore propose the following symbiont transmission model for associations of HMA sponges and their microbial consortia (Fig. 10): during sponge reproduction, gametes or later reproductive stages take up microbes from the adult mesohyl microbial community. Thereby, at least in some species, including *E. ferox*, nurse cells are involved in microbial incorporation and might actually represent the important factor fostering selection of this association. Sponge larvae are unable to feed and might therefore be regarded as a closed system with no exchange with the environment. Vertical transmission therefore causes the separation of symbionts in different sponges which might eventually result in genetic differentiation and would ultimately lead to cospeciation. After the settlement of sponge larvae and their subsequent morphosis into juveniles, the filter-feeding life phase begins, allowing the uptake of seawater microbes. In this phase, the horizontal transfer of symbionts between host individuals or environmental acquisition from seawater appears to be possible, at least to some degree, and evidence from sponges was recently discussed in detail by Taylor and colleagues (49). Symbionts might be released by sponges due to tissue damage by predators and environmental factors (currents and storms) or during spawning periods. The uptake of symbionts could possibly occur during filter-feeding periods of the host. It might also be conceivable that sponge larvae acquire microbial symbionts that colonize the outer surface of the larvae, after the release of larvae from the adult sponge. Such symbiont uptake and release mechanisms would result in symbiont dispersal among host individuals and even species, thus obscuring the pattern of cospeciation in phylogenetic trees (Fig. 10).

The importance of vertical transmission lies in the efficient passage of symbionts to the next generation. In HMA sponges, vertical transmission represents a bottleneck in the sense that only members of the specific adult microbial community, but apparently no seawater microorganisms, are transferred through reproductive stages. Horizontal transmission acts against symbiont separation and specialization. Instead of leading to cospeciation, this mechanism might maintain a uniformly shared microbial community in HMA sponges. We propose that a combination of vertical and horizontal symbiont transmission maintains the associations of HMA sponges with highly symbiont-specific, complex microbial communities.

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