Effects of Ocean Acidification on Resident and Active Microbial Communities of *Stylophora pistillata*

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Ocean warming and ocean acidification (OA) are direct consequences of climate change and affect coral reefs worldwide. While the effect of ocean warming manifests itself in increased frequency and severity of coral bleaching, the effects of ocean acidification on corals are less clear. In particular, long-term effects of OA on the bacterial communities associated with corals are largely unknown. In this study, we investigated the effects of ocean acidification on the resident and active microbiome of long-term aquaria-maintained *Stylophora pistillata* colonies by assessing 16S rRNA gene diversity on the DNA (resident community) and RNA level (active community).

Coral colony fragments of *S. pistillata* were kept in aquaria for 2 years at four different pCO₂ levels ranging from current pH conditions to increased acidification scenarios (i.e., pH 7.2, 7.4, 7.8, and 8). We identified 154 bacterial families encompassing 2,047 taxa (OTUs) in the resident and 89 bacterial families including 1,659 OTUs in the active communities. Resident communities were dominated by members of Alteromonadaceae, Flavobacteriaceae, and Colwelliaceae, while active communities were dominated by families Cyclobacteriaceae and Amoebophilaceae. Besides the overall differences between resident and active community composition, significant differences were seen between the control (pH 8) and the two lower pH treatments (7.2 and 7.4) in the active community, but only between pH 8 and 7.2 in the resident community. Our analyses revealed profound differences between the resident and active microbial communities, and we found that OA exerted stronger effects on the active community. Further, our results suggest that rDNA- and rRNA-based sequencing should be considered complementary tools to investigate the effects of environmental change on microbial assemblage structure and activity.

Keywords: climate change, coral microbiome, 16S rDNA, 16S rRNA, coral holobiont

INTRODUCTION

Coral reefs are among the most biodiverse and productive ecosystems in the world. Despite covering less than 0.2% of the marine environment, they harbor 34% of the described marine biodiversity (Reaka-Kudla, 2001). Hundreds of millions of people depend on them for their livelihood, since they provide goods and services such as fisheries, tourism, coastal protection,
pharmaceutical substances and aesthetic and cultural value (Moberg and Folke, 1999). The framework of reefs is built by scleractinian corals. They are referred to as holobionts, obligate symbiotic organisms that live in close association with microorganisms such as endosymbiotic photosynthetic algae, bacteria, archaea, viruses and fungi (Rohwer et al., 2002; Rosenberg et al., 2007). Many studies have looked into the symbiotic relationship with the dinoflagellate algae from the family Symbiodiniaceae (LaJeunesse et al., 2018), but less is known about the bacterial partners in this association. The coral microbiome has a range of functional roles, including nitrogen fixation (Lesser et al., 2004; Lema et al., 2012; Pogoreutz et al., 2017a,b), sulfur cycling (Raina et al., 2009), antibacterial activities (Ritchie, 2006) and overall holobiont health (Rosenberg et al., 2007; Robbins et al., 2019; Voolstra and Ziegler, 2020). As high throughput sequencing of microbial DNA becomes more accessible, studies based on analysis of phylogenetic markers such as the 16S rRNA gene become increasingly available.

Coral reefs are declining globally, mainly due to rising CO₂ emissions (Hoegh-Guldberg, 2007; Hughes et al., 2017). Besides causing ocean warming, rising CO₂ in the atmosphere is also linked to ocean acidification (OA), which is caused by elevated partial pressure of carbon dioxide (pCO₂) in the seawater that alters carbonate ion chemistry and reduces seawater pH (Doney et al., 2009). Seawater surface pH has decreased 0.1 units since pre-industrial values and is predicted to decrease another 0.2–0.4 units by 2100 (Rhein, 2014). There is evidence that OA can negatively impact calcifying organisms, such as scleractinian corals, by decreasing their calcification rate and photosynthesis (Anthony et al., 2008; Doney et al., 2009). However, it is still not clear how OA might affect the coral microbiome, and consequently holobiont physiology and health. Some studies showed that OA can shift coral microbiome composition from a healthy-associated community to one usually seen in diseased and stressed corals (Thurber et al., 2009; Meron et al., 2011; Webster et al., 2013), while others show a stable microbiome (Meron et al., 2012; Webster et al., 2016) or species-specific responses (Morrow et al., 2015; Grottoli et al., 2018).

Most studies that characterized microbial community composition were based on 16S rDNA amplicon sequencing, an established method that can successfully describe microorganisms present in the community (Lane et al., 1985; Weisburg et al., 1991). However, one limitation of this method is that it includes any DNA, including DNA from dormant and dead microbes, making it impossible to distinguish the metabolically active from the non-active population (Pietramellara et al., 2009; Blagodatskaya and Kuzyakov, 2013; Carini et al., 2016). Alternatively, sequencing 16S rRNA serves as an effective proxy to estimate active microbial populations, since it is a component of ribosomes, which are predicted to increase in number with increasing metabolic activity (Poulsen et al., 1993). In addition, once outside the cell, RNA degrades faster than DNA, ensuring sampling of only the active community. Thus, the ratio of rRNA/rDNA of a specific taxon can be used as a measure of its potential activity (Jones and Lennon, 2010), although this approach is not without its caveats (Blazewicz et al., 2013). Many studies have pointed out a mismatch between the DNA/RNA based communities (DeAngelis and Firestone, 2012; Blott et al., 2015; Steven et al., 2017). Hence, simultaneous sequencing and comparison of both components can unveil rare but highly active, as well as abundant and inactive taxa (Jones and Lennon, 2010; Campbell et al., 2011; Gaidos et al., 2011; Lanzén et al., 2011; Hunt et al., 2013).

In this study, we assessed the effect of long term increased pCO₂ treatments on the microbial community of the Red Sea coral Stylophora pistillata at both the 16S rDNA and rRNA level to obtain an integrative view of the effects on the resident and active microbial partners.

**MATERIALS AND METHODS**

**Coral Husbandry**

Nubbins of *Stylophora pistillata* have been kept in a long-term experimental aquarium setup since the early 2010’s at the Monaco Scientific Centre as described previously (Venn et al., 2013; Tambutté et al., 2015; Liew et al., 2018) and were sampled for this study in April 2014, after 2 years of OA treatment. Coral nubbins were exposed to four different pCO₂ treatments: 2,447.48 ± 7.16, 2,461.77 ± 5.97, 2,474.16 ± 5.89 and 2,496.62 ± 6.99 μmol kg⁻¹ total alkalinity. These treatments were equivalent to: pH 8.0 (control), 7.8, 7.4 and 7.2, respectively. Colonies were divided into four fragments and placed under the different treatments, with three fragments sampled from each treatment. Carbonate chemistry was manipulated by bubbling CO₂ to decrease pH to the desired target values (pCO₂ levels: 537.66 ± 4.88, 798.77 ± 14.36, 2109.3 ± 59.63, 3513.15 ± 67.20 μatm). Temperature and pH were monitored in the four treatment tanks with probes (Ponsel-Mesure, France) linked to a monitoring system (Enoleo, Monaco) that also controlled CO₂ bubbling. Mediterranean water was supplied at an exchange rate of 70% h⁻¹, adjusted to 38 g liter⁻¹ salinity and 25°C temperature. Light intensity was set to 170 μmol photons m⁻² s⁻¹ with a 12 h light/12 h dark photoperiod with HQI10000K metal halide lamps (BLV Nepturion). Total alkalinity (TA) and pH were measured daily, and carbonate seawater chemistry was calculated as described previously (Tambutté et al., 2015).

**DNA and RNA Extraction, Reverse Transcription, and 16S rRNA Gene and Transcript Amplicon Sequencing**

*Stylophora pistillata* DNA was extracted with DNeasy PowerBiofilm (Qiagen, Hilden, Germany) following manufacturer's instructions. Briefly, small pieces from nubbins of *S. pistillata* were collected in PowerBiofilm Bead tubes and homogenized with CryoMill (Retsch, Germany). Triplicates were extracted for each pH condition. Following centrifugation, supernatants were passed through silica spin filter tubes and after washing steps, DNA was eluted in 10 mM Tris.

Total RNA was isolated from triplicates collected for each pH treatments using RNeasy kit (Qiagen) according to manufacturer's instructions. First-strand cDNA was synthesized
Operational Taxonomic Units (OTUs) were defined based on the number of sequences possible while still keeping all replicates. After subsampling to 10,669 sequences, which was the highest number that allowed keeping all replicates. Sequences were reclassified against the Silva database (McDonald et al., 2012). Sequences were then classified against the Silva database using the remove.lineages command in mothur. Sequences were further excluded with UCHIME (Edgar et al., 2011), as were chloroplast, mitochondrial, archaeal, and eukaryotic sequences. Chimeric sequences were exclusion of singletons, sequences were aligned to the Silva database (Pruesse et al., 2007). Chimeric sequences were handled by removing LINEAGE and removing chimeras. Sequences were further excluded with UCHIME (Edgar et al., 2011), as were chloroplast, mitochondrial, archaeal, and eukaryotic sequences. Chimeric sequences were excluded with UCHIME (Edgar et al., 2011), as were chloroplast, mitochondrial, archaeal, and eukaryotic sequences. Chimeric sequences were then classified against the Silva database (McDonald et al., 2012). Sequences were reclassified against the Silva database after subsampling to 10,669 sequences, which was the highest number of sequences possible while still keeping all replicates. Operational Taxonomic Units (OTUs) were defined based on sequence clustering at a 97% similarity cutoff and used for following analysis.

We used R (version 3.5.1, 2018) and PRIMER E v6 (PERMANOVA+) software for multivariate analysis on Log (x + 1) transformed OTU count data (Clarke and Gorley, 2006). Difference between groups according to genetic material (i.e., DNA vs. RNA) and pH treatment were visualized with non-metric MultiDimensional Scaling (nMDS) plots, based on Bray-Curtis dissimilarity distances. One-factorial PERMutational ANOVA (PERMANOVAs) were used to test for differences between DNA vs. RNA material and pH treatments. We also performed an analysis of similarity percentages (SIMPER; Clarke, 1993) based on Bray Curtis resemblance matrix to identify OTUs mainly responsible for dissimilarity between groups. Alpha diversity indices (Chao1, Inverse Simpson, Simpson Evenness) were calculated for each sample in mothur. Samples were tested for normality and homoscedasticity with Shapiro-Wilk and Levene's test, before being analyzed for differences between pH treatments using ANOVA with Tukey post hoc tests in R (version 3.5.1, 2018). The putative core microbiome of S. pistillata was defined by identifying bacterial OTUs present in all treatment and replicates. Significantly differences in OTU abundance amongst treatments were analyzed by the linear discriminant analysis (LDA) effect size (LEfSe) method (Segata et al., 2011) on the website https://huttenhower.sph.harvard.edu/galaxy/root (LDA > 2).

RESULTS

Sample and Sequencing Overview

Illumina MiSeq sequencing of bacterial 16S rDNA and rrRNA amplicons generated a total of 4,157,907 sequences from 24 samples [average 173,247 ± 100,095 (SD)], being 1,482,981 for RNA samples and 2,674,938 for DNA samples (Supplementary Table 1). After quality filtering and exclusion of chimeras and unwanted sequences, 1,474,649 remained, with an average of 61,444 ± 38,313 (SD). From these, 14,829 distinct sequences were identified after clustering at 97% similarity and assigned to bacteria. Subsampling was done to 10,669 sequences, which was the highest number that allowed keeping all replicates. After quality filtering, DNA samples contained an average of 76,488 sequences (range from 13,735 to 165,630), while RNA samples contained an average of 46,399 sequences (range from 10,669 to 114,558, Supplementary Table 1). Rarefaction curves for DNA and RNA samples suggested that subsampling to 10,669 was indeed sufficient to capture most of bacterial diversity (Supplementary Figure 1A).

Distinct Bacterial Communities Based on 16S rDNA and 16S rrRNA Sequencing

We identified a total of 154 bacterial families for DNA- and 89 bacterial families for RNA-based sequencing. DNA samples were dominated by members of Alteromonadaceae (24.52%), Flavobacteriaceae (24.17%) and Colwelliaceae (18.51%), while RNA samples were dominated by families Cyclobacteriacea and

Library Preparation and Sequencing

Triplicate PCRs for each sample were pooled, and cleaned using the Mini Elute PCR purification kit (Qiagen, Hilden, Germany), following manufacturer instructions. PCR products were indexed with Nextera XT barcoded sequencing adapters (Illumina, San Diego, CA, United States) and cleaned again following manufacturer instructions. Indexed PCR products were then quantified with QuBit (dsDNA High Sensitivity Assay Kit, Thermo Fisher Scientific, Waltham, MA, United States) and cleaned again using the Mini Elute PCR purification kit (Qiagen, Hilden, Germany), following manufacturer instructions. PCR products were then indexed with Nextera XT barcoded sequencing adapters (Illumina, San Diego, CA, United States) and cleaned again following manufacturer instructions. Indexed PCR products were then quantified with QuBit (dsDNA High Sensitivity Assay Kit, Thermo Fisher Scientific, Waltham, MA, United States) and pooled in equimolar ratios. Pooled samples were inspected for quality on the BioAnalyzer (Agilent Technologies, Santa Clara, CA, United States) and sequenced at 6 pM on the Illumina MiSeq, 2 bp × 300 bp paired-end v3 chemistry according to the manufacturer instructions.

Sequence Data Processing and Analysis

Raw reads were quality-filtered (QV = 25) and decontaminated of Illumina adapter and phiX sequences using the BBduk tool from the BBMap v37.62 suite (Bushnell B’). Mothur v1.44.3 (Schloss et al., 2009) was used for sequence data processing and analysis. Briefly, forward and reverse reads were assembled into contigs and ambiguous reads removed before quality trimming and pre-clustering took place (Huse et al., 2010). After exclusion of singletons, sequences were aligned to the Silva database 138.1 (Pruesse et al., 2007). Chimeric sequences were further excluded with UCHIME (Edgar et al., 2011), as were chloroplast, mitochondrial, archaeal, and eukaryotic sequences using the remove.lineages command in mothur. Sequences were then classified against the Silva database (McDonald et al., 2012). Sequences were reclassified against the Silva database after subsampling to 10,669 sequences, which was the highest number of sequences possible while still keeping all replicates. Operational Taxonomic Units (OTUs) were defined based on

http://sourceforge.net/projects/bbmap/
Amoebophilaceae (57.97 and 20.69%, respectively, Figure 1A). For further analyses, sequences were clustered into Operational Taxonomic Units (OTUs) at a 97% similarity level. A total of 3,266 OTUs were identified from this dataset. DNA samples contained 2,047 OTUs, while RNA samples contained 1,659 OTUs. DNA samples were dominated by *Thalassotalea* sp. (18.37% relative abundance, OTU0002, 100% identical to *Thalassotalea* sp. GenBank accession no. MN822801.1), *Alteromonas* sp. (16.54% relative abundance, OTU0003, 100% identical to *Alteromonas macleodii*, GenBank accession no. CP018321.1) and unclassified Flavobacteriaceae (12.77% relative abundance, OTU0005, 98.95% identical to *Polaribacter* sp., GenBank accession no. MK818916.1, Figure 1B). In contrast, *Fulvivirga* sp. (OTU0001, 95.85% identical to *Fulvivirga* sp., GenBank accession no. JQ516517.1) dominated RNA samples (57.00% relative abundance), followed by *Candidatus amoebophilus*, (OTU0004, 17.69%, Figure 1B).

In each pH treatment, some OTUs were present exclusively in RNA samples while others were exclusively found in DNA samples (Supplementary Figures 1B–E). Yet, most of the top
abundant OTUs were present in both DNA and RNA samples, (Supplementary Table 2) despite a noticeable disparity in abundance. For instance, the most abundant bacterial OTU found in DNA, Thalassotalea sp., (OTU0002), only accounted for 0.18% in RNA samples. Similarly, Fulvivirga sp. (OTU0001) accounted for more than half of the bacterial community in RNA samples, yet, they only represented 1.47% of the relative abundance in DNA samples. The next three most abundant taxa found in DNA samples accounted for only between 0.007 and 0.04% of the relative abundance in RNA samples. Similarly, following the same pattern, the next three most abundant taxa in RNA samples accounted for only between 0.15 and 0.17% of the relative abundance in DNA samples (Supplementary Figure 2A). In fact, Stylophora pistillata’s bacterial community was distinct when comparing 16S rDNA and 16S rRNA sequencing results from the same samples (PERMANOVA, pMC = 0.0001, Figure 2A). Furthermore, Simper analysis indicated a dissimilarity value between DNA and RNA samples of 87.14% (Supplementary Table 3). Alteromonas sp. (OTU0004), Thalassotalea sp. (OTU0002) and Tenacibaculum sp. (OTU0010) contributed the most to the DNA-RNA dissimilarity (Supplementary Table 4).

Decreased pH Has Larger Impact on Active Bacterial Communities

We identified approximately even numbers of OTUs in DNA samples with a total of 730 OTUs at pH 8.0, 717 at pH 7.8, 796 at 7.4, and 871 at 7.2 in the DNA samples and 131 OTUs were shared between all pH groups (Supplementary Figure 2B). For RNA samples, the number of OTUs was decreasing with pH, with 1,078 bacterial OTUs at pH 8.0, 759 at pH 7.8, 166 at pH 7.4 and 169 at pH 7.2, with only 33 shared OTUs between all pH groups (Supplementary Figure 2C). When considering only the DNA data, there was a significant general effect of pH (PERMANOVA, pMC = 0.02, Pseudo F = 2.0182), with significant differences between pH 7.2 and 8.0 (pMC = 0.0312, t = 1.9779, Figure 2B). We observed increased abundance of families Flavobacteriaceae at pH 7.8 and 8.0 (40 and 29%), and Chlorobiaceae and Alteromonadaceae at pH 7.2 (19 and 37% respectively, Figure 1A). At OTU level, unclassified Flavobacteriaceae (OTU0005) was relatively more abundant at pH 7.8 and 8.0 (28 and 21%, respectively), while Alteromonas sp. (OTU0004) and Prostheocochloris sp. (OTU0007) was more abundant at 7.2 pH treatments (25 and 18%, respectively) (Figure 1B). LEfSe analysis indicated 14 representative OTUs for pH 8.0, four for pH 7.8, seven for pH 7.4 and seven for pH 7.2 in DNA samples (Figure 3A).

When considering only the RNA data, there was also a significant effect of pH (PERMANOVA, pMC = 0.0044, Pseudo F = 2.8021, Figure 2C), yet with significant differences between pH 7.2 and 8.0 (pMC = 0.0332, t = 1.9342) and between pH 7.4 and 8.0 (pMC = 0.0228, t = 2.1838). Cellvibrionaceae was more abundant at pH 7.2, while Cyclobacteriaceae decreased in relative abundance at pH 8.0. Families Alcaligenaceae, Amoebophilaceae, Rhodobacteriaceae and Florabacteriaceae were more abundant at pH 8.0 (Figure 1A). At OTU level, Castellaniella sp. and “Candidatus Amoebophilus” were more abundant at pH 8.0. Fulvivirga sp. was less abundant at pH 8.0, while unclassified Cellvibrionaceae was more abundant at pH 7.2 (Figure 1B). LEfSe analysis identified five microbial OTUs enriched at pH 8.0, two at pH 7.8 and one enriched at pH 7.4 (Figure 3B).

Alpha diversity measures (Chao estimate of species richness, Shannon diversity and Simpson evenness indices) were calculated using mothur. While Shannon diversity index was significantly higher in the control treatment (pH 8.0) compared to 7.2 and 7.4 pH (ANOVA, p < 0.05; Tukey, p < 0.05) at the RNA level, the same was not observed at the DNA level (Table 1). Although the control treatment showed a pattern of higher Chao estimated richness at RNA level than at low pH, and higher in pH 7.2 at DNA level than at high pH, this was not statistically significant (Kruskal–Wallis, p > 0.05). Community evenness was similar amongst pH treatments at DNA level (ANOVA, p < 0.05), while a significant decreasing pattern was observed from low to high pH at RNA level with the least even communities at control pH (ANOVA, p < 0.05, Table 1).

Core Microbiome

Bacterial OTUs that were present in all treatments and all replicates (from DNA samples only) were considered to constitute the putative core microbiome. This resulted in 23 taxa that were consistently found to be part of the microbiome of Stylophora pistillata (Table 2). Amongst them, Thalassotalea sp. (18.37% of relative abundance), Alteromonas sp. (16.54%) and unclassified Flavobacteriaceae (12.77%) were the most abundant taxa. The remaining taxa showed substantially lower relative abundances, ranging from 4.18 to 0.04%.

DISCUSSION

Discrepancy Between Resident and Active Bacterial Communities in Response to Ocean Acidification

This is the first research to assess the effects of OA on a coral microbiome considering both the resident and active bacterial communities. Sequencing of 16S rDNA gene is a de facto standard method to identify microbial communities. However, a major disadvantage of this method is that DNA can remain in the environment after cell death, making it challenging to distinguish active from dormant populations or even extracellular material (Carini et al., 2016). Since RNA is short-lived, and ribosome quantity is generally associated with active growth and cellular activity, some authors consider sequencing of reverse transcribed rRNA a more accurate representation of metabolically active microbial communities (Ramos et al., 2000).

In this study, we found differences in bacterial community composition and relative abundance when sequencing the resident (DNA) and active (RNA) communities from the same samples. The top OTUs most contributing to these differences were around 4–8 times more abundant in the resident than the active community. Interestingly, a previous study on marine sediments detected 16S rDNA gene abundances of bacterial
communities 1–2 orders of magnitude higher than the transcript abundances (Li et al., 2020). Conversely, some OTUs were more abundant in the active community. This decoupling could be explained by either a response to specific treatment or by a substantial fraction of bacteria becoming active when their abundance decreased. This could suggest that high abundance limits activity or that top-down processes, such as grazing and virus lysis, could be involved in promoting or regulating activity (Berdjeb et al., 2011).

Interestingly, we found many OTUs exclusively in either the resident or the active communities, with only 13.5% overlap between both sources. However, the majority of the most abundant OTUs were present in both fractions. Previous studies found that only 29–50% of 16S rDNA sequences were identical to sequences in the 16S rRNA library from the same soil (Nogales et al., 2001; Mengoni et al., 2005; Lillis et al., 2009). Other studies also detected stark differences between 16S rDNA and rRNA libraries from soil samples (Baldrain et al., 2012; Romanowicz et al., 2016) and the coral L. pertusa (Galand et al., 2018). Undetected OTUs may reflect low abundance below PCR amplification levels (Mengoni et al., 2005) or they could be excluded during subsampling. In a biological context, the presence of a specific taxa in only rDNA samples could indicate lack of activity, while exclusively detecting taxa on the rRNA level might indicate rare species that are highly active. It could also indicate colonization by opportunistic bacteria that are likely not active and have low ecological relevance (Galand et al., 2018).

Our results showed higher OTU richness and diversity for resident compared to active communities, indicating that the metabolically active bacterial components in S. pistillata constitute only a fraction of the total microbiome. A similar pattern was observed by other authors for bacterial populations in the rhizosphere of Chrysanthemum (Duineveld et al., 2001), coral reef and marine sediments (Gaidos et al., 2011), as well as other studies based on sequencing of bacterial communities from different environments (Lanzén et al., 2011; Weigold et al., 2016). The higher richness and diversity of DNA based libraries is likely a reflection of recovering a broader range of microbes, including active, dormant and even dead cells as opposed to only active cells detected by RNA based libraries (Salgar-Chaparro and Machuca, 2019).

However, it is important to note that bias introduced by PCR-induced artifacts and different extraction methods for DNA and RNA, including the use of RNA protection agents, is a recurrent problem when characterizing microbial communities (Acinas et al., 2005; McCarthy et al., 2015).

Ocean Acidification Effects Are Distinct Between Resident and Active Microbial Communities

Many studies have evaluated the effects of ocean acidification on the coral microbiome, but most of them focused only on sequencing of 16S rDNA (O’Brien et al., 2016). While these studies found some effect of OA on community composition and function (Webster et al., 2013, 2016; Morrow et al., 2015; Grottoli et al., 2018), others reported a stable microbiome (Meron et al., 2012; Zhou et al., 2016; O’Brien et al., 2018). In this study, we detected a significant difference in bacterial communities from pH 7.2 and pH 8.0 in both active and resident bacterial communities, and from pH 7.4 and 8.0 in the active community. Overall, we found stronger differences in response to OA in the active compared to the resident bacterial communities. A similar pattern was observed by other authors in response to pollution and toxic compounds (Mengoni et al., 2005; Hoshino and Matsumoto, 2007; Lillis et al., 2009). This pattern is, to some extent, expected since the resident community also encompasses dead and non-active cells that will not respond to changes in the environment. Thus, our data support the notion that RNA better describes the actively responding community, which should show the most direct effects of disturbance (Mengoni et al., 2005).

We observed a drop in richness and diversity in the active community with decreasing pH, suggesting growth and activity limitation at low pH for many bacterial OTUs. For instance, Castellaniella sp. (OTU 0009, 100% identical to Castellaniella denitrificans, GenBank accession no. MK039104.1), which is from the Alcaligenaceae family and known to be involved in
Denitrification processes (Rosenberg, 2013) was present in the resident community at all pH levels, but decreased with pH in the active community. Similarly, studies on denitrifying prokaryotes in soil, wastewater sludges and batch reactor cultures found a decrease in denitrification activity in low pH (<6.5) (Thörn and Sörensson, 1996; Ellis et al., 1998). Denitrifying prokaryotes are found in coral holobionts worldwide (Gaidos et al., 2011; Yang et al., 2013; Rädecker et al., 2015; Tilstra et al., 2019). Denitrification by members of the coral microbiome may have a potential role in contributing to a balanced nitrogen to phosphorous ratio by decreasing nitrogen levels in coral tissues. Maintaining a N-limited state may generally contribute...
TABLE 1 | Differences in alpha diversity indices (Chao estimate of species richness, Shannon and Simpson evenness) in microbiome of Stylophora pistillata kept in aquaria for 2 years at different pH treatments (7.2; 7.4, 6.8 and 8.0) for DNA and RNA samples.

| Treatment | Chao1 Mean | SD | SE | Shannon Mean | SD | SE | Simpson evenness Mean | SD | SE |
|-----------|------------|----|----|-------------|----|----|-----------------------|----|----|
| DNA 7.2   | 765.94     | 100.73 | 58.16 | 2.83 | 0.35 | 0.20 | 0.01 | 0.00 | 0.00 |
| 7.4       | 494.74     | 323.31 | 186.66 | 3.30 | 0.25 | 0.15 | 0.04 | 0.02 | 0.01 |
| 6.8       | 492.58     | 260.52 | 150.41 | 2.54 | 0.66 | 0.38 | 0.02 | 0.02 | 0.00 |
| 8         | 410.36     | 142.16 | 82.08 | 3.38 | 0.43 | 0.25 | 0.03 | 0.00 | 0.00 |
| RNA 7.2   | 147.16     | 45.09 | 26.03 | 1.08 | 0.14 | 0.08 | 0.03 | 0.00 | 0.00 |
| 7.4       | 164.47     | 59.28 | 34.22 | 0.74 | 0.23 | 0.14 | 0.02 | 0.01 | 0.00 |
| 7.8       | 492.65     | 552.17 | 318.79 | 1.34 | 0.82 | 0.47 | 0.01 | 0.01 | 0.00 |
| 8         | 684.66     | 130.10 | 75.11 | 2.42 | 0.43 | 0.25 | 0.01 | 0.00 | 0.00 |

TABLE 2 | Core microbiome of Stylophora pistillata kept in aquaria for 2 years at different pH treatments (7.2; 7.4, 7.8 and 8.0).

| OTU  | Taxon       | Average Abundance  | Relative abundance% | Blast on NCBI database |
|------|-------------|---------------------|---------------------|------------------------|
| 7.2  | Thalassotalea 1 | 1,295.7 | 1,927.3 | 2,312.3 | 2,304.0 | 18.37 | 536 | 98% | Thalassotalea sp. |
| 7.4  | Alteromonas sp. 3 | 2,703.7 | 2,082.0 | 1,368.7 | 904.3 | 16.54 | 536 | 100% | Alteromonas marina |
| 7.8  | Flavobacteriacea 5 | 218.0 | 81.7 | 2,946.3 | 2,202.3 | 12.77 | 527 | 100% | Flexibacter sp. |
| 8    | Alteromonas sp. 8 | 558.7 | 481.0 | 420.3 | 325.7 | 4.18 | 536 | 98% | Alteromonas sp. |
| 7.2  | Tenacibaculum holothuriorum 10 | 354.7 | 626.0 | 271.0 | 271.7 | 3.57 | 525 | 98% | Tenacibaculum holothuriorum |
| 7.4  | Thalassolituus oleivorans 12 | 401.3 | 368.7 | 224.7 | 149.7 | 2.68 | 525 | 98% | Thalassolituus oleivorans |
| 7.8  | Kordia sp. 11 | 296.7 | 481.7 | 23.0 | 189.7 | 2.32 | 523 | 100% | Kordia sp. |
| 8    | Shimia sp. 13 | 375.0 | 182.3 | 15.3 | 188.0 | 1.49 | 531 | 98% | Shimia sp. |
| 7.2  | Thalassobius alba 14 | 40.3 | 390.3 | 15.3 | 188.0 | 1.49 | 531 | 98% | Shimia sp. |
| 7.4  | Alteromonas sp. 15 | 111.7 | 177.7 | 156.7 | 396.7 | 1.97 | 534 | 98% | Mameleilla alba |
| 7.8  | Alteromonas sp. 16 | 40.3 | 390.3 | 15.3 | 188.0 | 1.49 | 531 | 98% | Alteromonas sp. |
| 8    | Thalassolituus oleivorans 17 | 218.0 | 81.7 | 2,946.3 | 2,202.3 | 12.77 | 527 | 100% | Flexibacter sp. |
| 7.2  | Alteromonas sp. 18 | 558.7 | 481.0 | 420.3 | 325.7 | 4.18 | 536 | 98% | Alteromonas sp. |
| 7.4  | Tenacibaculum holothuriorum 19 | 354.7 | 626.0 | 271.0 | 271.7 | 3.57 | 525 | 98% | Tenacibaculum holothuriorum |
| 7.8  | Unclassified bacteria 20 | 14.0 | 247.7 | 6.0 | 65.7 | 0.78 | 357 | 98% | Leptospira ruysenii |
| 8    | Alteromonas sp. 21 | 16.7 | 254.3 | 8.0 | 4.3 | 0.66 | 536 | 98% | Alteromonas sp. |
| 7.2  | Flavobacteriaceae 22 | 15.7 | 3.7 | 202.0 | 55.3 | 0.65 | 536 | 98% | Alteromonas sp. |
| 7.4  | Endozoicomonas 23 | 7.7 | 119.7 | 3.7 | 61.3 | 0.45 | 510 | 94% | Alteromonas sp. |
| 7.8  | Vibrio alginolyticus 24 | 27.0 | 54.0 | 70.0 | 25.7 | 0.41 | 536 | 98% | Vibrio alginolyticus |
| 8    | Microscilla 25 | 49.0 | 10.0 | 24.0 | 15.7 | 0.23 | 534 | 98% | Microscilla |
| 7.2  | Ruegeria sp. 26 | 9.7 | 33.0 | 22.0 | 16.3 | 0.19 | 529 | 98% | Ruegeria sp. |
| 7.4  | Microscilla 27 | 49.0 | 10.0 | 24.0 | 15.7 | 0.23 | 534 | 98% | Ruegeria sp. |
| 7.8  | Endozoicomonas 28 | 11.7 | 177.7 | 156.7 | 396.7 | 1.97 | 534 | 98% | Alteromonas sp. |
| 8    | Microscilla 29 | 49.0 | 10.0 | 24.0 | 15.7 | 0.23 | 534 | 98% | Microscilla |

Only DNA samples were considered when defining the core microbiome.

to holobiont functioning and health through the regulation of Symbiodiniaceae proliferation (Rädecker et al., 2015; Cui et al., 2019; Tilstra et al., 2019). In fact, previous studies using the same coral colonies did not observe a decrease in symbiont density with pH decrease (Tambutté et al., 2015). Consequently, a decrease in denitrifying bacteria at lower pH treatments could potentially impact nitrogen balance in coral holobionts.

Our LEfSe analysis indicated that Alteromonas sp. (OTU0008) and Lacimicrobium sp. (OTU0029) were representatives of the active community under control conditions at pH 8.0, and nearly absent in lower pH treatments. Many species of the family Alteromonadaceae, especially Alteromonas, are sulfate reducers and known to be involved in DMSP metabolism (Raina et al., 2009), which is linked to osmoregulation and antioxidant capacity (Gardner et al., 2016; Hopkins et al., 2016). Previous studies also associated this genus with disease resistance (Wright et al., 2017). Thus, the decrease in abundance of members of this genus at low pH treatments could indicate loss in resilience and biogeochemical functions. Nonetheless, the functional role and ecological relevance of many bacterial taxa is still unknown, and this research area is essential for understanding holobiont functioning during stress. Even though OA can negatively
influence microbial communities, the implication of those shifts on the host health, ecosystem function and adaptation or acclimation mechanisms are still being investigated. In contrast, we found that OA stimulated growth and activity of specific OTUs, such as unclassified Cellvibrionaceae (OTU0006, 98.28% identical to Exilibacterium trileocarpae, GenBank accession no. MN094885.1). This OTU was present in low abundance in the resident community, suggesting that it might represent an opportunistic bacterium that increased in activity in response to acidification in low pH treatments.

Interestingly, richness and diversity in the resident community did not differ statistically between treatments, and remained high even in low pH treatments, suggesting that many bacterial taxa remain in the community after environmental change as dormant members. Dormancy plays an important role in shaping bacterial communities and is a well-known strategy used by many organisms to survive through unfavorable conditions (Sussman and Douthit, 1973). Previous studies indicate that dormant bacteria can account for high proportion of taxa in stressed ecosystems (Jones and Lennon, 2010). Some resident representative OTUs identified by LEfSe analysis at pH 7.2 were members of the phylum Bacteroidetes, including the genus Tenacibaculum (OTU0279), a known fish pathogen (Faulde et al., 2013) and members of the Flavobacteriaceae family (OTU0041 and OTU0375). A higher abundance of Bacteroidetes has been reported previously for corals under low pH conditions (Thurber et al., 2009; Meron et al., 2011; Morrow et al., 2015) and in diseased coral (Bourne et al., 2009). Similarly, three indicator species for pH 7.2 were from the order Alteromonadales, which encompasses some potential coral pathogens (Krediet et al., 2013; Roden et al., 2014).

Thus, a change in OA can induce shifts in the bacterial community by inhibiting activity of many taxa while stimulating activity of a few specific, potentially opportunistic, taxa previously resident in low abundances. Despite the identified changes on microbial community composition in response to OA, the implications for host health are still unknown. In fact, changes in the coral microbiome due to environmental stress do not necessarily reflect holobiont stress, but could even reflect potential acclimation responses. Thus, more studies should be conducted to investigate consistent patterns in microbiome changes associated with environmental stressors, as well as acclimation potential and ecological functions of key microbial taxa in coral phenotypes.

Core Microbiome of Stylophora pistillata
To study patterns in microbiome composition, previous studies have focused on the core microbiome, which is defined as the members consistently shared among microbial consortia from similar habitats (Shade and Handelsman, 2012). Usually, the coral microbiome has a core species-specific bacterial group and a transient group that changes according to environmental parameters such as climate and geography (Ritchie, 2006).

S. pistillata has previously been shown to have specific associations with species from the genus Endozoicomonas (Bayer et al., 2013; Neave et al., 2017) and we also identify them as part of the core microbiome in this study (Table 2). However, their abundance is lower than reported in studies using colonies sampled from the wild, potentially due to the long-term rearing of the colonies used here in an aquarium environment. Coral microbiome studies greatly depend on corals maintained in aquaria that simulate natural conditions, but the effects of captivity on the coral microbiome are understudied. Previous research has observed differences in the microbiome of corals between colonies in situ and those reared in aquaria. For instance, shifts on the surface mucus layer (SML) microbiome were reported for Fungia granulosa (Kooperman et al., 2007) and Siderastrea siderea after as early as 1 day of aquarium rearing (Fratté et al., 2015). In the coral Lophelia pertusa, both skeleton and SML were different between in situ and ex situ habitats (Schöttner et al., 2009). In Eguchipsammia fistula, the associated microbiome also differed between native and aquarium settings. Functional profiling suggested that processes related to carbon- and nitrogen-limited environments were enriched in the native habitat, which could reflect diet changes between in situ and aquarium conditions (Röthig et al., 2017). Galand et al. (2018) investigated microbiome changes in aquaria across different time points, both on the active (rRNA) and resident (rDNA) fraction. Their results suggest a species-specific response. For Madrepora oculata, bacterial associations remained stable for at least 6 months of captivity, which is in agreement with a stable bacterial community in aquarium-reared Acropora loripes over 4 weeks (Hill and Ralph, 2006; Danjanovic et al., 2020). Interestingly, Galand et al. (2018) further found that the abundance of Endozoicomonas in the RNA fraction decreased after only 1 day of captivity but started to increase again after 5 days. For Lophelia pertusa, they observed microbiome changes after only 1 day of aquarium rearing. Some taxa disappeared completely and other appeared only after months in aquaria, which the authors considered an aquarium signature. They further concluded that opportunistic bacteria might colonize corals in captivity. Thus, considering the long-term nature of this study’s experimental setup, with corals kept in captivity for several decades, it is possible that their microbiomes diverged significantly from the ones found in natural environments. This illustrates a potential limitation of using aquarium specimens for microbial community studies, since it will limit the presence of potentially ecological relevant bacterial taxa associated to geographical location.

Importance of the Active Microbiome
In this study, we identified bacterial taxa with low relative abundance on DNA level, but high abundance on RNA level, indicating rare but highly active community members. The existence of a large diverse microbial community, with few abundant and many rare taxa is common in the natural environment (Sogin et al., 2006). While the relationship between bacterial abundance and activity is still unclear, one common hypothesis for this distribution is that abundant bacteria have high growth rates, while rare taxa are slow growing (Pedrós-Alió, 2006). However, multiple studies have found rare bacteria taxa to be disproportionally active and displaying high growth rates (Jones and Lennon, 2010; Campbell et al., 2011; Weigold et al., 2016).
There is an increasing number of studies highlighting the importance of rare bacterial taxa (Sogin et al., 2006; Pedrós-Alió, 2007, 2012; D Ainsworth et al., 2015; Lynch and Neufeld, 2015; Jousset et al., 2017). These often underrated and unexplored species may represent a source of genetic novelty in times of environmental change. Some might even be keystone species, being essential to understand community composition and function (Sogin et al., 2006; Elshahed et al., 2008).

CONCLUSION

The use of 16S rDNA sequencing to study bacterial communities is widespread. However, only few studies use 16S rRNA sequencing, which means much is unknown about the correlation between microbial activity and composition, particularly for coral holobionts. We show that OA can significantly impact the resident coral microbiome, although a stronger effect was observed on the active microbial community, which suggests that the microbial community responded primarily to the environmental change through modified activity patterns. Importantly, we also found that low pH might play a role in the shift of bacterial OTUs from active to dormant states and vice versa, allowing opportunistic bacteria to increase in activity while reducing the abundance of some members of the core microbiome. The microbiome is a critical component of coral resilience, and this study provides insights into the effects of ocean acidification on coral bacterial community composition and coral holobiont response to climate change. Moreover, our study confirms that rDNA and rRNA based sequencing can result in different community composition, with the RNA-based community reflecting the metabolically active population. Our analysis also showed that some low abundant resident bacterial taxa were highly active, suggesting they could play key functional roles. Thus, both methods, rDNA and rRNA sequencing, should be considered complementary tools to investigate microbial assemblage structure and activity.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ebi.ac.uk/ena, PRJEB44699.

AUTHOR CONTRIBUTIONS

DZ, ST, DA, ET, and AV performed the experiment in Monaco. MZ, CV, and MA designed the study. MB performed laboratory work, analyzed the data, and wrote the first draft of the manuscript. CA contributed to the bioinformatics analysis. All authors contributed to the writing and editing of the manuscript and read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021.707674/full#supplementary-material

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