1	Article type:	Original	Article
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- 4 high pCO_2 reef.
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14 Keywords: Microbiome, limestone-boring, ocean acidification, *Ostreobium*, coral skeletons, multi-

15 marker metabarcoding.

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21 Running title: Endolithic microbes and ocean acidification

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This is the author manuscript accepted for publication and has undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the <u>Version of Record</u>. Please cite this article as <u>doi: 10.1111/mec.14268</u>

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32 Abstract:

34 The health and functioning of reef-building corals is dependent on a balanced association with 35 prokaryotic and eukaryotic microbes. The coral skeleton harbours numerous endolithic microbes, but 36 their diversity, ecological roles and responses to environmental stress, including ocean acidification, 37 are not well characterized. This study tests whether pH affects the diversity and structure of prokaryotic 38 and eukaryotic algal communities associated with skeletons of Porites spp. using targeted amplicon 39 (16S rRNA gene, UPA and tufA) sequencing. We found that the composition of endolithic 40 communities in the massive coral *Porites* spp. inhabiting a naturally high pCO_2 reef (avg. pCO_2) 41 811 μ atm) is not significantly different from corals inhabiting reference sites (avg. pCO₂ 357 μ atm), 42 suggesting that these microbiomes are less disturbed by ocean acidification than previously thought. 43 Possible explanations may be that the endolithic microhabitat is highly homeostatic, or that the 44 endolithic microorganisms are well adapted to a wide pH range. Some of the microbial taxa identified 45 include nitrogen-fixing bacteria (Rhizobiales and cyanobacteria), algicidal bacteria in the phylum 46 Bacteroidetes, symbiotic bacteria in the family Endozoicomoniaceae, and endolithic green algae, 47 considered the major microbial agent of reef bioerosion. Additionally, we test whether host species has 48 an effect on the endolithic community structure. We show that the endolithic community of massive 49 Porites spp. is substantially different and more diverse than that found in skeletons of the branching 50 species Seriatopora hystrix and Pocillopora damicornis. This study reveals highly diverse and 51 structured microbial communities in Porites spp. skeletons that are possibly resilient to ocean 52 acidification.

53 Introduction

54 Ocean acidification (OA) is predicted to threaten the persistence of coral reefs by affecting the 55 balance between constructive forces (calcification and growth of reef builders) and destructive forces 56 (bioerosion and carbonate dissolution) (Tribollet 2008a; Andersson & Gledhill 2013). Acidification 57 lowers the saturation state of calcium carbonate (CaCO₃), making it more difficult for calcifying 58 organisms, such as stony corals, to build their skeletons (Orr et al. 2005; Hofmann et al. 2010). OA has 59 been shown to slow down calcification and cause structural deformities in juvenile corals (Crook et al. 60 2013; Foster et al. 2016). Some studies, however, indicate that corals are able to regulate the pH at the 61 tissue-skeleton interface, where calcification takes place, mitigating the potential consequences of OA 62 on the calcification process (McCulloch et al. 2012; Venn et al. 2013; Georgiou et al. 2015). Rates of 63 biological dissolution of CaCO₃ (bioerosion) tend to increase under low pH conditions, mostly due to 64 an increase in biomass of the boring organisms living inside coral skeletons (Manzello et al. 2008; 65 Tribollet et al. 2009; Crook et al. 2013; Fang et al. 2013; Reyes-Nivia et al. 2013; Enochs et al. 2016), potentially resulting in a shift from a net reef accretion condition to one of net erosion (Andersson &Gledhill 2013).

68 The skeletons of live and dead corals harbour bacteria, fungi, sponges and an abundant 69 population of limestone-boring algae, all having important roles in the reef's CaCO₃ budget (Le 70 Campion-Alsumard et al. 1995; Tribollet 2008a; Verbruggen & Tribollet 2011). For example, the 71 green alga Ostreobium can be responsible for 70-90% of carbonate dissolution within dead corals, 72 eroding as much as 1 kg of reef carbonate per m^2 per year (Tribollet 2008b; Grange *et al.* 2015). Green 73 algal biomass in live coral skeletons exceeds Symbiodinium biomass in coral tissues by about 16 times 74 (Odum & Odum 1955), making the limestone attractive to grazers and further increasing bioerosion 75 (Chazottes et al. 1995; Clements et al. 2016). However, endolithic algae also protect corals from high 76 light stress (Yamazaki et al. 2008) and provide vital nutrients to corals, potentially extending the time 77 they can survive without Symbiodinium during bleaching events (Schlichter et al. 1995; Fine & Loya 78 2002). Endolithic algae have exceptionally high levels of cryptic diversity (Marcelino & Verbruggen 79 2016; Sauvage et al. 2016; Del Campo et al. 2017), and although it is known that their biomass 80 increases substantially upon acidification and warming (Tribollet et al. 2009; Reves-Nivia et al. 2013), 81 it is not known which of the cryptic species increase in relative abundance.

82 The endolithic community, along with the coral host and its other symbionts, constitutes the 83 coral holobiont (Rohwer et al. 2002). The responses of the coral microbiome, including both 84 prokaryotic and eukaryotic members, to acidification has gained attention as we continue to uncover 85 vital roles played by microorganisms in holobiont health and resilience (Bourne et al. 2009; Sharp & 86 Ritchie 2012; Krediet et al. 2013; Blackall et al. 2015; Bourne et al. 2016). Because the ocean pH 87 naturally changes throughout seasons, along depth gradients, with productivity and other biological 88 factors, marine microorganisms may have the physiological plasticity required to cope with the 89 predicted levels of ocean acidification over the next 100 years (Joint et al. 2011). This notion is 90 supported by several studies showing stable coral prokaryotic community when shifted from ambient 91 to high CO₂ partial pressure (pCO_2) and therefore reduced seawater pH conditions (Meron *et al.* 2012; 92 Webster et al. 2016; Zhou et al. 2016). However other studies have demonstrated that a reduced 93 seawater pH can lead to the loss of Symbiodinium (coral bleaching) and trigger shifts from a healthy 94 microbiome composition to a microbial community typically associated with diseased corals (Anthony 95 et al. 2008; Vega Thurber et al. 2009; Meron et al. 2011; Webster et al. 2013; Morrow et al. 2015). 96 These different responses of a coral's microbiome to reduced seawater pH may reflect differences in 97 resilience across coral species to acidification or different experimental setups used in the various 98 studies.

99 Reefs at the Milne Bay Provence of Papua New Guinea (PNG) are in close proximity to 100 volcanic seeps (expelling ~ 99% pure CO_2) and constitute a good model system to study the impacts of 101 acidification *in situ* on the microbial community associated with corals. Both coral species composition 102 and the prokaryotic microbial community associated with coral tissue and mucus differ between high 103 pCO_2 seep sites and nearby reference sites with ambient pCO_2 (Fabricius *et al.* 2011; Morrow *et al.* 104 2015). However, little is known about the coral endolithic communities and how these may change 105 under various seawater pH conditions. Previous studies have screened 16S rRNA gene clone libraries 106 and demonstrated contrasting results, with significant effects of OA community composition within the 107 skeleton in an experimental system (Meron *et al.* 2011) but no significant changes in corals 108 transplanted to a natural CO_2 seep site (Meron *et al.* 2012). One limitation with the 16S rRNA gene 109 marker is that it underestimates the diversity of eukaryotic algae (Marcelino & Verbruggen 2016), and 110 as a consequence, the major microbial agents of bioerosion have been overlooked.

111 Here we use high-throughput amplicon sequencing to investigate the effects of ocean 112 acidification on the diversity and structure of endolithic microbial communities of corals inhabiting a 113 high pCQ_2 site in PNG. Our goals are to: 1) test whether the community composition of prokaryotes 114 and photosynthetic eukaryotes (assessed with the 16S rRNA gene, UPA and tufA markers) within the 115 skeletons of massive colonies of Porites spp. differs between high pCO2 sites and nearby reference 116 sites where pCO_2 is not affected by the volcanic seeps; 2) compare the endolithic communities of 117 Porites spp. with those of two branching coral species (Seriatopora hystrix and Pocillopora 118 damicornis) to investigate whether the microbiome in coral skeletons varies among host species; and 3) 119 describe the endolithic community diversity found in corals of Papua New Guinea and discuss the 120 potential functional roles of this microbiome under ocean acidification.

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- 123 Methods
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- 125 Field sites and sampling

126 Samples of massive *Porites* spp. (n = 24, six per site and month) were collected in April and127 November 2014 at two high pCO_2 (seep) and reference sites within the D'Entrecasteaux Islands, Milne 128 Bay Province, Papua New Guinea. High pCO_2 samples were collected at Illi Illi (Upa-U) Seep 129 (09.82425S, 150.81789E) and Dobu Seep (09.73646S, 150.86894E), and at nearby reference sites 130 (ambient pCO_2) not exposed to elevated pCO_2 conditions (Illi Illi Reference, 09.82806S, 150.82028E 131 and Dobu Reference, 09.75211S, 150.85410E) (Fabricius et al. 2011; Uthicke et al. 2013). High pCO₂ 132 and reference sites were ~500 m and ~3 km apart from one another at Illi Illi and Dobu respectively. 133 Samples of the branching corals Seriatopora hystrix (n = 3 at each site) and Pocillopora damicornis (n 134 = 3 at each site) were only collected in April 2014 at Illi Illi seep and reference sites (same as above). 135 Seawater carbonate chemistry varies in response to bubble activity and water motion at the seep sites; thus, at Illi Illi seep, corals experience a pH range (defined here as the 5th and 95th percentiles) of 7.28– 136 137 8.01 (avg. pCO₂ 624 μatm) and at Dobu seep a pH range of 7.08-7.99 (avg. pCO₂ 998 μatm). At Illi 138 Illi reference site the pH ranges from 7.91-8.09 (avg. pCO₂ 346 µatm) and at Dobu reference site the 139 pH ranges from 7.91-8.10 (avg. pCO₂ 368 µatm) (Fabricius et al. 2014), which is within the range of 140 future predictions for the year 2100 (Moss et al. 2010). 141 Coral fragments were collected using bone cutters or a hammer and chisel and placed into

142 individual sections within a plastic tackle box, which allowed for water flow whilst underwater. After

returning to the boat, samples were immediately placed into flowing seawater sourced directly from the collection site. Large pieces of *Porites* spp. were chipped into smaller fragments, rinsed thoroughly with sterile 0.02 µm-filtered seawater and placed in 50 mL Falcon tubes with RNAlater (Ambion). Samples were kept in a cooler with ice until returned to the laboratories at the Australian Institute of Marine Science (AIMS) where they were processed.

Fragments were removed from RNAlater and soaked in 0.2 μ m filtered calcium and magnesium free seawater for ~10 minutes at room temperature (CMFSW; 0.45M NaCl, 10mM KCl, 7mM Na₂SO₄, 0.5mM NaHCO₃ and milli-Q water; (Esteves *et al.* 2016)) to aid in the initial removal of tissues from the skeleton. Tissues were removed into the CMFSW using an air gun fitted with a sterile tip. Skeletons with tissues removed were placed back into the original RNAlater collection buffer and stored at -80°C until shipment to the University of Melbourne where DNA was isolated from the endolithic community (see below).

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156 DNA isolation, library preparation and sequencing

157 Total DNA was isolated from coral skeletons using the Wizard Genomic DNA Purification 158 Kit (Promega). The manufacturer's protocol for plant DNA was followed, with the exception of an 159 extended 3 hr incubation step with the first extraction buffer to allow the DNA to leak out from the 160 limestone into the solution. Amplified DNA products for library preparation were obtained with a two-161 step process described by Marcelino and Verbruggen (2016). During the first PCR step, three 162 metabarcoding markers were amplified: the 16S rRNA gene (Klindworth et al. 2013); the universal 163 plastid amplicon (UPA), which is a fragment of the 23S rRNA gene (Presting 2006; Sherwood & 164 Presting 2007) and the elongation factor Tu (tufA), which targets green algae (Ulvophyceae) (Fama et 165 al. 2002; Marcelino & Verbruggen 2016). During the second PCR step, barcodes and Illumina adapters 166 were attached to both 3' and 5' ends of the amplicons. One negative control was performed with each 167 amplification (6 in total, one per marker and per amplification step) and sequenced with the library, 168 even though no DNA was detected in any negative control during quantification. Two mock 'blank' 169 extractions were also performed along with the samples DNA isolation, processed through the 170 amplification process and sequenced to further control for possible contamination. DNA isolation and 171 PCR preparation were carried out inside a dedicated dead-air box (PCR workstation) sterilized with 172 UV light for 15 min prior to each use. Libraries were quantified using the Quant-It PicoGreen reagent 173 (Invitrogen) and pooled with other samples of another project. The library was sequenced using the Illumina MiSeq platform (2×300 bp paired end reads) at the Centre for Translational Pathology, 174 175 University of Melbourne. Further details about the primers and library preparation are provided in 176 Supplementary materials.

177

178 Data processing

179 The MiSeq run yielded one file containing all amplicons per sample, which were 180 demultiplexed based on the primer sequences. The 3' ends of reads were trimmed to improve

181 consensus quality; forward and reverse reads were merged using FLASH (Magoc & Salzberg 2011) 182 and sequences having average quality scores smaller than 35 or lengths shorter than a threshold (350 bp 183 for 16S rRNA gene, 320 bp for UPA and 400 bp for the tufA) were filtered out using PRINSEQ 184 (Schmieder & Edwards 2011). To verify that the data lost during quality control did not affect the 185 results, we also performed the analyses with a less stringent quality filtering (Supplementary 186 materials). Sequences were clustered into Operational Taxonomic Units (OTUs) using UPARSE 187 (Edgar 2013). A similarity threshold of 98% was set for the *tufA* marker, a threshold near species level 188 for this marker (Sauvage et al. 2016). For the other markers, the default threshold of 97% was used. 189 The 16S rRNA gene OTUs were aligned with PvNAST (Caporaso et al. 2010a) while the UPA and 190 tufA datasets were aligned with MAFFT (Katoh et al. 2002). A taxonomy affinity was assigned to the 191 OTUs using the Naïve Bayesian Classifier (RDP) implemented in OIIME v.1.9.1 (Wang et al. 2007; 192 Caporaso et al. 2010b). The Greengenes v.13.8 dataset (DeSantis et al. 2006) was used to classify the 193 16S rRNA gene sequences, and custom-made reference datasets (described and available in Marcelino 194 & Verbruggen 2017) were used for tufA and UPA. The resulting OTU table went through a filtering 195 process to remove OTUs found in the negative controls (3 OTUs in the 16S rRNA gene, 5 OTUs in the 196 UPA and 2 OTUs in the tufA datasets) and rare OTUs (i.e. OTUs with less than 5 reads across all 197 samples and OTUs from samples where they are present with 2 or less reads). OTUs were also filtered 198 based on their taxonomic classification to focus on the taxonomic groups that each marker best 199 characterizes: chloroplast sequences were excluded from the 16S rRNA gene dataset and bacterial 200 sequences were excluded from the tufA dataset. Further details about the data processing pipeline are 201 provided in the Supplementary materials.

202

203 Statistical analysis

204 There were no significant differences related to the time of collection (Supplementary 205 materials), therefore all *Porites* spp. samples (n = 24) were used to investigate the effects of pCO₂ in 206 endolithic communities associated with this coral genus. Rarefaction curves of the number of observed 207 OTUs per number of reads were constructed by randomly subsampling the reads in OIIME, allowing to 208 set a threshold for each marker where the curve reaches saturation (i.e. a plateau in the rarefaction 209 curve), which was 2,200 reads in the 16S rRNA gene, 1,400 in the tufA and 7,000 in the UPA dataset 210 (Supplementary figure 1). Samples with sequencing depths lower than these thresholds were excluded, 211 resulting in 20 samples in the 16S rRNA gene, and 22 samples in the UPA and tufA datasets 212 (Supplementary Table S1). Alpha diversity indices (Chao1 and observed OTUs) were calculated using 213 QIIME (Caporaso et al. 2010b). The relative abundance of individual OTUs and taxonomic groups (i.e. 214 OTUs grouped at phylum level for bacteria and genus level for algae) between sites were tested for 215 significant differences with a Kruskal-Wallis test (for OTUs) and ANOVA (for taxon groups) using 216 QIIME (Caporaso et al. 2010b).

To further investigate the distribution of green algal lineages, a maximum likelihood phylogeny was built with the green algal *tuf*A OTUs together with reference sequences (from GenBank) using a GTR+gamma model of sequence evolution in RAxML v.8.2.6 (Stamatakis 2006).

OTUs present in less than 3 samples were excluded, their relative abundances were normalized with
 cumulative sum scaling (Paulson *et al.* 2013) and visualized alongside a phylogenetic tree using the R
 package phytools (Revell 2012).

223 Potential differences in community composition between high pCO_2 and reference sites (beta-224 diversity) were investigated with a combination of statistical methods. Principal coordinate analysis 225 (PCoA) on weighted UniFrac distance matrices was performed using QIIME (Caporaso et al. 2010b; 226 Lozupone et al. 2011) and the results visualized using the ggplot2 package in R (Wickham 2009). A 227 multivariate generalized linear model (MGLM) was used to investigate potential differences in 228 community composition between high pCO_2 and reference sites. The MGLM was computed using the 229 mvabund R package (Wang et al. 2012), considering a negative binomial distribution. The null 230 hypothesis of no difference among sites was statistically tested with analysis of deviance using 999 231 bootstrap interactions (R scripts provided in Supplementary materials). To verify that the results are not 232 a consequence of PCR bias, PCoA and MGLM analyses were also performed with a distance matrix 233 based on Sørensen similarity, which is a presence/absence index (Supplementary materials).

234 The number of samples of the branching species (S. hystrix and P. damicornis) did not allow 235 statistical analyses to test for differences between high pCO_2 and reference sites, although it did permit 236 a comparison of the endolithic communities associated with the different coral hosts (Supplementary 237 Table 1). To investigate the community structure related to host species, a rarefaction threshold of 707 238 reads for the 16S rRNA gene, 713 for the tufA and 3257 for the UPA marker was used, allowing the 239 inclusion of a larger number of samples in the analysis. Samples with sequencing depths lower than 240 these thresholds were excluded, resulting in 35 samples in the 16S rRNA gene and UPA datasets and 241 27 in the tufA datasets (Supplementary Table S1). Alpha diversity, Kruskal–Wallis test (for OTUs), 242 ANOVA (for taxon groups), PCoA and MGLM were performed on this dataset as previously 243 described, but here, samples from different pCO₂ conditions from conspecific host species were 244 combined in order to investigate the community structure purely associated with coral host species.

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- 247 Results
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249 Effects of pCO₂ conditions on the Porites spp. endolithic microbiome

250 A total of 6,584,274 sequence reads were recovered for the samples analysed here, 4,405,336 251 belonging to Porites spp. samples (ca. 25% of these reads belonged to a marker that is not analysed in 252 this manuscript). After stringent pre and post OTU-clustering and filtering, a total of 119,367 (16S 253 rRNA gene), 109,948 (tufA) and 393,816 (UPA) reads remained in the analysed Porites spp. dataset. A 254 less stringent quality filtering which resulted in inclusion of more sequence reads did not alter the 255 community patterns (see Supplementary materials). The alpha diversity statistics, including Chao1 and 256 observed OTUs, indicated that the species richness of the endolithic communities associated with 257 *Porites* spp. was not significantly different between high pCO_2 and reference sites (Table 1). Although

- the relative abundance of some microbial taxa differed between high pCO_2 and reference sites (Figure 1, see also Supplementary figure 2 for a sample-based representation), the differences were not statistically significant (Bonferroni-corrected *p*-values = 1), neither at the OTU level (Supplementary table S2) nor at higher taxonomic levels (Supplementary table S3). Accordingly, principal coordinate analysis and MGLM did not reveal any significant pattern between sites with all three markers (Figure 2).
- We further investigated whether any of the different phylogenetic lineages in the endolithic algal communities differed in abundance at high pCO_2 and reference sites. A phylogenetic heatmap of relative abundances (Supplementary figure 3) indicated that phylogenetic relatedness among green algae is not correlated with different abundances in high pCO_2 or reference sites. Notably, three algal OTUs were present in either high pCO_2 or reference sites, but not in both (Supplementary figure 3).
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270 Taxonomic profiling of the Porites spp. endolithic community

271 The microbial community observed in the skeletons of *Porites* spp. was highly diverse and 272 variable among samples within pCO_2 conditions. Prokaryotic members of the microbiome (observed in 273 the 16S rRNA gene dataset) accounted for most of the species diversity (Table 1), and no bacterial 274 OTUs were present in all Porites spp. samples. The most abundant phylum recovered was 275 Proteobacteria, followed by Bacteroidetes and Archaea (Figure 1A, Supplementary figure 2). The 276 relative abundance of the nitrogen-fixing order *Rhizobiales* (Alphaproteobacteria) in all *Porites* spp. 277 samples was $9.1\% \pm 4\%$ standard deviation (hereafter \pm only), and the phylum of green sulphur 278 bacteria, Chlorobi, was $0.4\% \pm 1\%$. Members of the Bacteroidetes were twice as abundant at high 279 pCO_2 sites (12.1% ± 10% in reference versus 23.9% ± 17% in high pCO_2 sites), mostly due to a higher 280 abundance within the classes Cytophagia ($4.8\% \pm 7\%$ versus $10.4\% \pm 9\%$), Flavobacteria ($2.2\% \pm 2\%$) 281 versus $3.7\% \pm 3\%$) and Saprospirae (2.4% ± 5% versus 9.4% ± 15%). We also observed a lower 282 abundance of the Archaeal class *Parvarchaea* in the high pCO_2 site (1.6% \pm 0.9% versus 11.6% \pm 12% 283 in high pCO_2 and reference sites, respectively). A representation of the relative abundances of bacterial 284 phyla on a sample-by-sample basis can be found in Supplementary figure 2. These differences were, 285 however, not statistically significant based on ANOVA and Kruskal-Wallis tests (Bonferroni-corrected 286 p-values = 1, Supplementary tables S2 and S3).

287 The *tufA* dataset (Figure 1B) suggested that the algal community was dominated (64.7% \pm 288 36%) by lineages of the Ostreobiaceae (Chlorophyta, Bryopsidales). Although no tufA OTUs were 289 found to be omnipresent in *Porites* spp., siphonous green algae (order Bryopsidales) were present in all 290 samples analysed here, indicating that they are ubiquitous members of the coral skeleton core 291 microbiome. Ostreobium clade #1 showed the highest relative abundance $(33\% \pm 40\%)$, followed by 292 Ostreobium clades #4, #3 and #2. While the relative abundance of clade #1 was similar between sites, 293 the relative abundance of the other Ostreobium clades varied substantially, but not statistically 294 significant, between reference and high pCO_2 sites. A high abundance of an unclassified group of 295 OTUs belonging to the green algal order Bryopsidales was also observed, particularly in the reference 296 site (Figure 1B).

297 The prevalence of green algal lineages in the skeletons of *Porites* spp. was also suggested by 298 the UPA dataset (Figure 1C), which shows that $86.8\% \pm 15\%$ of the reads belong to the green algal 299 order Bryopsidales. There are no UPA reference sequences for Ostreobium clades #1 and #2, therefore 300 possible sequences of these clades might have been classified as clades #3, #4 or "unclassified 301 Bryopsidales" by the RDP classifier, which may explain the differences in the abundances of 302 Ostreobium clades between the tufA and UPA datasets. One UPA OTU was found across all Porites 303 spp. samples (OTU_3). This OTU was only classified at the kingdom level as a eukaryote 304 (Supplementary table S2) and showed little similarity to known red algae and Stramenopiles species 305 (blastn e-values \leq 4e-79 but Identity \leq 79%). Differences in the relative abundances between reference 306 and high pCO_2 sites were minimal (Figure 1C, Supplementary figure 2).

307

308 Endolithic communities across different coral host species

309 The prokaryotic endolithic communities of Seriatopora hystrix and Pocillopora damicornis 310 were significantly less rich than those found in *Porites* spp., as indicated by Chao1 and observed OTUs 311 indices (*p*-value = 0.003, Table 2). A significant difference was detected in the relative abundances of 312 certain OTUs belonging to the Endozoicimonaceae family between coral species, with the highest 313 abundance in *P. damicornis* (Kruskal-Wallis, *p*-values < 0.0002, Supplementary table S4). The *Porites* 314 spp. samples had a higher relative abundance of an OTU related to the order Rhizobiales (genus 315 Afifella), and P. damicornis showed a significantly (p-value = 0.03) higher abundance of an OTU 316 related to the phylum Bacteroidetes (order Cytophagales; Supplementary table S4). At higher 317 taxonomic levels, the relative abundance of the phyla Planctomycetes, Bacteroidetes, and the bacterial 318 phylum OD1 were significantly different among coral hosts (ANOVA, p-values = 0.007 and 0.01 319 respectively, Figure 3A, Supplementary figure 2, Supplementary table S5). Principal coordinate 320 analysis (PCoA) showed evident differences in the prokaryotic microbiome associated with different 321 coral hosts: Porites spp. samples clustered together, clearly separated from the two branching species 322 (Figure 4A). The MGLM analysis confirmed a significant difference between the prokaryotic 323 communities of different coral hosts (Figure 4A).

324 The alpha diversity of green algae (i.e. Chao 1 and Observed OTUs within the *tufA* dataset) 325 was significantly different between S. hystrix and Porites spp., but no significant differences were 326 observed within the taxonomically broader spectrum of eukaryotic algae amplified with the UPA 327 marker (Table 2). The Kruskal-Wallis test suggested no significant difference in the relative 328 abundances of particular OTUs between host species (neither within the *tufA* nor within the UPA 329 dataset), at least when corrected (Bonferroni) p-values were taken into consideration (Supplementary 330 table S4). After rarefying the sequences, the *tufA* dataset was reduced to a single *P. damicornis* sample 331 (Supplementary table S1), therefore we could not test for differences in alpha diversity or relative 332 abundances in the P. damicornis community compared to other host corals. We observed a 333 significantly different relative abundance of Ostreobium spp. (order Bryopsidales) among coral hosts 334 within the UPA dataset (ANOVA, p-value = 0.000002), but not in the *tufA* dataset (p-values > 0.05, 335 Figure 3B and 3C, Supplementary figure 2, Supplementary table S5). Seriatopora hystrix had a high

336 and variable $(83\% \pm 57\%)$ relative abundance of endolithic lineages related to the macroalga Halimeda 337 spp. within the *tufA* dataset, while this group constituted a minimal fraction $(0.04\% \pm 0.2\%)$ of the 338 endolithic community of Porites spp. (Figure 3B). No pattern was observed within the PCoA plot of 339 the community composition using the *tufA* marker (Figure 4B). The UPA marker, which comprised 340 more samples of the branching species, showed that Porites spp. samples cluster together and away 341 from S. hystrix and P. damicornis, although two outliers belonging to branching samples cluster with 342 Porites spp. (Figure 4C). The MGLM analysis suggested no significant differences between the algal 343 communities of different coral hosts (Figure 4B and C), except when a presence-absence distance 344 matrix was used (Supplementary materials, Supplementary figure 4).

345

346 Discussion

347 Our results show that the prokaryotic and eukaryotic microbiome in the skeletons of Porites 348 spp. are highly diverse but indistinguishable between corals inhabiting naturally high pCO_2 reefs and 349 ambient conditions. Ocean acidification is predicted to affect the coral reef CaCO₃ budget and its 350 biological associations (Meron et al. 2011; Andersson & Gledhill 2013; Morrow et al. 2015), and 351 depending on the experiment, endolithic communities were shown to either exacerbate or buffer the 352 effects of these environmental changes (Fine & Loya 2002; Tribollet et al. 2009; Reyes-Nivia et al. 353 2013). Our results suggest that the composition of endolithic communities, at least in *Porites* spp., is 354 virtually unaffected by the surrounding high pCO_2 water from a natural volcanic seep, and therefore 355 less likely to be disturbed by OA than we previously thought. Although homogeneous between high 356 pCO_2 and reference sites, we show that the endolithic community is highly diverse and structured 357 among coral host species.

358

359 A stable microbiome

The mechanisms influencing the structure of the endolithic microbiome (regardless of variable pCO_2 conditions), are currently unknown. We raise here two hypotheses that may explain our results. The "stable habitat" hypothesis assumes that the endolithic environment is highly homeostatic so that pH is maintained inside the skeletons regardless of external changes in the surrounding water. The "tolerant endolith" hypothesis is based on the notion that endolithic microorganisms have a wide pH tolerance range, potentially wider than the microbes associated with the tissues and mucus.

The first hypothesis is supported by the ability of some corals to up-regulate the pH at the tissue-skeleton interface, which allows them to calcify and grow even under high pCO_2 (McCulloch *et al.* 2012; Venn *et al.* 2013; Georgiou *et al.* 2015). The pH within coral cells remains relatively constant throughout the day (7.05-7.46 units), likely due to membrane transporters that extrude the excess of byproducts of photosynthesis and respiration to maintain a stable intracellular pH (Laurent *et al.* 2013). This process may indirectly create a stable microhabitat within the coral skeleton that is protected from shifting pH in the surrounding seawater (see also Shashar *et al.* 1997). The observation that

373 radioactivity in seawater impacted corals' living tissue but did not reach their endolithic zone (Odum &374 Odum 1955) supports this notion.

375 One problem with the stable habitat hypothesis is that the pH in the skeletons of *Porites* 376 (compressa) can vary daily from 7.7 to 8.5 pH units, mostly due to the by-products of respiration and 377 photosynthesis of the coral and Symbiodinium that are exported to the skeleton (Shashar & Stambler 378 1992). This daily variation is well above the projections of OA for the near future, which predicts a pH 379 drop of 0.4 units by 2100, and up to 0.7 units by 2300 (Raven et al. 2005; Hoegh-Guldberg et al. 380 2007). A reasonable counterargument is that the direction of the movement (pH reduction due to OA) 381 may be more important than the daily variation within skeletons – for example, microorganisms living 382 under 7.7 to 8.5 pH units not necessarily withstand a seawater pH shift from 7.8 to 7.6 pH units. The 383 resilience of at least some microbes might also be related to their boring mechanism, which involves a 384 sophisticated control of intracellular pH (associated with calcium pumps and protons counter-transport) 385 in endolithic cyanobacteria (Garcia-Pichel 2006; Garcia-Pichel et al. 2010). Therefore our second 386 hypothesis that organisms exposed to daily pH fluctuations within the skeleton are adapted to cope 387 with a wide range of pCO_2 conditions may be more accurate. Experimental work and genomic data of 388 endolithic organisms will help to test the tolerant endolith and the stable habitat hypotheses. For 389 example, specialization to the low light experienced in the endolithic habitat has been observed in the 390 plastid genome of Ostreobium quekettii (Marcelino et al. 2016) and the presumed pH tolerance may 391 also be reflected in the genomes of endolithic organisms.

392 The lack of discernible differences in the endolithic community composition between high 393 pCO_2 and reference corals observed here is in agreement with the results of an experiment where 394 Balanophyllia europaea and Cladocora caespitose corals were transplanted to a naturally high pCO_2 395 area (Meron et al. 2012). In an aquarium-based experiment conducted over a shorter time, the bacterial 396 community composition present in the tissue, skeleton and mucus of Acropora eurystoma were found 397 to be affected by high pCO_2 , but further analysis using clone libraries suggested that only the 398 prokaryotic communities of the mucus and tissue, not the skeleton, were affected by low pH (Meron et 399 al. 2011). These different observations might be associated with the different time spans and 400 experimental setups of the two studies, and it is likely that the microbial community associated with 401 different coral taxa have different responses to acidification. The resilience of endolithic algae to 402 acidification has also been observed: the net photosynthesis and respiration of algae growing at the 403 surface of dead coral blocks was severely impacted upon exposure to high pCO_2 treatments, while the 404 endolithic flora was unaffected (Tribollet et al. 2006). Studies have demonstrated that endolithic algae 405 actually benefit from low pH and tend to increase in biomass under high pCO_2 conditions (Tribollet et 406 al. 2009; Reyes-Nivia et al. 2013; Enochs et al. 2016).

407 The observation that high pCO_2 did not impact the endolithic community of *Porites* spp. does 408 not necessarily imply that coral holobionts are immune to ocean acidification. It is possible that the 409 methods used here are not sufficiently powerful to detect the effects of high pCO_2 on endolithic 410 microbial communities. However, the fact that we detected significant differences among coral hosts 411 even though the sampling size for the branching corals was smaller, indicates that our methods and

412 sampling design are adequate and it is unlikely that differences among high pCO_2 and reference sites 413 were present but went undetected. It is possible though that high pCO_2 impacts the endolithic 414 communities of other coral species that were not examined. It is interesting to note that massive Porites 415 spp. dominate the reef near volcanic seeps while the presence of branching species (e.g. Acropora spp.) 416 was largely reduced (Fabricius et al. 2011). Our analyses are restricted to the volcanic seeps of Milne 417 Bay, which have relatively small areas under high pCO_2 and are surrounded by ambient seawater. 418 Further studies at additional sites impacted by high pCO_2 and across a wider range of coral species is 419 required to evaluate the results in our study across the broader ecological context of effects of OA on 420 coral microbiomes.

421

422 Diversity and potential functional roles of the endolithic microbiome

423 Bacteria related to *Endozoicomonas* spp. (class Gammaproteobacteria) are predicted to have a 424 key role in the coral holobiont. These bacteria have been shown to be endosymbionts, forming 425 aggregations within coral tissues (Neave et al. 2017), potentially contributing to nutrient cycling and 426 structuring of the microbiome through the production of quorum-sensing signalling metabolites and 427 antimicrobial compounds (Meyer et al. 2014; Morrow et al. 2015 and references therein). The relative 428 abundance of *Endozoicomoniaceae* within coral tissues appears to be sensitive to high pCO_2 (Morrow 429 et al. 2015; Webster et al. 2016), but in the skeletons of Porites spp. analysed here, they did not differ 430 significantly between samples from different pCO_2 conditions (Supplementary Table S2). We observed 431 a significantly higher relative abundance of two Endozoicomoniaceae OTUs in the skeletons of P. 432 damicornis when compared to the other two coral species, possibly reflecting stable associations of 433 Endozoicomoniaceae species with this coral host (see Neave et al. 2017). Although some of the 434 sequences retrieved here may derive from other parts of the coral, other studies have also detected 435 members of Endozoicomonaceae in the endolithic community (Williams et al. 2015; Marcelino & 436 Verbruggen 2016; see also Ainsworth et al. 2015).

437 Bacteria in the phylum *Bacteroidetes* are often associated with coral disease and have been 438 shown to increase in relative abundance under reduced pH (Vega Thurber et al. 2009). The average 439 relative abundance of this group doubled in endolithic communities from high pCO_2 sites, but this 440 difference was not significant likely due to the high level of variation in community composition 441 among colonies within sites. This increase was mostly due to a higher abundance of the classes 442 Saprospirae, Flayobacteria and Cytophagia, which contain most of the known marine algicide bacteria 443 (Furusawa et al. 2003; Mayali & Azam 2004; Zozaya-Valdes et al. 2015). It is plausible that a higher 444 relative abundance of these microorganisms is associated with an increase in endolithic algal biomass 445 under high pCO₂ (see Reyes-Nivia et al. 2013, Johnson et al. 2017). Rather than compromising coral 446 health, these bacteria may control excessive endolithic algal growth and may help to maintain a stable 447 community composition under ocean acidification.

448 Microorganisms involved in nitrogen cycling may be fundamental to coral resilience to ocean 449 acidification and climate change (Rädecker *et al.* 2014; Santos *et al.* 2014; Radecker *et al.* 2015). We 450 found a diverse community of nitrogen fixing (diazotrophic) microorganisms inhabiting coral

451 skeletons. The majority (in terms of relative abundances) belonged to the order *Rhizobiales*, a group 452 that appears to form stable symbiotic associations with corals (Lema et al. 2014). Green sulphur (also 453 diazotrophic) bacteria in the phylum Chlorobi, previously documented as prevalent members of the 454 endolithic community in the coral Isopora (Yang et al. 2016), were found at low relative abundances in 455 the samples analysed here and in a previous study (Marcelino & Verbruggen 2016). Cyanobacterial 456 OTUs captured with the UPA marker, while not abundant, were very diverse and mostly unclassified at 457 lower taxonomic levels. Cyanobacteria have been shown to fix nitrogen in coral tissues (Lesser et al. 458 2004; Radecker et al. 2015) and can be responsible for a large fraction of the nitrogen fixation 459 observed in their skeletons (Crossland & Barnes 1976; Davey et al. 2007).

460 Endolithic algal biomass has been shown to increase under high pCO_2 , as phototrophic 461 organisms benefit from the increased availability of carbon dioxide for photosynthesis (Tribollet et al. 462 2009; Reyes Nivia et al. 2013). Indeed, we observed a higher relative abundance of all Ostreobium 463 clades in *Porites* spp. samples from high pCO_2 sites, but the variability among replicates (i.e. *Porites* 464 spp. samples within sites) was also high, making it difficult to draw conclusions about whether 465 Ostreobium spp. are competitively superior to other endolithic algal lineages under OA. Whether the 466 increase in algal biomass is a threat to corals under OA depends on whether the associated bioerosion 467 levels will exceed reef accretion (calcification). Besides increasing bioerosion, excessive endolithic 468 algal growth can penetrate the coral living tissue, possibly increasing their susceptibility to infections 469 (Peters 1984; Fine et al. 2006). An increase in endolithic algae may also be beneficial to the coral by 470 providing them with vital nutrients, which is especially important during coral bleaching events 471 (Schlichter et al. 1995; Fine & Loya 2002).

472 The possibility that the endolithic microbiome contributes to the resilience of corals under 473 future OA conditions deserves further attention. Massive Porites spp. may be considered more 474 competitive under OA than branching species based on their prevalence at naturally high pCO₂ sites 475 (Fabricius et al. 2011). We visibly observed that our massive Porites spp. samples had higher 476 colonisation with endolithic algae compared to the branching species. The photosynthetic activity of 477 Symbiodinium plays an important role in maintaining pH homeostasis within corals (Gibbin et al. 478 2014), and it is possible that endolithic algae provide a similar service within the skeleton. The biomass 479 of endolithic algae may exceed that of Symbiodinium by 16-fold (Odum & Odum 1955) and can 480 contribute significantly to the buffering capacity of the holobiont (see Yamazaki et al. 2008; Reyes-481 Nivia et al. 2014). It is noteworthy that several functionally important microorganisms (e.g. 482 Endozoicomoniaceae and Bacteroidetes) often found in coral tissues and mucus also occur in coral 483 skeletons (Sweet et al. 2010; Ainsworth et al. 2015; Willians et al. 2015; Marcelino & Verbruggen 484 2016; this study). It is possible that some of these microorganisms were initially associated with polyp 485 tissues, but remained after removing the tissues from the skeleton with pressurized air, especially in 486 Porites spp. which is a perforate coral with tissues that penetrate the skeleton, or that they penetrated 487 the coral skeleton when the samples were placed in the storage buffer. It is noteworthy however that 488 multiple studies have found tissue-associated microbes within coral skeletons, and that the majority of 489 OTUs commonly found in healthy corals have also been found in bare coral skeleton, but not in 490 seawater or in a diseased coral tissue (Fernando et al. 2015). It is possible therefore that the coral

491 skeleton serves as a reservoir for the microbiome and provides a source of beneficial bacteria to coral 492 tissues, analogous to the human appendix which functions as a safe house for symbiotic microbes that 493 repopulate the intestine following acute illness (Randal Bollinger *et al.* 2007). Acute environmental 494 stress can disrupt symbiotic relationships among hosts and symbionts (see Hawkins *et al.* 2013), and a 495 stable endolithic community may assist in the recovery of the coral microbiome after environmental 496 (and/or physiological) conditions stabilize.

497

498 Different host species harbour distinct endolithic communities

499 The endolithic communities of the branching corals Seriatopora hystrix, Pocillopora 500 damicornis and the massive Porites spp. contain significantly different relative abundances of 501 functionally important members of the microbiome (including species of Endozoicimonaceae and 502 Bacteroidetes) and appear to separate based on morphology or taxonomy (as both branching species 503 belong to the family *Pocilloporidae*). First, the two branching species harbour a reduced diversity of 504 bacteria and algae. The low relative abundance of Ostreobium spp. in the endolithic communities of 505 branching species is surprising, considering the generally ubiquitous nature of this alga in coral 506 skeletons (Odum & Odum 1955; Tribollet 2008a; Gutner-Hoch & Fine 2011). Instead of Ostreobium 507 spp., the coral S. hystrix has a high relative abundance of OTUs related to a macroalga (Halimeda 508 spp.), which has only recently been reported in coral skeletons. It is possible that Halimeda spp. occur 509 in the coral skeleton in the form of rhizoids that have penetrated the limestone, or most likely, as an 510 unknown microscopic and endolithic life stage of two Halimeda species (H. discoidea and H. 511 micronesica) that are commonly present in metabarcoding studies of endolithic communities 512 (Marcelino & Verbruggen 2016; Sauvage et al. 2016; this study).

513 The observed differences in endolithic community composition among coral hosts may be a 514 result of specialization to particular host traits or reflect co-evolution between coral hosts and 515 endolithic species. The living tissue of Porites (lobata) is about five times thicker, penetrates the 516 skeleton and contains a higher density of Symbiodinium than the living tissue of P. damicornis and S. 517 hystrix (Yost et al. 2013). Tissue thickness would influence the amount of light that penetrates and 518 reflects within the inner parts of the skeleton and may influence the composition of the endolithic 519 community. Branching coral species also tend to grow faster than massive corals (Gates & Ainsworth 520 2011), and the branch tips collected in this study may have a younger population of endoliths in 521 comparison to more mature sections of the colony base (a pattern reported in Pica et al. 2016). Future 522 studies would benefit from examining the microbiome associated with different areas of the colony and 523 possible specialization to skeletal features. Alternatively (and not mutually exclusively), endolithic 524 lineages might form stable community assemblies that have co-evolved with the coral host, or the host 525 species have some control over the composition of the endolithic community by selecting beneficial 526 taxa. Mutualistic relationships between corals and their endolithic associates have been suggested in 527 several studies (Odum & Odum 1955; Schlichter et al. 1995; Schlichter et al. 1997; Fine & Loya 2002; 528 Försterra & Häussermann 2008; Titlyanov et al. 2009), and future research would benefit from 529 characterizing possible co-evolutionary processes among coral species and endolithic microorganisms.

531 Conclusions

532 This study reports a diverse microbiome within the skeletons of Porites spp., and 533 demonstrates that little discernible patterns exist in this microbiome across ambient and naturally high 534 pCO_2 environments. We show that the endolithic community shares several functionally important 535 microbes with the coral tissue layer. Environmental stress can induce corals to lose their symbiotic 536 microorganisms, and a diverse endolithic microbial community might serve as a reservoir to recolonise 537 the microbiome in the coral tissue after the re-establishment of their physiological equilibrium. We 538 found functionally important members in the endolithic community, including members in the 539 Endozoicimonaceae and Bacteroidetes, forming distinct associations with the different host coral 540 families, an observation consistent with the endolithic reservoir proposition. The diversity and 541 community structure observed in this study form the baseline for future studies aiming to investigate 542 the roles of endolithic microorganisms in enabling corals to endure climate change.

543

544 Acknowledgements

545 This work was supported by the Australian Biological Resources Study (RFL213-08), the Australian 546 Research Council (FT110100585, DP150100705), the Holsworth Wildlife Research Endowment, the 547 Albert Shimmins Fund and the University of Melbourne (ECR grant to HV and scholarship to VRM), 548 Melbourne Bioinformatics (project UOM0007) and the Nectar Research Cloud (National Collaborative 549 Research Infrastructure Strategy). We are thankful to Emmanuelle Botté for coral collections in Papua 550 New Guinea and Sara Bell for laboratory assistance in processing samples. The expedition to conduct 551 the fieldwork and collections was funded by the Australian Institute of Marine Science. We thank the 552 reviewers (including from Axios) for their constructive feedback.

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813 Data Accessibility

- B14 DNA sequences have been deposited in NCBI's Sequence Read Archive (SRA) under the accession
 B15 IDs SAMN07251731 SAMN07251770.
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817 Author Contributions

KMM and DGB conceived and conducted the sampling. VRM performed DNA extractions, library
preparation, analyses and drafted the manuscript. MvO, KMM, DGB and HV reviewed the analyses
and contributed to writing.

- 821 822
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- 824 Tables:
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831

826 **Table 1:** Diversity indices based on the microbiome of *Porites* spp. skeletons from reference and high

827 pCO_2 sites and standard deviations (±). Chao1 and Number of Observed OTUs between high pCO_2

828 and reference sites were compared with a two-sample t-test, *p*-values ≤ 0.05 suggest significant

829 differences between sites, but were not observed. N = number of samples after rarefaction. Seqs =

830 rarefaction threshold. OTUs = number of OTUs retrieved in each dataset after quality filtering.

			-	Chao1		Obs. OTUS			
	Ν	Seqs	OTUs	Reference	high pCO ₂	<i>p</i> -value	Reference	high pCO ₂	<i>p</i> -value
16S	20	2200	890	141.2 ± 52.4	148.5 ± 45.8	0.79	133.6 ± 49.9	140.7 ±44.8	0.77
tufA	22	1400	59	7.8 ± 3.3	7.4 ± 2.8	0.78	7.7 ± 3.2	7.3 ± 2.8	0.74
UPA	22	7000	164	21.6 ± 7.1	24.4 ± 5.4	0.33	20.6 ± 6.7	23.2 ± 5.1	0.33
8.	32								
8	33								
8.	34								
8.	35								
8.	36								

837

16S rRNA gene	e Chao1						
Group1	Group2	Group1 mean	Group1 std	Group2 mean	Group2 std	t stat	<i>p</i> -valu
S. hystrix	P. damicornis	29.087	29.494	21.514	15.705	0.507	1.000
S. hystrix	Porites spp.	29.087	29.494	125.296	49.889	-4.363	0.003
P. damicornis	Porites spp.	21.514	15.705	125.296	49.889	-4.854	0.003
16S rRNA gene	e observed OTU	S					
Group1	Group2	Group1 mean	Group1 std	Group2 mean	Group2 std	t stat	<i>p</i> -valu
S. hystrix	P. damicornis	21.767	19.758	19.083	12.331	0.258	1.000
S. hystrix	Porites spp.	21.767	19.758	101.274	37.282	-4.865	0.003
P. damicornis	Porites spp.	19.083	12.331	101.274	37.282	-5.138	0.003
tufA Chao1							
Group1	Group2	Group1 mean	Group1 std	Group2 mean	Group2 std	t stat	<i>p</i> -valu
S. hystrix	P. damicornis	2.033	0.858	3.450	0.000	1.168	1.000
S. hystrix	Porites spp.	2.033	0.858	9.606	3.849	-3.264	0.006
P. damicornis ¹	Porites spp.	3.450	0.000	9.606	3.849	-1.531	0.309
tufA observed	OTUs						
Group1	Group2	Group1 mean	Group1 std	Group2 mean	Group2 std	t stat	<i>p-</i> valu
S. hystrix	P. damicornis	1.933	0.736	3.000	0.000	1.024	1.000
S. hystrix	Porites spp.	1.933	0.736	9.330	3.623	-3.388	0.003
P. damicornis ¹	Porites spp.	3.000	0.000	9.330	3.623	-1.673	0.192
UPA Chao 1							
Group1	Group2	Group1 mean	Group1 std	Group2 mean	Group2 std	t stat	p-valu
S. hystrix	P. damicornis	19.353	14.158	13.386	16.038	0.624	1.00
S. hystrix	Porites spp.	19.353	14.158	21.732	6.187	-0.591	1.00
P. damicornis	Porites spp.	13.386	16.038	21.732	6.187	-1.921	0.16
UPA observed	OTUs						
Group1	Group2	Group1 mean	Group1 std	Group2 mean	Group2 std	t stat	<i>p</i> -valu
S. hystrix	P. damicornis	18.100	12.882	12.567	14.816	0.630	1.000
S. hystrix	Porites spp.	18.100	12.882	19.539	5.572	-0.394	1.000
P. damicornis	Porites spp.	12.567	14.816	19.539	5.572	-1.754	0.327

hystrix and Pocillopora damicornis corals. Chao1 and Number of observed OTUs were compared with a two-sample t-test, p-values ≤ 0.05 indicate significant differences between host species and are

- 1 - The tufA dataset has only one P. damicornis sample (after rarefaction), therefore the significance cannot be calculated. See
- Supplementary table S1 for number of samples.



Reference High pCO2 This article is protected by copyright charves of the microbiome based on the tura marker; C) biodiversity survey using the Universal Plastid Amplicon.



Figure 2: Principal Coordinate Analysis of microbial communities present in limestone skeletons of *Porites* spp. from high *p*CO₂ and reference sites. The analyses were based on weighted UniFrac distance matrices calculated with OTU-level abundances for each metabarcoding marker: A) prokaryotic 16S rRNA gene marker; B) eukaryotic green algae *tuf*A marker; C) Universal Plastid Amplicon marker. The results of the MGLM analysis (Deviance and *P* value) are shown.



Figure 3: Relative abundances of the most common microbial taxa in coral skeletons of *Pocillopora damicornis*, *Seriatopora hystrix* and *Porites* spp. Results were based on all samples from each coral species, averaging the relative abundances at the taxonomic level displayed in the legend.

A) Biodiversity survey targeting prokaryotes based on the 16S rRNA gene; This article is protected **Byscopsyrighte Allkaglotic grsenvalg**al members of the microbiome based on the *tuf*A marker; C) biodiversity survey using the Universal Plastid Amplicon



Figure 4. Principal Coordinate Analysis of microbial communities present in limestone skeletons of three coral host species collected in high pCO_2 and reference sites. The analyses were based on weighted UniFrac distance matrices calculated with OTU-level abundances for each metabarcoding marker: A) prokaryotic 16S rRNA gene marker; B) eukaryotic green algae *tuf*A marker; C) Universal Plastid Amplicon marker. The results of the MGLM analysis (Deviance and *P* value) are shown.