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Different nitrogen sources speed recovery from corallivory and uniquely alter the microbiome of a reef-building coral

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ABSTRACT

Corals are in decline worldwide due to local anthropogenic stressors, such as nutrient loading, and global stressors, such as ocean warming. Anthropogenic nutrient loading, which is often rich in nitrate, inhibits coral growth and worsens corals' response to warming while natural sources of nitrogen, such as ammonium from fish excretion, promotes coral growth. Although the effects of nutrient loading and ocean warming have been well-studied, it remains unclear how these factors may interact with biotic processes, such as corallivory, to alter coral health and the coral microbiome. This study examined how nitrate vs. ammonium enrichment altered the effects of increased seawater temperature and simulated parrotfish corallivory on the health of Pocillopora meandrina and its microbial community. We tested the effects of nitrogen source on the response to corallivory under contrasting temperatures (control: 26 °C, warming: 29 °C) in a factorial mesocosm experiment in Moorea, French Polynesia. Corals were able to maintain growth rates despite simultaneous stressors. Seawater warming suppressed wound healing rates by nearly 66%. However, both ammonium and nitrate enrichment counteracted the effect of higher temperatures on would healing rates. Elevated seawater temperature and ammonium enrichment independently increased Symbiodiniaceae densities relative to controls, yet there was no effect of nitrate enrichment on algal symbiont densities. Microbiome variability increased with the addition of nitrate or ammonium. Moreover, microbial indicator analysis showed that Desulfovibrionaceae Operational taxonomic units (OTUs) are indicators of exclusively temperature stress while Rhodobacteraceae and Saprospiraceae OTUs were indicators of high temperature, wounding, and nitrogen enrichment. Overall, our results suggest that nitrogen source may not alter the response of the coral host to simultaneous stressors, but that the associated microbial community may be distinct depending on the source of enrichment.

Subjects Ecology, Marine Biology, Microbiology

Keywords Corallivory, Seawater warming, Nitrate, Ammonium, Coral microbiome, Nutrient loading

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Additional Information and Declarations can be found on page 18

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INTRODUCTION

Corals typically thrive in ecosystems with oligotrophic waters due to tight nutrient cycling between corals and algal symbionts (*Muscatine & Porter, 1977*). However, anthropogenic nutrient pollution has increased nutrient availability on many reefs worldwide, especially with respect to nitrogen (N) availability (*D'Angelo & Wiedenmann, 2014*; *Fabricius, 2011*). Anthropogenic-derived sources of N, often in the form of nitrate, have deleterious consequences on coral growth and physiology (*D'Angelo & Wiedenmann, 2014*; *Shantz & Burkepile, 2014*) and can make corals more susceptible to seawater warming (*Burkepile et al., 2019*; *Fabricius et al., 2013*; *Vega Thurber et al., 2014*). These patterns are concerning given that seawater warming events have been increasing in frequency and intensity due to global climate change (*Hughes et al., 2018*).

Alternatively, natural sources of N (e.g., ammonium from fish excretion) can benefit corals by increasing coral growth and calcification (*Holbrook et al., 2008; Meyer & Schultz, 1985; Meyer, Schultz & Helfman, 1984; Shantz & Burkepile, 2014*). In fact, corals under ammonium enrichment receive more translocated carbon from Symbiodiniaceae (formerly *Symbiodinium; LaJeunesse et al., 2018*) than corals enriched with nitrate (*Ezzat et al., 2015*). Ammonium can even alleviate the negative impacts of seawater warming by maintaining the response mechanisms of coral immunity at the molecular level (*Zhou et al., 2017*). Indeed, anthropogenic- vs. naturally occurring N seem to have contrasting impacts on coral physiology and susceptibility to seawater warming (*Burkepile et al., 2019*). Yet these effects on physiology can vary with N concentration (*Ferrier-Pages et al., 2000; Marubini & Davies, 1996; Marubini & Thake, 1999*), which likely influences how corals respond to other stressors (*Fabricius et al., 2013*).

Coral predation (i.e., corallivory) is a common biotic process on reefs with many corallivores removing coral mucus, tissue, and skeletal structure. Scraping and excavating corallivory by parrotfishes and pufferfishes removes coral tissue and varying degrees of skeletal structure, which can substantially reduce coral growth rates (Cole, Pratchett & Jones, 2008; Rice, Ezzat & Burkepile, 2019; Rotjan & Lewis, 2008). This impact can exacerbate corals' response to concurrent stressors and even prevent recovery from anthropogenic perturbations (Rice, Ezzat & Burkepile, 2019). For instance, parrotfish corallivory inhibited the recovery of Orbicella spp. colonies after a bleaching event (Rotjan et al., 2006). Nutrient loading can also interact with fish corallivory to drive changes in coral mortality. In the Florida Keys, parrotfish corallivory increased Porites mortality by 62% when corals were simultaneously exposed to anthropogenic nutrient sources (Zaneveld et al., 2016). These patterns may be driven by increases in opportunistic bacteria and wounding driving changes in the coral microbiome directly (Zaneveld et al., 2016) or resulting from compromised host immunity and thus inability of the coral to regulate its microbiome (Zaneveld, McMinds & Vega Thurber, 2017).

The coral microbiome is dominated by bacteria, protozoans, and archaea that perform a multitude of functions from nutrient cycling to protecting the host against opportunistic

bacteria (Bang et al., 2018; Bourne, Morrow & Webster, 2016; Ritchie, 2006; Rohwer et al., 2002). Some commensal microbes produce antibacterial compounds to prevent opportunistic bacteria from colonizing the host (Bourne, Morrow & Webster, 2016; Ritchie, 2006). When the microbial community is disturbed, populations of opportunistic bacteria can become established, which can compromise the holobiont immunity and lead to coral mortality (Glasl, Herndl & Frade, 2016). This mechanism has been proposed for Porites corals under simultaneous thermal stress and nutrient enrichment, and even parrotfish corallivory (Zaneveld et al., 2016). Yet, commensal microbes can help corals resist and recover from abiotic stress and are critical for enabling their host to cope with challenging environmental conditions (Bang et al., 2018; Bourne, Morrow & Webster, 2016). In a previous analysis of the coral microbiomes included in this study, we demonstrated that stressors primarily act additively or antagonistically, not synergistically, to alter microbial community composition with high temperature and simulated corallivory wounding independently causing the strongest responses (Maher et al., 2019). Additionally, we showed that changes in community structure with stress are driven by increases in opportunistic taxa, rather than the depletion of symbionts. When considering holobiont health, it is vital to understand how the interactions between corals and their microbial counterparts respond to anthropogenic forcing (*Rädecker et al., 2015*; for review see McDevitt-Irwin et al., 2017).

It remains unclear how N source (nitrate vs. ammonium) may alter corals' response to elevated temperatures and corallivory. To that end, our study seeks to evaluate how different N sources (nitrate vs. ammonium) may mediate changes in: (1) coral growth rates, (2) wound healing rates, (3) Symbiodiniaceae densities, and (4) bacterial community dynamics in response to concurrent seawater warming and corallivory. We hypothesized that nitrogen source would differentially mediate the effects of seawater warming and simulated corallivory. Further, we predicted that varying nitrogen source would produce distinct microbial communities with indicator taxa that suggest potential functional responses to multiple stressor regimes.

MATERIALS AND METHODS

Study species

We used a full factorial mesocosm experiment to test how nitrogen source (nitrate vs. ammonium) may alter a coral's response to seawater warming and corallivory. The experiment was conducted at the Richard B. Gump South Pacific Research Station in Moorea, French Polynesia (17°29′26.04″S, 149°49′35.10″W). Research was completed under permits issued by the French Polynesian Government and the Haut-commissariat de la République en Polynésie Francaise (Protocole d'Accueil 2005–2018). *Pocillopora meandrina* was chosen as the study species because it is one of the most abundant corals on the fore reef in Moorea (*Edmunds, 2018*) and is heavily preyed on by parrotfishes in the Pacific (*Cole, Pratchett & Jones, 2008*). We distinguished this taxon according to its distinct morphology, although we acknowledge that definitive taxonomy of *Pocillopora* spp. is challenging in this region (*Edmunds et al., 2016*).

Experimental design

In September of 2016, the experiment was conducted in twelve independent 150 L flow-through, temperature controlled mesocosms on a 12:12 light:dark cycle (Aqua Illumination Hydra 52 LEDS) at ~700 mmol m² s⁻¹. Seawater was pumped from Cook's Bay and filtered with a 20 μ m sediment filter before entering the mesocosms. A total of 10 healthy *P. meandrina* colonies with no observed corallivory were collected at three to four m depth on the north shore fore reef and transported in seawater by boat to the Gump Research Station. A total of 12 nubbins (2.7 ± 0.05 cm height) were fragmented from each colony and epoxied (using Z-spar A-788) onto one cm² plastic mesh. Nubbins were allowed to recover for ~24 h in the mesocosms at ambient temperatures (26 ± 1 °C).

After the acclimation period, half of the coral nubbins were mechanically injured on the branch tip using eight mm snub nose pliers to mimic parrotfish bites. The pliers were sterilized with ethanol and heat after each nubbin to prevent the transfer of microbes across replicates. The injuries were $45.2 \pm 1.5 \text{ mm}^2$ and $\sim 2 \text{ mm}$ deep, resulting in a single wound that removed the tissue layer and some skeletal structure. The injuries resembled a wound similar to a scraping parrotfish bite. Coral nubbins were randomly assigned to the following treatment tanks (n = 2 tanks per treatment combination): (1) $26 \degree C$, (2) $29 \degree C$, (3) ammonium and $26 \degree C$, (4) nitrate and $26 \degree C$, (5) ammonium and $29 \degree C$, and (6) nitrate and $29 \degree C$. Five intact and five wounded coral nubbins were in each tank (n = 10 nubbins per treatment; Fig. S1), and there were no differences in the initial weight ($29.1 \pm 0.7 \text{ g}$) of the coral nubbins across treatments (Kruskal–Wallis; $\chi^2 = 8.2$, P = 0.7).

After ~24 h of acclimation, half of the mesocosms were gradually raised from 26 to 29 °C, over a 24-h period (~1 °C change per 8 h), to reach temperatures observed during summer seawater warming in Moorea (*Pratchett et al., 2013*). To establish the pulse nitrogen treatments, the mesocosms assigned to nitrogen treatments were enriched every ~12 h to 4 μ M NO₃⁻ or 4 μ M NH₄⁺ from stock solutions of KNO₃ and NH₄Cl, respectively. Background nutrient concentrations in the seawater system during this time period were 0.34 μ M NO₃⁻, 0.21 μ M NH₄⁺, and 0.15 μ M soluble reactive phosphorus. The flow to tanks was ceased for 1 h during the enrichment. Nubbins were haphazardly moved within the tank every 2 days to avoid position effects. The experiment was maintained for 21 days.

Coral growth rates, Symbiodiniaceae densities, and wound healing rates

At the beginning and end of the experiment, coral nubbins were buoyant weighed to determine changes in mass for growth rates (*Davies, 1989; Jokiel, Maragos & Frankzisket, 1978*). For measurements of wound healing rate, initial and final photos were taken using an Olympus TG-4 camera and ruler and processed in ImageJ for scar area. A wound was considered healed if there were visible polyps in the wound area. After the experiment, coral nubbins were frozen at -40 °C for microbiome analysis. From each of the 12 treatments, six nubbins were randomly selected for microbial analysis while controlling for parent colony and tank effects. The tip of each nubbin was clipped off using sterilized

bone cutters, and frozen at -80 °C until DNA extractions. The remaining coral tissue was removed using 0.7 µm filtered seawater (FSW) and an air brush and collected into Falcon tubes. The tubes were centrifuged at 3,000 rpm for 10 min. The supernatant was removed and the Symbiodiniaceae pellet was resuspended with 10 mL of 0.7 µm FSW. Symbiodiniaceae densities were quantified using compound microscopy and a hemocytometer (n = 4 counts per replicate). The coral skeletons were dried at 60 °C for 7 days, allowed to cool to room temperature, and wax-dipped at 60 °C to determine surface area by regressing the difference in weight between single and double wax dippings against the surface area of known objects (*Stimson & Kinzie, 1991*). Growth rates and Symbiodiniaceae densities were normalized by the surface area of each nubbin.

16S library preparation, sequencing, and initial data processing

DNA was extracted from 72 samples (n = 6 per treatment) representing a subset of the experiment using the MoBio Powersoil[®] DNA Isolation Kit. Amplicon libraries were prepared for the V4 region of the 16S rRNA gene using the primer pair 515F (5'-GTG CCA GCM GCC GCG GTA A-3') and 806Rb (5'-GGA CTA CHV GGG TWT CTA AT-3') that targets bacterial and archaeal communities (*Apprill et al., 2015; Parada, Needham & Fuhrman, 2016*). Amplicons were barcoded with Schloss-indexed barcoding primers with Nextera adapters, pooled in equal volumes for sequencing (*Kozich et al., 2013*) and purified with AMPure XP beads. Paired-end sequencing was performed on the Illumina MiSeq platform, 2×300 bp end version 3 chemistry according to the manufacturer's specifications at the Oregon State University's Center for Genome Research and Biocomputing Core Laboratories.

QIIME (v1.9) (*Caporaso et al., 2010b*) was used to process all 16S sequence libraries. Demultiplexed raw reads were trimmed and pair-end sequences merged. Chimeric sequences and sequences with a total expected error of >1 for all bases were discarded. 97%-similarity operational taxonomic units (OTUs) were picked using USEARCH 6.1 (*Edgar, 2010*), QIIME's subsampled open-reference OTU-picking protocol (*Rideout et al., 2014*), and the 97% GreenGenes 13_8 reference database (*McDonald et al., 2012*) to create a starting OTU table. Taxonomy was assigned using UCLUST, and reads were aligned against the GreenGenes database using PyNAST (*Caporaso et al., 2010a*). The aligned reads were then used to reconstruct a phylogenetic tree using FastTreeMP (*Price, Dehal &*-*Arkin, 2010*).

Operational taxonomic units were removed if their representative sequences failed to align with PyNAST to the GreenGenes database or if they were annotated as mitochondria or chloroplasts. After this step, the OTU table had 3,383 unique OTUs, and the number of reads per sample ranged from 1 to 87,262 with a median of 9,742 per sample. OTUs with less than 100 reads across the table were removed resulting in a total of 430 unique OTUs. We did not find that any low count OTUs were associated with one particular sample. After these quality control steps, ten samples were found to contain fewer than 1,000 reads and were thus removed from the dataset (Table S1).

In *R* (v3.4.0) the package *phyloseq* (v1.20.1) (*McMurdie & Holmes, 2013*) was used to rarefy the resulting table to exactly 1,070 sequences per sample, and to calculate from this

rarefied table beta diversity metrics including Bray Curtis, Binary Jaccard, Weighted Unifrac, and Unweighted UniFrac dissimilarities. For beta diversity metrics, the OTU table was first log-transformed in *phyloseq*. Inclusion of all four distance measures allows for a robust analysis of community dynamics including ecological and phylogenetic changes in bacterial abundance (Bray Curtis and Weighted Unifrac) and in the presence or absence of certain bacterial species (Binary Jaccard and Unweighted Unifrac). Also from this rarefied table, alpha diversity metrics including Faith's phylogenetic diversity (*Faith, 1992*), Chao1 statistic (*Chao & Chiu, 2016*), and Simpson's diversity index (*Heip, Herman & Soetaert, 1998*) were calculated in *phyloseq*.

Growth rates, wound healing, and Symbiodiniaceae densities data analysis

All data analysis was conducted in R (v3.4.3) (R Development Core Team, 2017) and all figures were produced using ggplot2 (Wickham & Wickham, 2009). Treatment effects on coral growth and Symbiodiniaceae densities were assessed with linear mixed-effects models (LMMs) with the *lmer* function in R (*Bates et al., 2015*) with temperature, nitrogen, wounding, and the interactions as fixed effects and tank and parent colony as random effects. Wound healing rates were analyzed similarly but with temperature, nitrogen, and their interaction as fixed effects and tank and colony as random effects. Random effects were dropped if not significant in the model according to Chi-squared tests, resulting in final LMMs that have the most parsimonious random effects structure (Zuur et al., 2009). For all models, only colony was a significant random effect, thus tank was excluded from all final models. Model residuals were visually assessed for Gaussian distribution and homoscedasticity. The significance of fixed effects was determined using the anova function from the *lmerTest* package with Kenward-Roger correction for degrees of freedom (Zuur et al., 2009). Multiple comparisons were done with least-squares means using the lsmeans function (Lenth, 2016). An outlier in growth rate (6.93 mg cm⁻² day⁻¹) was removed from the analysis because it was >1.5 larger than the interquartile range of the data. The removal of the outlier did not change the results or interpretation.

Microbial community data analysis

We previously evaluated how these individual and multiple stresses affected bacterial community taxonomic composition, evenness, and diversity (for details see *Maher et al., 2019*). In this study, however, microbial analyses were focused to investigate potential microbiome-dependent mechanisms underlying significant changes in host responses to the treatments and the bacterial community response to differences in nitrogen regimes, two aspects that were not explored in the previous study. Microbial community alpha and beta diversity were evaluated for associations with host responses (i.e., growth rates, wound healing rates, Symbiodiniaceae densities). First, alpha diversity metrics were regressed against host responses using LMMs with host response as the fixed effect and tank and parent colony as random effects. Next, associations between microbial community beta diversity and host responses were assessed with a PERMANOVA using

the *adonis* function in the package *vegan* (v2.4.3) (*Oksanen et al.*, 2007) for each of the four beta diversity metrics. Associations between microbial community beta diversity of wounded corals and treatment effects of temperature, nitrogen, and the interaction were then assessed with *adonis*. Homogeneity of group dispersions for wounded corals was independently assessed for temperature, nitrogen source, colony, and tank with PERMDISP using the *betadisper* function in the package *vegan* (*Oksanen et al.*, 2007). Significant results were ordinated and visualized using NMDS in *phyloseq*. The core microbiome was evaluated from a relative abundance, unrarefied table with the package *microbiome* (v1.5.31) and defined as those taxa present in \geq 50% of samples (*Lahti et al.*, 2017).

Taxa indicative of any treatment combination (e.g., ammonium enriched and wounded under ambient temperature) were investigated using indicator species analysis (*De Caceres & Legendre, 2009; De Caceres, Legendre & Moretti, 2010*). Indicator species analysis involves calculating an indicator value between a species and each group that reflects both the exclusivity, occurring only in a single treatment group, and fidelity, occurring in all samples of a treatment group. The rarefied OTU table was used in the function *multipatt* from the package *indicspecies* (v1.7.6) (*De Caceres & Legendre, 2009; De Caceres, Legendre & Moretti, 2010; Dufrêne & Legendre, 1997*). The *multipatt* function identifies species that are associated with a particular treatment group by calculating an Indicator Value index with a correction for unequal group sizes using the function *IndVal.g.*

RESULTS

Growth rates, wound healing rates, and Symbiodiniaceae densities

Growth rates of individual *P. meandrina* nubbins ranged from 0.49 to 2.38 mg cm⁻² day⁻¹. We did not observe main effects or interactions among seawater warming, nitrogen source, or simulated wounding on *P. meandrina* growth rates (Fig. 1; Table S2). For wound healing, there were no main effects of temperature (LMM; F = 3.57, P = 0.065) or nitrogen enrichment (LMM; F = 2.09, P = 0.14), yet there was a significant interaction between temperature and nitrogen enrichment (LMM; F = 6.51, P < 0.01; Fig. 2; Table 1). Pairwise comparisons revealed that healing rates were reduced ~66% at 29 °C compared to 26 °C under ambient nutrient conditions (P < 0.01; Table S3). At 26 °C, coral nubbins exposed to ammonium enrichment had faster wound healing rates than controls at 29 °C (P < 0.05; Table S3), but coral nubbins exposed to nitrate at 26 °C did not (P = 0.063; Table S3). For corals at 29 °C, nitrogen enrichment removed the negative effect of seawater warming on healing rate regardless of nitrogen source. When comparing the wound healing rates at 29 °C, we found that both ammonium and nitrogen enrichment increased healing rates by ~63% compared to ambient conditions (P < 0.05; Table S3). However, there were no differences in the wound healing rates for P. meandrina nubbins enriched with either ammonium or nitrate across temperature treatments (P > 0.9 for all comparisons; Table S3).

Symbiodiniaceae densities ranged from 1.03 to 10.7×10^5 cells cm⁻². Seawater warming increased Symbiodiniaceae density (LMM; *F* = 7.91, *P* < 0.01; Fig. 3; Table 2). There was

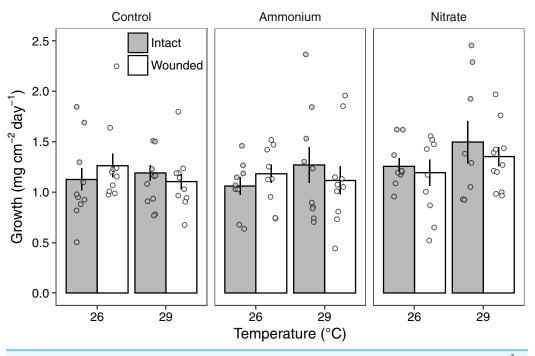


Figure 1 The growth rates of *Pocillopora meandrina* corals across treatments. Growth rates $(mg cm^{-2} day^{-1}; mean \pm SE)$ of intact (gray bars) and wounded (white bars) *Pocillopora meandrina* nubbins under temperature (26 °C, 29 °C) and nutrient (control, ammonium, nitrate) treatments. The points show the distribution of the data. Full-size DOI: 10.7717/peerj.8056/fig-1

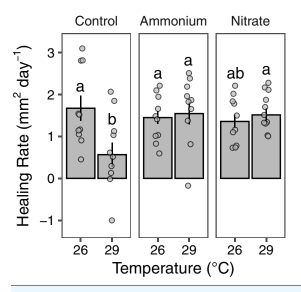


Figure 2 Wound healing rates of *Pocillopora meandrina*. The wound healing rates $(mm^2 day^{-1}; mean \pm SE)$ of wounded *Pocillopora meandrina* nubbins under different temperature and nutrient treatments. The points show the distribution of the data. Letters represent significant difference among treatments. Full-size DOI: 10.7717/peerj.8056/fig-2

also a significant effect of nitrogen source (LMM; F = 4.55, P < 0.05; Fig. 3; Table 2) with ammonium enrichment increasing Symbiodiniaceae densities by 30% relative to control conditions (P < 0.01; Table S4). We did not observe differences in Symbiodiniaceae

Kenward-Roger approximation for degrees of freedom.				
Fixed effects	df	F	Р	
Temperature	1	3.566	0.0653	
Nutrient	2	2.091	0.135	
Temperature × Nutrient	2	6.505	<0.01	

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Note:

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P-values defined as significant at a threshold of 0.05 are highlighted in bold.

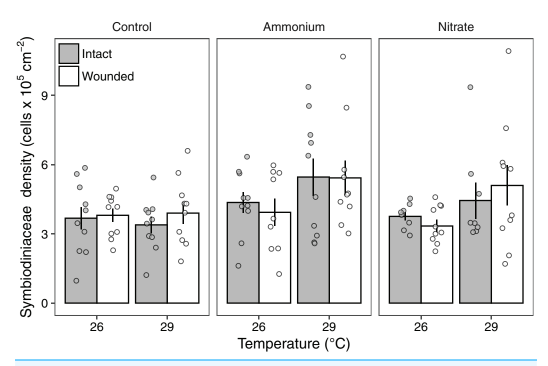


Figure 3 Symbiodiniaceae densities of Pocillopora meandrina corals across treatments. Symbiodiniaceae densities $(10^5 \text{ cells cm}^{-2}; \text{ mean } \pm \text{ SE})$ of intact (gray bars) and wounded (white bars) **Pocillopora meandrina** nubbins under temperature (26, 29 °C) and nutrient (control, ammonium, nitrate) treatments. The points show the distribution of the data. Full-size DOI: 10.7717/peerj.8056/fig-3

Table 2 Linear mixed-effects model results for Symbiodiniaceae densities $(10^5 \text{ cells cm}^{-2})$ with Kenward-Roger approximation for degrees of freedom.

Fixed effects	df	F	Р
Temperature	1	7.909	<0.01
Nutrient	2	4.545	<0.05
Wounded	1	0.0612	0.805
Temperature × Nutrient	2	2.522	0.0857
Temperature × Wounded	1	0.804	0.372
Nutrient × Wounded	2	0.346	0.708
Temperature \times Nutrient \times Wounded	2	0.0405	0.960

Note:

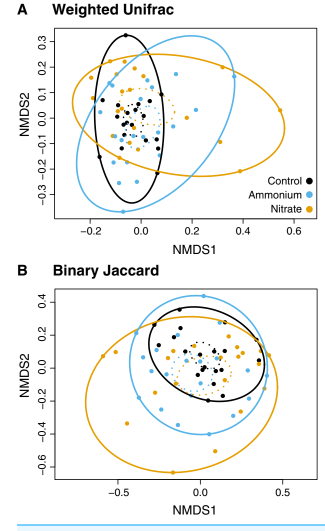
P-values defined as significant at a threshold of 0.05 are highlighted in bold.

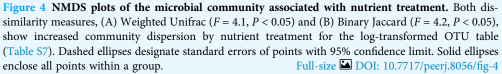
densities for corals enriched with nitrate relative to controls (P = 0.51; Table S4) or nitrate relative to ammonium enrichment (P = 0.18; Table S4). Further, there was no interaction between seawater warming and nitrogen source on Symbiodiniaceae densities (LMM; F = 2.52, P = 0.086; Fig. 3; Table 2). Simulated corallivory had no effects on Symbiodiniaceae densities and there also was no three-way interaction between temperature, nitrogen, and wounded treatments on Symbiodiniaceae densities (Fig. 3; Table 2).

Response of microbial diversity to coral treatments

In addition to measuring changes in the responses of the host and algal symbionts, we also evaluated the coral-associated bacterial communities to fully assess the effects of these stressors on the holobiont. In summary, there were 428 unique bacterial OTUs across the entire rarefied dataset. On average, coral nubbins contained a low diversity of bacterial taxa, around 48.0 ± 4.0 unique OTUs. The most abundant family in the dataset was Endozoicomonadaceae with a mean relative abundance of $67.76\% \pm 3.51\%$ and ranging from 2.90% to 99.44%. Only two samples under increased seawater temperature and ammonium enrichment, one intact and another scarred, contained less than 10% mean relative abundance of Endozoicomonadaceae. Although the Greengenes database identifies this family as Endozoicomonaceae, here the updated taxonomic assignment of Endozoicomonadaceae is used (Bartz et al., 2018; Neave et al., 2016). Other abundant families included Desulfovibrionaceae $(5.51\% \pm 1.55\%$ and ranging from 0.00% to 59.91%), Enterobacteriaceae ($3.79\% \pm 1.39\%$ and ranging from 0.00% to 76.73%), Rhodobacteraceae $(5.88\% \pm 0.89\%$ and ranging from 0.00% to 32.71%), and Moraxellaceae $(2.83\% \pm 0.78\%$ and ranging from 0.00% to 41.22%). On average, coral nubbins with ambient nutrients had a Chao1 index of 68.56 ± 6.06 , although this did not differ with coral nubbins under ammonium or nitrate enrichment (F = 2.053, P = 0.14) which had indices of 47.37 ± 5.56 and 71.47 ± 12.51 , respectively. Coral nubbins in ambient nutrient conditions also were not significantly different (F = 0.377, P = 0.69) via Simpson's diversity (0.472 ± 0.069) when compared to ammonium (0.386 ± 0.055) or nitrate (0.446 ± 0.068) .

While there were no clear associations between alpha or beta diversity with Symbiodiniaceae densities, host growth rates, or host healing rates (Tables S5 and S6) there were differences in the microbial community structure across treatment regimes (Figs. 4 and 5). Differences in beta diversity between treatment groups were identified from the log-transformed community data. While PERMANOVA tests for distinct communities were significant for temperature, wounding, and nitrogen with various dissimilarity measures, all R^2 values were less than 0.1 (Table S7). Therefore, these results were not considered representative of biologically distinct communities. No treatment interactions produced significantly distinct communities (Table S7). However, unlike community dissimilarity measures, there were significant differences between nitrogen treatment group dispersions for Binary Jaccard (PERMDISP, F = 4.210, P < 0.05) and Weighted Unifrac (PERMDISP, F = 4.140, P < 0.05) measures of community dissimilarity (Fig. 4; Table S8). Pairwise comparisons for associations showed that for the Binary Jaccard and Weighted Unifrac measures, corals under both nitrate and ammonium treatments were





significantly more variable compared to coral microbiomes under ambient nutrients conditions (P < 0.05 and P < 0.05, respectively), but nitrate and ammonium were not significantly different from one another (P = 0.63 and P = 0.61, respectively). Group dispersions were also significantly different by temperature with the Binary Jaccard dissimilarity measures (PERMDISP, F = 6.730, P < 0.05, Fig. S2).

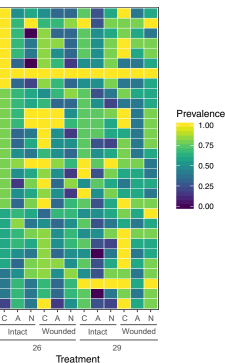
Prevalence and associations of microbial taxa across treatments

A total of 30 OTUs were identified as comprising the core microbiome which were present in at least 50% of samples. Taxa in the core microbiome were evaluated for prevalence across samples in the different treatments (Fig. 5A). A single OTU (ID: 109431) of the family Endozoicomonadaceae was found in every sample (Fig. 5A). Eight OTUs of the families Moraxellaceae, Enterobacteriaceae, and Endozoicomonadaceae were found in Α

В

Core Taxa

OTU89: Gammaproteobacteria, Enterobacteriales, Enterobacteriaceae OTU70: Gammaproteobacteria, Oceanospirillales, Endozoicomonaceae (Endozoicomonadaceae) 4423027: Gammaproteobacteria, Enterobacteriales, Enterobacteriaceae, Klebsiella 4403542: Gammaproteobacteria, Enterobacteriales, Enterobacteriaceae, Klebsiella 4318935; Gammaproteobacteria, Enterobacteriales, Enterobacteriaceae 3633321: Gammaproteobacteria, Pseudomonadales, Moraxellaceae, Acinetobacter, johnsonii ★109431: Gammaproteobacteria, Oceanospirillales, Endozoicomonaceae (Endozoicomonadaceae) 1085703: Gammaproteobacteria, Pseudomonadales, Moraxellaceae, Acinetobacter, johnsonii OTU65: Gammaproteobacteria, Oceanospirillales, Endozoicomonaceae (Endozoicomonadaceae) OTU229: Gammaproteobacteria, Oceanospirillales, Endozoicomonaceae (Endozoicomonadaceae) 816411: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae, Roseivivax 60398: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae 593848: Gammaproteobacteria. Xanthomonadales. Xanthomonadaceae. Stenotrophomonas 4456340: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae, Rhodovulum Таха 4333020: Gammaproteobacteria, Enterobacteriales, Enterobacteriaceae 321533: Cytophagia, Cytophagales, [Amoebophilaceae], SGUS912 256116: Gammaproteobacteria, Pseudomonadales, Pseudomonadaceae 231908: Synechococcophycideae, Synechococcales, Synechococcaceae, Synechococcus 200552: Gammaproteobacteria, Pseudomonadales, Pseudomonadaceae 1147246: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae OTU530: Gammaproteobacteria, Oceanospirillales, Endozoicomonaceae (Endozoicomonadaceae) OTU123: Gammaproteobacteria, Oceanospirillales, Endozoicomonaceae (Endozoicomonadaceae) 706432: Gammaproteobacteria, Pseudomonadales, Moraxellaceae, Acinetobacter 4327730: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae 4306551: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae 251481: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae 3180137: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae * 2932342: Deltaproteobacteria, Desulfovibrionales, Desulfovibrionaceae 163471: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae 1123147: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae



Indicator Taxa

*109431: Gammaproteobacteria, Oceanospirillales, Endozoicomonaceae (Endozoicomonadaceae) OTU216: Gammaproteobacteria, Oceanospirillales, Endozoicomonaceae (Endozoicomonadaceae) OTU58: ABY1 OTU24: Gammaproteobacteria, HTCC2188, HTCC2089 OTU59: Gammaproteobacteria, HTCC2188 * 2932342: Deltaproteobacteria, Desulfovibrionales, Desulfovibrionaceae 2962543: Gammaproteobacteria, Alteromonadales, OM60, Congregibacter 4384063: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae 495067: Actinobacteria, Actinomycetales, Corvnebacteriaceae, Corvnebacterium 765596: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae 1116027: Planctomycetia, Pirellulales, Pirellulaceae 3207: Oscillatoriophycideae, Chroococcales, Xenococcaceae Таха OTU25: Oscillatoriophycideae, Chroococcales, Cyanobacteriaceae OTU3: Oscillatoriophycideae, Chroococcales OTU57: Deltaproteobacteria, Myxococcales 3713320: Acidimicrobiia, Acidimicrobiales, ntu14 3772088: Planctomycetia, Pirellulales, Pirellulaceae, planctomycete, MS30D1 250136: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae, Octadecabacter OTU53: [Saprospirae], [Saprospirales], Saprospiraceae, Lewinella OTU185: [Saprospirae], [Saprospirales], Saprospiraceae 4355907: Alphaproteobacteria, Sphingomonadales, Erythrobacteraceae, Erythrobacter 4294811: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae, Octadecabacter 4405550: Alphaproteobacteria, Rhodobacterales, Rhodobacteraceae, Anaerospora 149505: Alphaproteobacteria, Rhodobacterales, Hyphomonadaceae, Oceanicaulis 1040220: Bacilli, Bacillales, Staphylococcaceae, Staphylococcus ċ A N A Intact 26

Prevalence 1.00 0.75 0.50 0.25 0.00 ċ Ν A N Ċ А Ν Wounded Intact Wounded 29 Treatment

Figure 5 Prevalence of core and indicator microbial taxa by treatment. (A) The core microbiome consists of those OTUs that occur in at least 50% of the samples. (B) Indicator taxa were identified for treatment combinations with Indicator Species Analysis. OTUs were considered indicators for a treatment group if they had a significant indicator value and are designated with an X. Not all treatment combinations contain indicator taxa. Taxa marked with an asterisk are both core and indicators of a specific group. Prevalence is determined by the proportion of samples within a treatment group containing that OTU. Full-size DOI: 10.7717/peerj.8056/fig-5

every control sample. Nearly every coral sample at 29 °C contained an OTU in the family Desulfovibrionaceae (ID: 2932342), whereas this OTU was less prevalent in samples at 26 °C (Fig. 5A). OTUs from the family Rhodobacteraceae were prevalent in wounded coral samples at ambient nutrient levels, however, the specific Rhodobacteraceae species differed based on whether the sample had been exposed to seawater at 26 or 29 °C. OTUs in this family were also less prevalent in wounded coral samples exposed to nitrogen enrichment (Fig. 5A).

Using a rarefied OTU table, indicator species analysis identified associations between bacterial taxa and treatment combinations (Fig. 5B). A total of 25 OTUs were significant indicators with a P < 0.05, however, one of these OTUs had an indicator value < 0.5(Table S9). The control group (ambient nutrients, intact, 26 °C) had a single indicator OTU of the family Endozoicomonadaceae. The indicator value of 0.344 for this OTU (Table S9) is likely due to the presence of this OTU in every sample (Figs. 5A and 5B). While this taxon dominates control corals with >95% relative abundance in all samples, it is not exclusive to the control group. Eight taxa were indicators for high temperature only, including an OTU of the family Desulfovibrionaceae which was also a member of the core microbiome (Fig. 5A). OTUs of the family Rhodobacteraceae were indicators for various treatment groups with wounding, high temperature, or with the combination of the two with nitrogen enrichment (Fig. 5B). The treatment groups of ammonium enrichment, ammonium or nitrate enrichment with wounding, and ammonium enrichment with high temperature did not have any significant indicator taxa (Fig. 5B).

DISCUSSION

Our study tested the hypothesis that different nitrogen (N) sources would have contrasting effects on *P. meandrina* growth, Symbiodiniaceae densities, wound healing, and the coral microbiome in response to seawater warming and simulated parrotfish corallivory. In contrast to our hypothesis, we observed that N source did not have divergent effects on the impacts of seawater warming and simulated corallivory on *P. meandrina* growth rates. However, N source did mediate the impacts of these stressors on Symbiodiniaceae densities and wound healing rates. In fact, intact and wounded corals were able to maintain growth rates under warmer temperatures and across nitrogen regimes. However, ammonium enrichment increased Symbiodiniaceae densities across temperature treatments. We also observed that warmer temperatures reduced tissue regeneration, but that ammonium enrichment counteracted this effect. At the microbial scale, community variability increased with nitrogen enrichment. We also observed distinct bacterial taxa that were indicators of corals under seawater warming, wounding, and the combination of these treatments with nitrogen enrichment.

Potential trade-offs between growth and wound healing under seawater warming

Corals can experience a trade-off between metabolic processes (e.g., growth, gametogenesis) and tissue regeneration, which is often prioritized over coral growth (reviewed by *Henry & Hart*, 2005). Yet our study provides evidence that these tradeoffs are

complex and depend on the abiotic conditions. The P. meandrina nubbins in our study maintained growth rates despite simulated corallivory under higher temperature and nitrogen enrichment. In contrast, Cameron & Edmunds (2014) found that simulated scraper corallivory decreased P. meandrina growth rates by ~42%. The discrepancies in results is perplexing given that our study had similar wound characteristics as those reported by Cameron & Edmunds (2014) and a similar experiment duration (21 days), but may be explained by differences in experimental approaches. Our experiment took place in mesocosms while Cameron & Edmunds (2014) conducted a field study in the back reef. Disparities in species responses can arise from different experimental approaches, and a response measured in the lab does not always translate to the field. However, the present study and others demonstrate that P. meandrina corals are able to maintain growth and calcification rates at ~29 °C (Medellin-Maldonado et al., 2016; Muehllehner & Edmunds, 2008). Moreover, Lenihan & Edmunds (2010) observed that injured P. verrucosa corals under seawater warming (~29 °C) outgrew intact conspecifics but had reduced tissue regeneration. These data are in agreement with our results showing that P. meandrina nubbins under seawater warming maintained growth (Fig. 1) but had lower wound healing rates (Fig. 2).

Wound healing for *P. meandrina* nubbins was ~66% lower at 29 °C than at 26 °C, suggesting that corals at warmer temperatures are less likely to recover from predation events. Seawater warming can reduce energy reserves in *Pocillopora* corals (*Rodriguez-Troncoso, Carpizo-Ituarte & Cupul-Magana, 2010*), which may explain the reduction in wound healing rates we observed. Pocilloporid corals may shift cellular resources to growth over tissue regeneration under warm water conditions. Evidence for this pattern has been observed in *Oculina patagonica* where growth is maintained with seawater warming while wound healing is suppressed (*Serrano, Ribes & Coma, 2017*). These patterns suggest that for *Pocillopora* spp. corals under warmer temperatures, a trade-off may exist between growth and tissue regeneration where growth is prioritized.

Nitrogen supply alleviated this trade-off by counteracting the effects of seawater warming on tissue regeneration rates. However, the effects of nitrogen likely depend on the concentration of nutrients. When considering ammonium enrichment alone, anthropogenic-driven concentrations (~20 μ M) was shown to reduce coral wound healing rates (*Koop et al., 2001*). Yet we showed here that at naturally-occurring concentrations typical of fish excretion (~1–4 μ M) (*Holbrook et al., 2008; Meyer & Schultz, 1985; Shantz et al., 2015*), ammonium supply can increase wound healing rates regardless of temperature (Fig. 2). This pattern indicates that *P. meandrina* may be more robust at recovering from predation events under warmer temperatures when nitrogen from fish excretion is readily available. Similarly, nitrate counteracted the effects of seawater warming on wound healing at 29 °C. In contrast, *Renegar, Blackwelder & Moulding (2008)* found that nitrate enrichment (~10 μ M) reduced wound healing between ~10–60% depending on the coral species. The disparity in our findings can likely be explained by the lower nitrate concentrations used in our study (~4 μ M), which are more environmentally relevant and thus less deleterious to corals. Altogether, our data suggest

that *P. meandrina* corals prioritize growth over tissue regeneration under warm water conditions but that nitrogen supply can neutralize this effect.

Seawater warming and nitrogen source mediate the effects on Symbiodiniaceae densities

A priori, we reasoned that Symbiodiniaceae densities would decrease under seawater warming and increase in response to nitrogen supply. However, under ambient nutrient conditions, seawater warming had little effect on Symbiodiniaceae densities. The lack of temperature effect is surprising given that Symbiodiniaceae densities tend to decline with seawater warming. For instance, *Schmidt et al.* (2016) found that seawater warming (~30 °C) reduces Symbiodiniaceae densities in *P. meandrina* corals. When seawater temperatures are approximately 30 °C in Moorea, Symbiodiniaceae densities for *P. meandrina* can also be reduced by upward of 35% (*Putnam & Edmunds, 2011*). Moreover, *Pocillopora* bleaching in Moorea has been observed for ~4.6 accumulated heat stress weeks (C°-weeks) when temperatures >29.0 °C (*Pratchett et al., 2013*). Such effects may not have been observed in our study due to its short duration (21 days) and that our warm water treatment did not exceed the 29.0 °C threshold required for *Pocillopora* bleaching in this region.

In regard to nitrogen supply, ammonium enrichment increased Symbiodiniaceae densities by ~30% compared to ambient conditions. Yet we did not observe significant changes in Symbiodiniaceae densities in response to nitrate. These results are in accordance with several studies showing that ammonium supply increases Symbiodiniaceae populations compared to corals without nitrogen enrichment, while nitrate enrichment tends to lower Symbiodiniaceae densities (*Chase et al., 2018; Shantz & Burkepile, 2014*). These trends may be due to the energetic costs of nitrate assimilation by Symbiodiniaceae for photosynthesis (*Grover et al., 2003; Patterson et al., 2010*).

We observed no effect of simulated corallivory wounds on Symbiodiniaceae densities compared to intact corals. This contradicts previous research showing that wounds decrease Symbiodiniaceae densities (*Rotjan et al., 2006; Shirur, Jackson & Goulet, 2016*). For example, *Rotjan et al. (2006)* found that parrotfish corallivory reduced Symbiodiniaceae densities of grazed *Orbicella* spp. in Belize. Simulated damage on gorgonians can also reduce Symbiodiniaceae densities in recovering tissues and tissues neighboring lesions (*Shirur, Jackson & Goulet, 2016*). The disparity in our observed results is likely attributable to the degree of damage. In our study, we simulated a single parrotfish scar on each coral nubbin while the coral colonies observed by *Rotjan et al. (2006*) typically had >30 parrotfish bites per colony. Thus, the degree of corallivory damage likely determines the impacts to Symbiodiniaceae populations.

Varying N source increases microbial community variability and produces distinct indicator taxa

Although different nitrogen sources did not produce distinct microbial communities, microbial community dispersion increased significantly with the addition of nitrate or ammonium. Control corals under no stress had stable microbial communities with several core microbial members consistent in every sample (Fig. 5A). Similarly, corals under ambient nutrient regimes shared a degree of stability that was lost with the addition of ammonium or nitrate (Fig. 4). The addition of nitrogen increased sample to sample variability, suggesting a reduction in the host's ability to regulate its microbial associates under stress (*Zaneveld, McMinds & Vega Thurber, 2017*). While our results suggest that microbiome composition changes under nitrogen enrichment stress, these changes are not deterministic and do not result in predictable stressed community states.

We also found that different stress regimes had indicator taxa that contribute to nitrogen cycling and may play an important role in regulating nitrogen availability in the host. In fact, nitrogen-fixation rates in the holobiont may moderate the hosts response to stress (Rädecker et al., 2015). For instance, the abundance of nitrogen fixing bacteria and total nitrogen fixation in the coral increases with higher temperatures (*Cardini et al.*, 2016; Santos et al., 2014). It has been suggested that under increased nitrogen availability, nitrogen fixation rates would be reduced while nitrification and denitrification would increase to reduce internal nitrogen levels and maintain internal nitrogen limitation (*Rädecker et al.*, 2015), which is necessary for a stable symbiosis with Symbiodiniaceae (Muscatine et al., 1989; Yellowlees, Rees & Leggat, 2008). However, one study found that both nitrogen-fixing and denitrifying bacteria in the coral Acropora hemprichii increased in response to increased nitrogen (*Jessen et al., 2013*). Likewise, bacteria of the order Chroococcales, notably Cyanobacteria, are known nitrogen-fixing taxa (Lesser, 2007; Wegley et al., 2007), and in the present study are indicators of wounded corals and corals exposed to excess nitrogen. Bacteria of the family Pirellulaceae are ammonium-oxidizers in sponges and may be conducting nitrification in corals (Gade et al., 2004; Kellogg, Ross & Brooke, 2016; Mohamed et al., 2010). Counterintuitively, taxa of this family are indicators of wounded corals and corals under high temperature and nitrate enrichment in the present study, rather than of ammonium enriched corals. While indicator species analysis of stress treatments identified several potential players in coral nitrogen metabolism, further functional studies are necessary to correlate community composition with changes in host nitrogen-cycling.

Indicator taxa are characteristic of control and disturbed environmental regimes on reefs

Indicator taxa observed in this study support evidence of previous associations between bacterial taxa and holobiont stress (*Maher et al., 2019*; *McDevitt-Irwin et al., 2017*). Bacteria from the order Oceanspirillales are hypothesized to provide a beneficial function to the coral holobiont (*Pantos et al., 2015*), perhaps through their contribution to sulfur cycling (*Raina et al., 2010*). A single taxon from this order is an indicator of control corals and, although it is present in every sample, the relative abundance of this taxon decreases with stress (*Maher et al., 2019*). Another taxon from this order is also an indicator for high temperature along with a taxon from the order Actinomycetales (Fig. 5B), which is proposed to contain antibacterial properties (*Mahmoud & Kalendar*, *2016*; *Nithyanand, Manju & Pandian, 2011*). These potentially beneficial taxa may moderate the host response to stress. Microbes associated with coral mucus have been hypothesized to produce antibiotic activities that select against potentially invasive microbes; however, antibiotic activity from these microbes is typically reduced during a period of high temperature (*Ritchie*, 2006). Therefore, any potential antibacterial properties of the coral tissue in the present study may be compromised in the high temperature treatment we applied here, although we can cannot confirm this hypothesis using these data alone.

Several indicator taxa also suggest increased opportunism in a community compromised with stress. A taxon from the family Desulfovibrionaceae is an indicator for high temperature and nearly all coral samples exposed to seawater at 29 °C contained this taxon (Fig. 5B). Desulfovibrionaceae is a sulfate-reducing bacterium (Bourne, Muirhead & Sato, 2011) and has been associated with increased seawater temperature and coral disease (Gajigan, Diaz & Conaco, 2017; Webster et al., 2011). Similarly, the family Saprospiraceae, which in the present study contains indicators of corals under nitrate enrichment, high temperature, and wounding, has been associated with corals exposed to fertilizer and municipal wastewater and other polluted environments (Jessen et al., 2013; Xia et al., 2008; Ziegler et al., 2016). Several taxa from the family Rhodobacteraceae were indicators of various treatments (Fig. 5B). These taxa are fast-growing and opportunistic (McDevitt-Irwin et al., 2017); however, the degree to which they proliferated with stress depends on the specific combination of stressors (*Maher et al., 2019*). While indicator species analysis can elucidate important patterns in bacterial associations, functional insights into consequences for the microbial community and host are limited. Additionally, further investigation is required to determine whether indicator taxa in a mesocosm experiment reflect indicator taxa on the reef. Of note, the identification of indicator or core microbiome members are limited here by the taxonomic resolution of the Greengenes database used here for taxonomic classification. For instance, the Greengenes database has not been updated since May 2013 while the SILVA v128 database was recently updated in 29/09/2016. Therefore, taxonomic classifications should utilize updated databases so as to avoid discarding sequences not annotated by an outdated database or with vague annotations suggesting contamination.

CONCLUSIONS

As anthropogenic perturbations become more common on reefs, it is crucial to understand how these disturbances may change corals' ability to cope with ongoing biotic processes. Corallivory is a common process on reefs that can exacerbate the response of corals to human impacts (*Rice, Ezzat & Burkepile, 2019*). The current study suggests that nitrogen source can alter the effects of concurrent seawater warming and corallivory on corals, while nitrogen enrichment can have distinct impacts on microbial community variability. Moreover, our results suggest that coral growth may be prioritized over tissue regeneration under warmer temperatures. However, how nitrogen availability and concentration may interact with corallivory and concurrent warming to drive changes to the coral microbial community warrants further research. Moreover, empirical studies are needed to understand how coral immune pathways involved in tissue regeneration respond to these anthropogenic stressors and across nutrient regimes. We observed increased microbiome variability with the addition of nitrogen and identified bacteria that are indicators of different stress regimes. Future research may investigate the functional capabilities of these indicator taxa, particularly in reference to nitrogen cycling, and how their function varies with anthropogenic forcing.

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ADDITIONAL INFORMATION AND DECLARATIONS

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Competing Interests

The authors declare that they have no competing interests.

Author Contributions

- Mallory M. Rice conceived and designed the experiments, performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Rebecca L. Maher analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Rebecca Vega Thurber analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Deron E. Burkepile conceived and designed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.

Field Study Permissions

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Data Availability

The following information was supplied regarding data availability:

Data is available at GitHub: https://github.com/mallorymrice/Poc_2016.git

Microbial data is available at NCBI Sequence Read Archive under the BioProject ID PRJNA549489.

Supplemental Information

Supplemental information for this article can be found online at http://dx.doi.org/10.7717/ peerj.8056#supplemental-information.

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