

Corals Form Characteristic Associations with Symbiotic Nitrogen-Fixing Bacteria

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The complex symbiotic relationship between corals and their dinoflagellate partner *Symbiodinium* is believed to be sustained through close associations with mutualistic bacterial communities, though little is known about coral associations with bacterial groups able to fix nitrogen (diazotrophs). In this study, we investigated the diversity of diazotrophic bacterial communities associated with three common coral species (*Acropora millepora*, *Acropora muricata*, and *Pocillopora damicornis*) from three mid-shelf locations of the Great Barrier Reef (GBR) by profiling the conserved subunit of the *nifH* gene, which encodes the dinitrogenase iron protein. Comparisons of diazotrophic community diversity among coral tissue and mucus microenvironments and the surrounding seawater revealed that corals harbor diverse *nifH* phylotypes that differ between tissue and mucus microhabitats. Coral mucus *nifH* sequences displayed high heterogeneity, and many bacterial groups overlapped with those found in seawater. Moreover, coral mucus diazotrophs were specific neither to coral species nor to reef location, reflecting the ephemeral nature of coral mucus. In contrast, the dominant diazotrophic bacteria in tissue samples differed among coral species, with differences remaining consistent at all three reefs, indicating that coral-diazotroph associations are species specific. Notably, dominant diazotrophs for all coral species were closely related to the bacterial group rhizobia, which represented 71% of the total sequences retrieved from tissue samples. The species specificity of coral-diazotroph associations further supports the coral holobiont model that bacterial groups associated with corals are conserved. Our results suggest that, as in terrestrial plants, rhizobia have developed a mutualistic relationship with corals and may contribute fixed nitrogen to *Symbiodinium*.

Corals are described as holobionts, where the coral animal hosts an array of mutualistic microorganisms, including the endosymbiotic dinoflagellate *Symbiodinium*, bacteria, archaea, and fungi (28, 44, 47, 48). These microorganisms inhabit the range of ecological niches provided by corals, such as the surface mucus layer, tissue layers, and the skeleton (6, 28, 30, 45, 48). Symbiotic relationships are based on the mutual exchange and control of nutritional resources. While interactions between *Symbiodinium* and coral hosts are well established, with *Symbiodinium* known to provide over 95% of fixed carbon to at least some coral hosts (42), the functional roles of diverse microbial communities that inhabit the coral holobiont are poorly understood. Documenting the bacterial communities associated with corals and elucidating the functional roles that they play in the corals' multipartner symbioses are the essential first steps in understanding processes involved in the establishment and structure of microbial communities and are critical to understanding the importance of bacterial communities for coral fitness.

Coral reefs have evolved in oligotrophic waters that are particularly poor in nitrogen (15). Although the success of coral reefs in such oligotrophic environments is largely dependent on symbioses between corals and *Symbiodinium*, the growth and abundance of these autotrophic dinoflagellates are also nitrogen limited (17). Gaseous nitrogen (N_2) is present in relatively high concentrations in seawater, and therefore nitrogen fixation, the reduction of N_2 to ammonia, is an important functional role that certain microorganisms, termed diazotrophs, might play in coral symbioses (36, 52, 58). Evidence is accumulating that diazotrophic organisms are more than just passive members of the coral microbial community and may potentially interact in a tight physiological relationship with both the coral animal and its associated microalgae and microbiota (26, 35, 36, 43, 52, 57). For example, acetylene reduc-

tion assays in early studies detected nitrogen fixation within coral samples (52, 58). More recently, cyanobacteria (32, 36) and other bacteria that possess genes for nitrogen fixation (26, 43, 57) have also been detected in coral tissues. Interestingly, Lesser et al. (35) and Olson et al. (43) both found that diazotrophs may have a close relationship with *Symbiodinium*. Lesser et al. (35) observed that the distribution of corals with symbiotic cyanobacteria was positively correlated with depth and suggested that cyanobacteria could sustain *Symbiodinium* nutrition in low-light environments. Olson et al. (43) found that the abundances of *Symbiodinium* and the dominant diazotrophic bacteria, which were closely related to the *Vibrio* genus, were positively correlated. Recent studies have also detected that both endosymbiotic algae and the coral host possess enzymes enabling ammonium assimilation (34, 55, 59); therefore, both could benefit from nitrogen fixation products. Additionally, coral microbiota, including archaea, are potentially involved in ammonium assimilation and other processes of nitrogen cycling, such as nitrification, ammonification, and denitrification (26, 53, 57).

Nitrogen fixation in diazotrophs is facilitated by the highly conserved nitrogenase enzyme complex (61). *nifH* is a segment of the gene that encodes the dinitrogenase iron protein, a subunit of the nitrogenase complex, and provides an ideal molecular target

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for gene-based phylogenetic characterization of diazotrophs because of agreement with 16S rRNA-based phylogeny (23, 56, 60, 61). In addition, there is little evidence of lateral gene transfer (61), which means that retrieval of *nifH* sequences has become a widely used approach for characterizing the diversity of diazotrophic communities in complex samples (12, 24, 38–40, 43, 56, 63).

In this study, we explored the diversity of N₂-fixing bacteria associated with three coral species common on the Great Barrier Reef (GBR), using retrieval and analyses of *nifH* sequences. We tested for species specificity in coral-diazotroph associations through comparative analysis of the diversity of diazotrophic communities among three coral species located on the same reef, among three reef locations for each coral species, and finally within coral microhabitats, namely, mucus and tissue communities.

MATERIALS AND METHODS

Sample collection and processing. Coral tissue and mucus samples were collected from three common coral species, *Acropora millepora*, *Acropora muricata*, and *Pocillopora damicornis*, at three midshelf reefs, Kelso reef (18°25'59"S, 146°59'40"E), Knife reef (18°34'31"S, 147°34'5"E), and Davies reef (18°49'31"S, 147°38'50"E), in the central Great Barrier Reef (GBR) region in January 2009. Two coral colonies (biological replicates) of each species were sampled at each site. All coral colonies sampled were separated by at least 5 meters and were collected from depths of 5 to 10 meters.

To collect coral tissue, branches from the center of colonies were collected and placed in plastic bags underwater. At the surface, branches were first rinsed with autoclaved artificial seawater to remove exogenous microorganism contaminants from the ambient seawater column and mucus and then air brushed (80 lb/in²) to collect coral tissue as a slurry. The tissue slurry was homogenized, aliquoted into cryovials, and centrifuged at 13,000 × *g* to pellet the cellular material (supernatant was removed). Samples were then snap-frozen with liquid nitrogen and stored at –80°C until DNA extraction. Coral mucus was collected from only one of the two replicate colonies (for each species at each site). Mucus was aspirated *in situ* with a syringe and, at the surface, filtered through a Sterivex (0.2- μ m) filter column (Millipore, MA), and frozen at –20°C shipboard before storage at –80°C. Seawater surrounding the sampled area (1 liter) was collected from Knife and Davies reefs and also filtered through Sterivex filters before storage at –80°C.

DNA extraction and purification. Pellets of cellular material derived from coral tissue samples were resuspended in 0.5 ml sucrose extraction buffer (0.75 M sucrose, 40 mM EDTA, 50 mM Tris, pH 8.3), and total DNA was extracted following the protocol described by Bourne et al. (5). Extracted crude DNA was suspended in 30 μ l sterile Milli-Q water and subsequently purified by passage through a 1.2% low-melting-point agarose gel, and high-quality DNA (2 kb) was cut from the gel and cleaned using the QIAquick gel extraction kit (Qiagen, Hilden, Germany), following the manufacturer's instructions. DNA was recovered in 30 μ l sterile Milli-Q water, quantified using the NanoDrop ND1000 (NanoDrop Technologies), and stored at –20°C.

DNA from the coral mucus and seawater samples was extracted following a modified protocol described by Ceh et al. (8) and based on that described by Schauer et al. (50). DNA was purified as outlined for coral tissue samples.

PCR amplification of *nifH* gene and clone library preparation. Clone libraries were prepared for each of the 18 coral tissue, 8 coral mucus, and 2 seawater samples. For clone library construction, a 359-bp fragment of the *nifH* gene (nitrogenase Fe protein gene) was amplified using degenerate primers and the nested PCR approach described by Zehr and McReynolds (62). Briefly, to amplify the *nifH* fragment, 2 μ l of template was added to a first PCR mixture (50 μ l) containing 1× PCR buffer [Tris-Cl, KCl, (NH₄)₂SO₄], 4.0 mM MgCl₂, 0.2 mM each deoxynucleoside triphosphate, 40

ng · μ l⁻¹ bovine serum albumin (New England BioLabs), 2.5 U of *Taq* polymerase (Scientifix), and 0.4 μ M (each) primers NifH3 (5'-ATR TTR TTN GCN GCR TA-3') and NifH4 (5'-TTY TAY GGN AAR GGN GG-3'). PCR amplification was performed in an Eppendorf Mastercycler (Eppendorf, Hamburg, Germany) thermocycler programmed with an initial heating step for 3 min at 95°C, followed by 25 cycles consisting of 95°C for 30 s, 57°C for 30 s, and 72°C for 45 s, followed by a final extension for 7 min at 72°C. After thermal cycling, the products (2 μ l) were subjected to a second round of PCR, identical to the first but with primers nifH1 (5'-TGYGAYCCNAARGCNGA-3') and nifH2 (5'-ANDGCCATCATYTCCNCC-3') and with 30 PCR cycles. For all PCR runs, negative controls were tested and no amplification was obtained.

PCR products from each sample were visualized by electrophoresis, excised, and purified using a QIAquick gel extraction kit (Qiagen) according to the manufacturer's instructions. Purified PCR products were ligated into the TOPO-TA cloning vector (Invitrogen, Carlsbad, CA), with subsequent transformation into competent *Escherichia coli* cells and selection by blue and white screening, according to the manufacturer's instructions. Individual white colonies were transferred to 96-well plates containing LB agar and ampicillin. Plates with the transformants were sealed and submitted to Macrogen Inc. (Seoul, South Korea) for plasmid extraction and sequencing using a model 3739 × I automatic sequencer (Applied Biosystems, Foster City, CA). Previous pilot sequencing and diversity indices demonstrated that the diversity of *nifH* genes in corals is low; therefore, only 48 clones were sequenced from each coral sample (mucus and tissue) from each species, and 96 clones were sequenced for each of the two seawater samples.

Sequencing and phylogenetic analysis. DNA sequences were edited initially to exclude primer and vector sequences using the Sequencher program (Gene Codes Corp., MI). Sequences were imported into the ARB software package (version 5.2; Department of Microbiology, Technical University of Munich [<http://www.arb-home.de/>]), where they were translated into protein amino acid sequences and aligned to create a Phylip-formatted distance matrix. Analysis was performed at the protein level because of the low resolution for identity that exists at the DNA level for this gene. A distance matrix was analyzed using MOTHUR (version 1.20.1; Department of Microbiology and Immunology, The University of Michigan [<http://www.mothur.org/wiki/>]), where deduced amino acid sequences from each library were grouped into operational protein units (OPUs) with a distance threshold of 0.10 (90% similarity between sequences). A representative sequence of each OPU was selected and aligned to the large *nifH* database available (Marine Microbiology, University of California [<http://www.es.ucsc.edu/~wwwzehr/research/database/>]) and maintained in ARB. The OPU were checked against the closest related sequence in GenBank using the online BLASTP function (<http://www.ncbi.nlm.nih.gov/GenBank/index.html>), and if not found in the ARB *nifH* database, they were imported and aligned. Phylogenetic trees from protein amino acid sequences were generated in ARB using a Phylip maximum-likelihood method (Phylip PROML), including bootstrap support of 1,000 replicates.

Statistical analysis. Diversity parameters of the *nifH* gene for each sample (tissue, mucus, and seawater) were generated in MOTHUR (version 1.20.1; Department of Microbiology and Immunology, The University of Michigan [<http://www.mothur.org/wiki/>]) using a 90% protein sequence similarity and included the Chao1 nonparametric richness estimates (9) and the Shannon-Weaver (51) and Simpson (54) indices of diversity. To determine patterns emerging from diazotrophic bacterial profiles from coral tissue, mucus, and seawater samples, principal-component analysis (PCA) was plotted using PAST statistical software (version 2.12) (22; http://palaeo-electronica.org/2001_1/past/issue1_01.htm). For tissue samples, the two biological replicates for each coral species at each site were pooled together. For all data analyses, only OPU representing greater than 5 sequences were included.

Accession numbers. The *nifH* sequences from the representative OPU in this study have been deposited in GenBank under nucleotide and

TABLE 1 Number of OPUs and diversity estimates for *nifH*-deduced protein sequences from mucus and tissue samples of three coral species

Library	No. of clones analyzed	No. of OPUs observed	ACE estimator	Chao1 estimator	Simpson index	Shannon-Weaver index
<i>Acropora millepora</i>						
Tissue	280	6	6	6	0.24	1.51
Mucus	145	10	10.19	11	0.23	1.70
<i>Pocillopora damicornis</i>						
Tissue	257	15	70.51	22.50	0.15	2.07
Mucus	133	6	6	6	0.22	1.60
<i>Acropora muricata</i>						
Tissue	230	17	38.43	24	0.17	1.99
Mucus	107	8	11.21	11	0.23	1.59
Seawater	197	23	33.22	34.25	0.26	1.96

protein sequence accession numbers [JN601397](#) to [JN601421](#) and [AEU12166](#) to [AEU12190](#), respectively.

RESULTS

Diversity of *nifH* sequences. A total of 1,344 high-quality *nifH* sequences were retrieved from coral mucus, coral tissue, and surrounding seawater samples. Retrieved *nifH* sequences had low relative diversity, and therefore sequences from replicate sample types were grouped, resulting in a total of 26 distinct OPUs across all samples. For mucus samples, between 6 and 10 OPUs were retrieved, with many being shared by 2 or more of the 3 coral species. Richness estimators (ACE and Chao1) revealed that the sampling effort was sufficient and that the diversity of the *nifH* sequences was well covered in these mucus samples (Table 1). Shannon-Weaver and Simpson indices provided corroborative evidence that the diversity in coral mucus samples was low, with values ranging from only 1.59 to 1.70 and 0.22 to 0.23, respectively, among coral species. Diversity parameters for coral tissue samples displayed more variability than those for mucus samples, with between 6 and 17 OPUs identified, many of which were shared across coral species. Interestingly, tissue samples of *A. millepora* revealed the lowest number of OPUs (6 OPUs) and the lowest Shannon-Weaver index (1.5), despite having the highest

sequencing effort (280 retrieved *nifH* sequences). In contrast, *Pocillopora damicornis* (15 retrieved OPUs) and *Acropora muricata* (17 OPUs) showed a greater diversity and had the highest ACE and Chao estimator values, suggesting that the full diversity of the *nifH* gene was not encompassed by the sampling effort for these two species. Not surprisingly, *nifH* gene diversity in seawater samples was the highest overall, with 23 OPUs recovered. The high diversity of *nifH* sequences recovered from seawater samples was also observed in rarefaction analysis, which did not reach saturation (see Fig. S1 in the supplemental material). In contrast, mucus samples appeared to be highly saturated, and tissue samples approached an asymptote with sampling effort (see Fig. S1 in the supplemental material).

Phylogeny of *nifH* sequences. Comparisons of the class composition of diazotrophic bacteria among sample types (Fig. 1) and phylogenetic analysis (Fig. 2) both revealed that diazotrophic bacterial communities differed among coral tissue, coral mucus, and seawater libraries. Mucus libraries comprised very heterogeneous diazotrophic types and shared several OPUs (OPUs 1, 2, 16, 17, and 18) with seawater libraries (Fig. 2). In contrast, the *Alphaproteobacteria* class dominated coral tissue libraries combined for the three coral species (Fig. 1), and these did not share any OPUs with

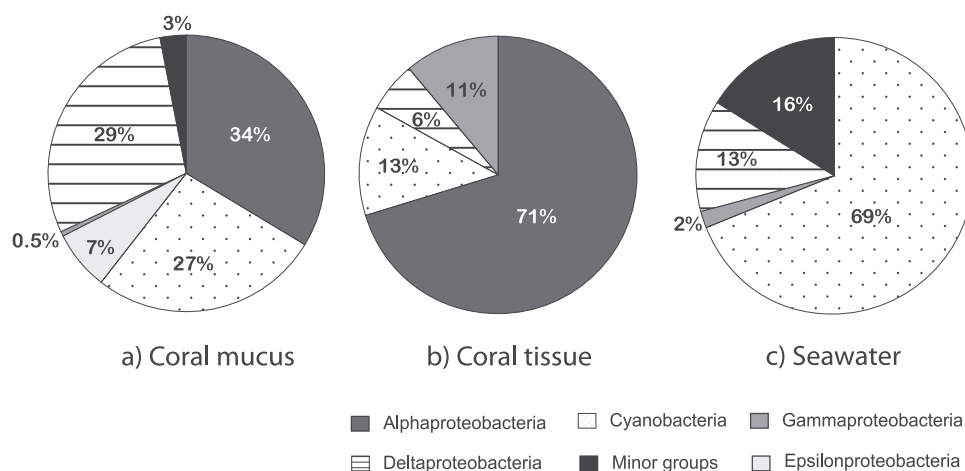


FIG 1 Class-level compositions of coral *nifH* clone libraries. Percent abundances per clone library from coral mucus (414 clones) (a), coral tissue sequences (767 clones) (b), and seawater (195 clones) (c) are shown. Minor groups represent OPU that had fewer than 5 sequences and were not further analyzed.

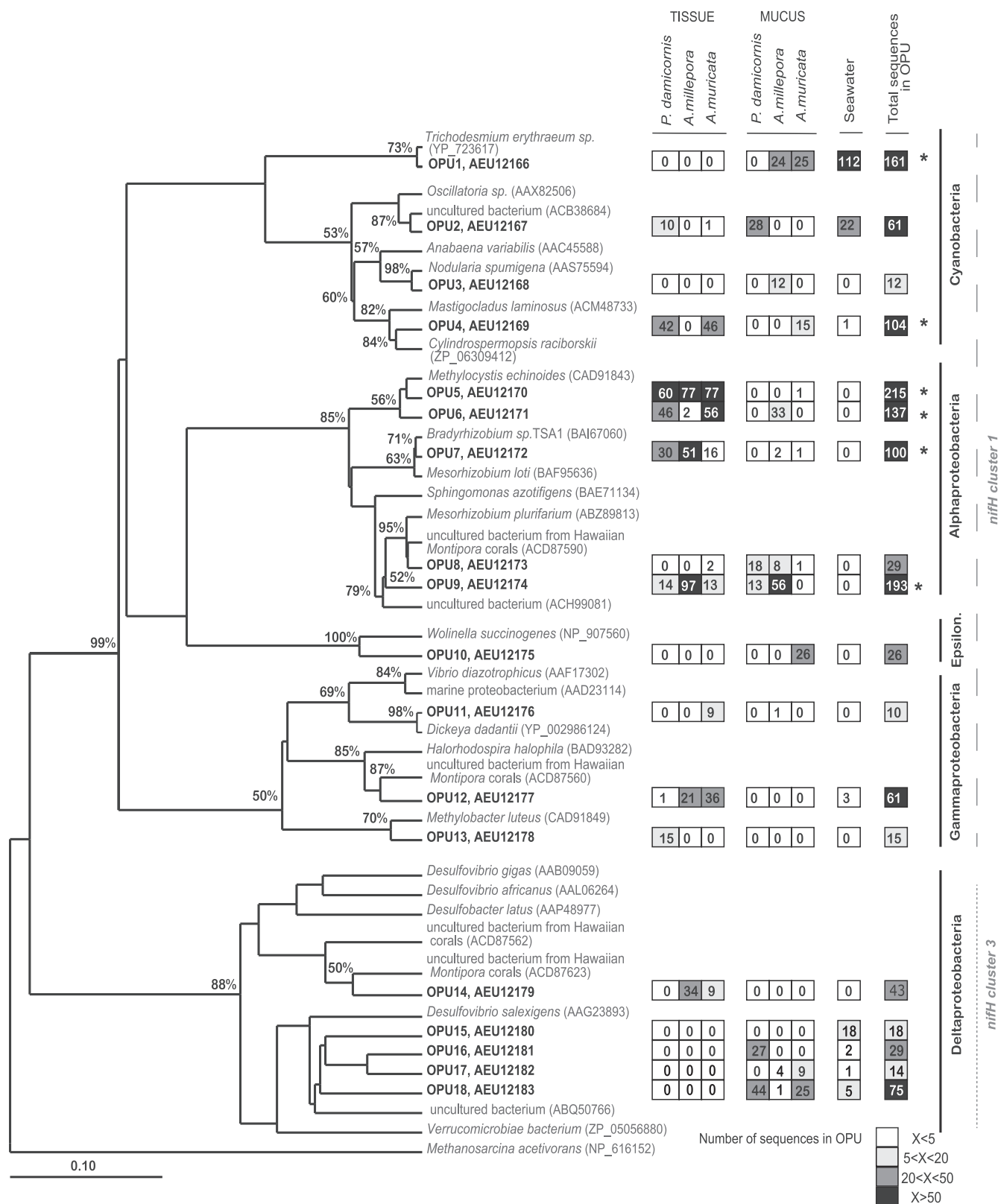


FIG 2 Phylogeny and composition of the *nifH* operational protein units (OPU) representing more than 10 sequences (cutoff, 0.1) ($n = 1,347$ sequences). OPUs from this study are represented in bold with associated protein accession numbers. The maximum-likelihood phylogenetic tree was generated in ARB and includes bootstrap support of 1,000 iterations. The tree is rooted with the *nifH* protein sequence of an archaeon (*Methanosarcina acetivorans*). The heat map to the right of the tree represents the total number of sequences retrieved for each OPU for each coral species (mucus and tissue separately) and seawater, with highest values (>50) in black and lowest (<5) in white. OPUs representing over 100 sequences are indicated with asterisks. Sequences from known phylogenies are indicated by species names and protein GenBank accession numbers in parentheses. Classes of bacteria are indicated to the far right of tree (Epsilon. stands for *Epsilonproteobacteria*), and *nifH* clusters are indicated with different dashed lines.

TABLE 2 Affiliations of representative *nifH*-deduced protein sequences grouped into OPUs (all samples pooled) and which contain 10 or more sequences

Total no. of sequences in OPU	Source OPU, protein accession no.	Closest relative (accession no.)	Alignment (bp)	Similarity (%) ^a	Taxonomic description
161	OPU 1, AEU12166	<i>Trichodesmium erythraeum</i> (YP_723617)	117/118	99	<i>Cyanobacteria</i>
61	OPU 2, AEU12167	Uncultured bacterium (ACB38684)	116/118	98	<i>Cyanobacteria</i>
12	OPU 3, AEU12168	<i>Nodularia spumigena</i> (AAS75594)	106/108	98	<i>Cyanobacteria</i>
104	OPU 4, AEU12169	<i>Cylindrospermopsis raciborskii</i> (ZP_06309412)	114/119	96	<i>Cyanobacteria</i>
215	OPU 5, AEU12170	<i>Methylocystis echinoides</i> (CAD91843)	115/118	97	<i>Alphaproteobacteria</i>
137	OPU 6, AEU12171	<i>Methylocystis echinoides</i> (CAD91843)	115/118	97	<i>Alphaproteobacteria</i>
100	OPU 7, AEU12172	<i>Bradyrhizobium</i> sp. (BAI67060)	117/118	99	<i>Alphaproteobacteria</i>
29	OPU 8, AEU12173	Uncultured bacterium from Hawaiian <i>Montipora</i> corals (ACD87590)	115/118	97	<i>Alphaproteobacteria</i>
193	OPU 9, AEU12174	Uncultured bacterium (ACH99081)	116/118	98	<i>Alphaproteobacteria</i>
26	OPU 10, AEU12175	<i>Wolinella succinogenes</i> (NP_907560)	118/132	89	<i>Epsilonproteobacteria</i>
10	OPU 11, AEU12176	<i>Dickeya dadantii</i> (YP_002986124)	118/119	99	<i>Gammaproteobacteria</i>
61	OPU 12, AEU12177	Uncultured bacterium from Hawaiian <i>Montipora</i> corals (ACD87560)	114/119	96	<i>Gammaproteobacteria</i>
15	OPU 13, AEU12178	<i>Methylobacter luteus</i> (CAD91849)	114/119	96	<i>Gammaproteobacteria</i>
43	OPU 14, AEU12179	Uncultured bacterium from Hawaiian <i>Montipora</i> corals (ACD87623)	112/119	94	<i>Deltaproteobacteria</i>
18	OPU 15, AEU12180	<i>Desulfovibrio salexigens</i> (AAG23893)	96/109	88	<i>Deltaproteobacteria</i>
29	OPU 16, AEU12181	Uncultured bacterium (ABQ50766)	103/119	87	<i>Deltaproteobacteria</i>
14	OPU 17, AEU12182	Uncultured bacterium (ABQ50766)	103/119	87	<i>Deltaproteobacteria</i>
75	OPU 18, AEU12183	Uncultured bacterium (ABQ50766)	119/119	100	<i>Deltaproteobacteria</i>

^a Protein sequences were aligned to their closest relative using protein BLAST (Blastp) results for affiliations.

seawater libraries (Fig. 2), which were dominated by *Cyanobacteria* (Fig. 1).

Coral mucus-derived sequences were taxonomically diverse, with the majority of sequences distributed among the *Cyanobacteria*, *Deltaproteobacteria*, and *Alphaproteobacteria* classes (Fig. 1). In these three classes of bacteria, the most abundant OPU in terms of sequence numbers were OPU 18 (71 sequences), which demonstrated 100% amino acid sequence identity with an uncultured member of *Deltaproteobacteria* (ABQ50766), OPU 9 (69 sequences), which was closely related to an uncultured member of *Alphaproteobacteria* (ACH99081) (98% similarity), and OPU 1 (50 sequences), which was closest (99% similarity) to the *Trichodesmium erythraeum nifH* sequence (Fig. 2; Table 2). Of the remaining mucus-derived sequences, only one was found in the *Gammaproteobacteria* class, and 26 sequences were found in OPU 10, the only OPU belonging to the *Epsilonproteobacteria* class (Fig. 1 and 2). These last sequences had 89% similarity to a *Wolinella succinogenes nifH* sequence (Table 2).

The seawater libraries were dominated by *Cyanobacteria* (Fig. 1), with more than half of the total sequences (112 of 192 sequences) belonging to OPU 1, which was affiliated with *Trichodesmium erythraeum* (99% amino acid sequence identity), and 22 other sequences being affiliated with an uncultured member of *Cyanobacteria* (ACB38684) (Fig. 2; Table 2). The remaining seawater sequences were found in the *Deltaproteobacteria* class (*nifH* cluster 3), with OPU 15, which had the highest numbers of seawater sequences in this class, being only distantly affiliated with *Desulfovibrio salexigens nifH* (88% protein sequence similarity) (Fig. 2 and Table 2). Sixteen percent of seawater libraries were represented by single retrieved clones, further indicating high sequence diversity in seawater samples (Fig. 1), though this diversity was not explored further.

Seventy-one percent of sequences retrieved from coral tissues (539 sequences) belonged to the *Alphaproteobacteria* class (Fig. 1). Interestingly, all *Alphaproteobacteria* sequences were found in only 4 OPUs (OPU 5, OPU 6, OPU 7, and OPU 9), and these OPU were all affiliated with *Rhizobiales*, a group of symbiotic diazotrophs that are found in the root nodules of legume plants (Fabaceae) (Fig. 2). OPU 7 (97 tissue sequences) was most closely affiliated (99%) to the *nifH* amino acid sequence of *Bradyrhizobium* sp. strain TSA1, and OPU 9 (124 tissue sequences) was closest taxonomically to an uncultured *Alphaproteobacteria* (ACH99081) *nifH* sequence (Fig. 2; Table 2). Interestingly, OPU 5 (214 tissue sequences), which was the most abundant OPU in all libraries, and OPU 6 (104 tissue sequences), which was the third most abundant group, were both affiliated (97% amino acid identity) with a known methanotroph (type I), *Methylocystis echinoides*. OPU 13, a member of *Gammaproteobacteria*, was also closely affiliated with the methanotroph (type II) *Methylobacter luteus* (CAD91849) *nifH* sequence, although this OPU comprised only 15 retrieved sequences. The remaining 29% of coral tissue-derived *nifH* sequences were distributed among 3 classes, the *Cyanobacteria*, the *Gammaproteobacteria*, and the *Deltaproteobacteria* (Fig. 1). Eighty-eight sequences derived from coral tissues were closely related (96% sequence identity) to a *nifH* sequence from the freshwater cyanobacterium, *Cylindrospermopsis raciborskii*. Interestingly, OPU 14 and OPU 12 were both most closely related to sequences that were found in the only other study to investigate *nifH* sequences in corals (43) (Fig. 2; Table 2). Another OPU that was closely related to a sequence found by Olson et al. (43) was OPU 8, which constituted 36 mucus-derived sequences and 2 tissue-derived sequences.

Specificity of coral diazotrophic assemblages. PCA compari-

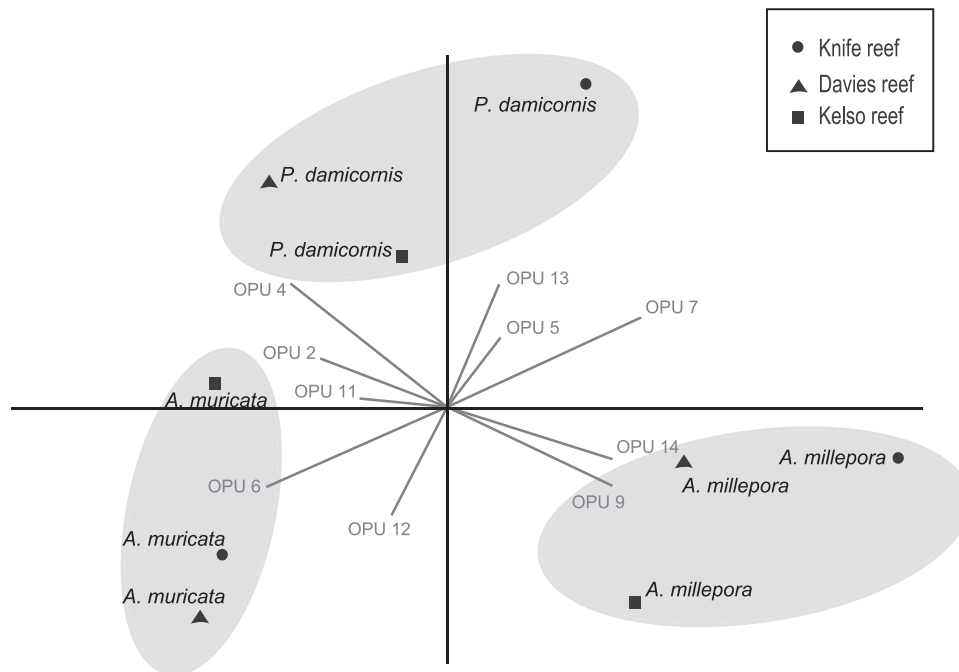


FIG 3 PCA biplot representing relative abundances of *nifH*-deduced protein sequences derived from tissue clone libraries of three coral species sampled from each of three midshelf reefs. Gray lines show vectors from representative OPUs that drive the differences among clone libraries. The corresponding taxonomic affiliation and GenBank accession number of each OPU can be found in Fig. 2 and Table 2. For this analysis, only OPUs with more than 5 sequences are represented.

son of diazotroph diversity associated with replicate coral tissue from the three coral species (Fig. 3) and with mucus from the three coral species and seawater samples from reefs samples (see Fig. S2 in the supplemental material) revealed clear patterns in bacterial communities among coral species. Tissue samples derived from colonies of the same coral species grouped together regardless of reef sampling location, with the first two components explaining 65% of the variation (Fig. 3). Although tissue samples from the three species were dominated by rhizobium-like sequences and all possessed similar abundances of sequences grouped in OPU 5, the relative abundances of other coral species-specific OPUs separated diazotrophic communities into three distinct groups corresponding to the three species (Fig. 3). For example, OPU 13 (15 sequences) was retrieved only from *P. damicornis* libraries, whereas OPU 11 (9 sequences) was retrieved only from *A. muricata* tissue libraries (Fig. 2 and 3). Both *P. damicornis* and *A. muricata* libraries contained sequences belonging to OPU 4, which was affiliated with a cyanobacterium closely related to *Cylindrospermopsis raciborskii*, but no cyanobacterial sequences were retrieved from *A. millepora* tissue samples. Sequences closely related to a methanotroph-derived *nifH* sequence grouped in OPU 6 and were abundant in both *P. damicornis* and *A. muricata* tissue libraries. *A. millepora* samples were clearly separated from those of the two other coral species in the PCA by their higher abundance of sequences grouped in OPU 9 and OPU 14. In addition, *A. millepora* and *P. damicornis* sequences were correlated strongly with OPU 7, while *A. millepora* and *A. muricata* sequences were correlated with OPU 12.

In contrast to patterns found for tissue samples, PCA analysis of retrieved *nifH* sequences derived from mucus and seawater samples showed no distinct patterns, with no evidence of mucus

libraries grouping in relation to either coral species or reef location (see Fig. S2 in the supplemental material). Interestingly, the most abundant *nifH* sequences from seawater, which grouped in OPU 1 and were affiliated with a *Trichodesmium* sp., were also abundant in mucus libraries of *A. millepora* and *A. muricata* from Knife reef (Fig. 2). The presence of this OPU in these mucus samples likely represents contamination with seawater and results in these samples being separated from other mucus sequences in the PCA (see Fig. S2 in the supplemental material). The first two components of the PCA explained 40% of the variation in mucus and seawater clone library sequences.

DISCUSSION

Coral tissues are dominated by rhizobia. Phylogenetic analysis of *nifH* gene clone libraries derived from tissues of three common coral species on the GBR revealed that diazotrophic communities are characterized by low diversity and a striking dominance of phylotypes from the *Alphaproteobacteria* class, with 71% of the total *nifH* sequences ($n = 767$ sequences) retrieved from coral tissues belonging to this class. Interestingly, all *Alphaproteobacteria* phylotypes were closely related to bacterial species belonging to the *Rhizobiales* order (between 97% and 99% amino acid identity) and were found in all three coral species investigated. Rhizobia are soil bacteria that inhabit nodules in the roots of legume plants. Rhizobia fix nitrogen, enabling plants to thrive and reproduce in nitrogen-poor environments, and in return, the rhizobia receive carbon and amino acids (19, 37). Rhizobium-like ribotypes have also been documented in 16S rRNA gene-based surveys of other coral species (29, 41, 47), as well as in functional gene arrays and metagenomic studies (26, 57), highlighting the potential importance of this group of bacteria for supplementing the nutritional

requirements of the coral holobiont in nitrogen-limited oligotrophic waters. Plant rhizobia require specific signals to enter into symbiosis and for nitrogen fixation (11, 18). Similarly, close symbiotic pathways between diazotrophs and *Symbiodinium* and/or the coral could exist, as both are capable of ammonium assimilation (34, 55, 59). In plants, the cortex layer of nodules protects rhizobia from high concentrations of oxygen by acting as an oxygen diffusion barrier, while the plants' leghaemoglobin facilitates diazotroph respiration at low oxygen concentrations (16). It is currently unclear how rhizobia might be protected from high concentrations of oxygen arising from dinoflagellate photosynthesis in coral tissues (31). Coral microhabitats including bacterial aggregates within the gastrodermis, as reported by Ainsworth and Hoegh-Guldberg (2), or microaerophilic regions in the gastrovascular cavities of coral polyps, as observed by Agostini et al. (1), may host diazotrophic communities under oxygen-depleted conditions.

Rhizobium-affiliated sequences were clustered into four identified OPU (OPUs 5, 6, 7, and 9). OPU 7 was closely related (99% amino acid similarity) to *Bradyrhizobium* spp., one of the most commonly occurring rhizobia that form symbioses in the nodules of legume plants. OPU 9 was closest to an uncultured bacterium (ACH99081) that is affiliated with a mesorhizobium. Interestingly, OPUs 5 and 6 were affiliated with a *nifH* sequence of *Methylocystis echinoides*, a methanotroph type II organism that also belongs to the *Rhizobiales* group. Moreover, OPU 5 represented the largest number of sequences retrieved and derived from all coral species sampled (215 sequences). The ability to fix nitrogen is an important phenotypic trait of most currently known methanotrophic bacteria, and *nifH* has been used to distinguish representatives at the clade level (4, 14). There are only a few reports of the presence of methanotrophs in corals. Using a functional gene array, Kimes et al. (26) found several genes involved in methane oxidation, with some belonging to diverse methanotrophs from type I and II and also from archaea. Siboni et al. (53) investigated archaea associated with the mucus of *Acanthastrea*, *Favia*, and *Fungia* species and found that 8% of the euryarchaeotal sequences were affiliated with anaerobic methanotroph type II organisms. In our study, a methanotroph type I (OPU 13) was also found, although only in *P. damicornis* tissue libraries. Our finding that this type of bacteria is the dominant diazotrophic group in three common coral species is an indication that methane is potentially abundant in coral tissues. Accordingly, our results highlight the need for further investigations of the fate of methane in the coral holobiont.

Diazotrophs from coral tissues are specific to coral species. Although all coral species had high abundances of rhizobium-like sequences in their tissues, PCA of *nifH* sequences grouped coral samples from 3 different reefs according to coral species. We conclude that differences in the relative abundances of OPUs were responsible for grouping diazotrophic communities according to coral species and that diazotrophic communities display some coral species-specific associations. Previous studies targeting the 16S rRNA gene have reported that bacterial communities on a coral species are similar at geographically separated locations (5, 46, 47; K. B. Ritchie and G. W. Smith, presented at the 8th International Coral Reef Symposium, Panama City, Panama, 1997). These studies have contributed to the development of the coral holobiont model, in which the coral is viewed as a structured symbiotic system composed of the animal host and an array of

microscopic partners, including bacteria (28, 44, 48). This study is the first to report that bacteria with a specific functional role can also be specific to a coral host, and it therefore provides additional support for the holobiont model. It is important to note that all reefs sampled in our study were mid- or outer-shelf reefs separated by distances ranging from 50 km to 200 km, and they thus experienced similar water quality parameters. Accordingly, it is possible that diazotrophic communities from coral species might differ between more distant reefs, particularly between locations with differing nutrient inputs, such as inshore versus offshore reefs. Findings that bacterial communities associated with corals differ among locations that differ in environmental characteristics (27, 44) highlight the need for further investigations of coral diazotrophic communities exposed to different water quality parameters.

The ephemeral nature of coral mucus. In contrast to the consistent patterns found for diazotrophic communities associated with coral tissues, a variety of *nifH* groups were retrieved from coral mucus samples, and these belonged to five different classes of bacteria (*Alphaproteobacteria*, *Gammaproteobacteria*, *Epsilonproteobacteria*, *Deltaproteobacteria*, and *Cyanobacteria*). The differences in the consistency of diazotrophic communities between coral tissue and mucus samples found in our study are corroborated by previous 16S rRNA studies, which also found differences in general bacterial communities between coral tissue and mucus samples (6, 20, 21, 30, 33, 47). The overlap found between the coral mucus microbiota and that of the surrounding seawater is not surprising, as small amounts of seawater are taken up when coral mucus samples are collected *in situ*. Indeed, the highest number of sequences recovered from mucus grouped in OPU 1, which also possessed the highest number of seawater-derived sequences and were closely affiliated to a *nifH* sequence derived from *Trichodesmium erythraeum*. In addition, PCA analysis showed no relationship between *nifH* gene diversity recovered from mucus libraries and either coral species or location (see Fig. S1 in the supplemental material). The location of mucus bacterial communities at the interface between seawater and coral tissues undoubtedly promotes continuous exchange and contributes to the ephemeral, complex, and dynamic nature of mucus microbial communities (7, 25). Indeed, bacterial groups within coral mucus exhibit high spatial heterogeneity because of the many different factors that may have an impact on this habitat, such as various degrees of light exposure, nutrient availability, sedimentation, mucus age, and competition with other microbes (3, 13, 21, 49).

A large number of *nifH* cluster III sequences (10), all closely related to sulfate reducers, including *Desulfovibrio* spp., represented the second largest group retrieved from mucus samples in our study and were also found in coral tissues (OPU 14). Kimes and colleagues (26) determined that 22% of *dsr* genes in corals belong to the *Deltaproteobacteria* subclass and suggested that inorganic sulfate might be another potential source of sulfur for the coral holobiont. Interestingly, *nifH* sequences close to *Desulfovibrio* and other uncultured sulfate reducers were also found in the only other study that investigated *nifH* in corals (43). As found by Olson and colleagues (43), our coral-derived *nifH* sequences were closest to *Desulfovibrio salexigens* and had low amino acid identity (only 87 to 88%) with all other sequences, indicating the presence of potentially novel species. OPU 14, which was the only *Deltaproteobacteria nifH*-affiliated sequence derived from coral tissue in this study, had a 94% similarity to a *nifH* sequence from the study

by Olson et al. (43). Interestingly, other OPUs were observed to be closely affiliated with *nifH* sequences retrieved from corals from Hawaii, including OPU 12, a tissue gammaproteobacterium, and OPU 8, a mucus alphaproteobacterium. The similarity of ribotypes between this and other studies further supports that some diazotrophic communities are consistent in corals and therefore that they represent important members of the coral holobiont symbiosis.

Conclusions. Tissues of three common coral species on the GBR displayed species-specific patterns in diazotrophic communities, suggesting that this bacterial group is likely to have a symbiotic role in the coral holobiont. The presence of consistent and dominant populations of rhizobia in all three coral species at all reefs suggests that coral rhizobia are important symbiotic members of coral tissues, although further studies are required to demonstrate that symbionts are actively fixing nitrogen. The specificity of tissue diazotrophs to each of the coral species provides further support for the holobiont model of coral symbioses. In contrast, coral mucus is a complex microbial microhabitat with an ephemeral nature, given its location at the interface between coral tissues and seawater and potential for microbial exchange, which appears to give rise to a diverse and unstructured diazotrophic community. Overall consistency in the identification of diazotrophic *nifH* sequences associated with corals in our and other studies suggests that these microbial groups may be essential in nitrogen cycling within the coral holobiont. Importantly, changes in diazotrophic communities could directly reflect shifts in environmental parameters, such as nutrient inputs, and could be used to detect changes in coral fitness in response to environmental change.

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