Heat stress reduces the contribution of diazotrophs to coral holobiont nitrogen cycling

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Efficient nutrient cycling in the coral-algal symbiosis requires constant but limited nitrogen availability. Coral-associated diazotrophs, i.e., prokaryotes capable of fixing dinitrogen, may thus support productivity in a stable coral-algal symbiosis but could contribute to its breakdown when overstimulated. However, the effects of environmental conditions on diazotroph communities and their interaction with other members of the coral holobiont remain poorly understood. Here we assessed the effects of heat stress on diazotroph diversity and their contribution to holobiont nutrient cycling in the reef-building coral Stylophora pistillata from the central Red Sea. In a stable symbiotic state, we found that nitrogen fixation by coral-associated diazotrophs constitutes a source of nitrogen to the algal symbionts. Heat stress caused an increase in nitrogen fixation concomitant with a change in diazotroph communities. Yet, this additional fixed nitrogen was not assimilated by the coral tissue or the algal symbionts. We conclude that although diazotrophs may support coral holobiont functioning under low nitrogen availability, altered nutrient cycling during heat stress abates the dependence of the coral host and its algal symbionts on diazotroph-derived nitrogen. Consequently, the role of nitrogen fixation in the coral holobiont is strongly dependent on its nutritional status and varies dynamically with environmental conditions.

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INTRODUCTION

The association with microbial symbionts is central to the ecological success of reef-building corals in the oligotrophic tropical ocean [1, 2]. The close metabolic coupling between heterotrophic corals and their phototrophic algal symbionts supports their immense productivity, has given rise to their rapid radiation, and has led to the formation of coral reef ecosystems [3–7]. Consequently, the destabilization of this symbiosis in times of anthropogenic environmental change is posing a direct threat to the functioning of coral holobionts, i.e., the ecological unit of corals and their associated microorganisms, and the reefs they support [8–10]. In recent years, ocean warming has repeatedly caused mass coral bleaching, the breakdown of the coral-algal symbiosis, frequently followed by the death of the coral host and subsequent ecosystem-wide reef degradation [11, 12]. An in-depth understanding of the processes underlying the functioning and destabilization of the coral-algal symbiosis is thus required to predict or mitigate the effects of climate change on coral reefs [13–16].

Efficient recycling of organic and inorganic carbon in the coral-algal symbiosis depends on the nutrient-limited state of the algal symbionts [17–19]. In a stable state of the symbiosis, low bioavailable nitrogen availability limits algal growth and results in the accumulation of photosynthates in algal cells [20, 21]. The release of these excess nutrients, in turn, fuels the energy metabolism of the coral host. Although low levels of nitrogen assimilation are required to support the holobiont’s net productivity and growth [22], increases in the coral host’s catabolic activity during heat stress or other environmental stressors may increase nitrogen availability for the algal symbionts. Excess nitrogen availability may destabilize nutrient recycling in the coral-algal symbiosis resulting in a breakdown of coral holobiont functioning and eventually host starvation [15, 23–26]. The stability and functioning of the coral holobiont under changing environmental conditions thus depend on its ability to maintain a limited nitrogen availability for the algal symbionts [16, 25].

Importantly, nitrogen assimilation in the coral holobiont is not limited to the coral host and its algal symbionts. In addition to heterotrophic feeding and the uptake of nutrients from seawater, most coral holobionts also show detectable rates of nitrogen fixation, i.e., the prokaryotic conversion of atmospheric dinitrogen.

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Following 7 days of acclimation, five of the aquaria (one per coral colony) were gradually ramped up to a temperature of 32.9 °C (annual maximum at the collection site in 2017) over the course of 3 days, while the remaining five aquaria were maintained at a constant 29.1 °C (annual mean at the collection site in 2017) [46]. After 7 days at the maximum temperature (i.e., on day 10 of the experiment), replicate fragments were removed from all colonies and were used in triplicate for all molecular and physiological analyses. For molecular analyses, one nubbin per colony and treatment was immediately flash-frozen in liquid nitrogen and stored at −80 °C until further processing. For physiological analyses, one nubbin per colony and treatment was used for each of the incubations, which were performed immediately after the collection. Incubated response parameters included community composition (16S rRNA gene amplicon sequencing), diazotroph community composition (nifH amplicon sequencing), nitrogen fixation activity (acetylene reduction assay), net assimilation of diazotroph-derived nitrogen ($^{15}$N$_2$ isotope labeling + bulk analysis), and partitioning of diazotroph-derived nitrogen in the coral holobiont (NanoSIMS imaging of $^{15}$N$_2$ isotope assimilation). For details on individual response parameters, see below.

DNA extraction, amplification, and sequencing

Individual coral fragments were placed into Ziploc bags, covered in 0.6 mL of AP1 buffer (Qiagen, Hilden, Germany) and air-blasted to remove their tissue from the coral skeleton using air pressure for 1 min. RNase A solution (6 μL from a 100 mg/mL stock) was immediately added to the retrieved tissue slurry in AP1 and 400 μL of the resulting slurry was directly used for DNA extraction with the DNeasy Plant Mini Kit (Qiagen) according to the manufacturer’s instructions. Indexed samples were then cleaned and normalized using the SeqalPrep Normalization Plate Kit (Invitrogen, Carlsbad, CA, USA), pooled at equimolar ratios, and verified on the Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA). Amplicon libraries were sequenced at the KAUST Bioscience core laboratory on the MiSeq platform (Illumina) using 2 × 301 bp overlapping paired-end reads with 20% spiked-in phiX.

For the characterization of the diazotroph community composition, the primers IGK3 5′-CTGGTGGGAGCGTGACGCGTGAATGTGATATATTTAAAGAGGACAGAGTAGCATCCGGCTGA-3′ and DVW 5′-GTGCTGAGCGGGCTTCGAGATGTGATATATTTAAAGAGGACAGAGTAGCATCCGGCTGA-3′ (Illumina overhang adaptor sequences are underlined) [47, 48]. PCRs were run in triplicate using a reaction volume of 10 μL with 5 μL of Qiagen multiplex PCR master mix, 10 ng of DNA template, and final primer concentrations of 0.5 μM for each primer, respectively. PCR cycling conditions consisted of an initial activation step at 95 °C for 15 min, followed by 27 cycles of 94 °C for 30 s, 55 °C for 30 s, and 72 °C for 30 s, followed by a final extension step at 72 °C for 10 min. Successful amplification of all samples was confirmed by gel electrophoresis before products of triplicate PCRs for each sample were pooled. PCR products were subjected to an indexing PCR (8 cycles) using the Nextera XT Index Kit (Illumina, San Diego, USA) according to the manufacturer’s instructions. Indexed samples were then cleaned and normalized using the SeqalPrep Normalization Plate Kit (Invitrogen, Carlsbad, CA, USA), pooled at equimolar ratios, and verified on the Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA). Amplicon libraries were sequenced at the KAUST Bioscience core laboratory on the MiSeq platform (Illumina) using 2 × 301 bp overlapping paired-end reads with 20% spiked-in phiX.

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**Sequence processing and analysis**

For the 16S rRNA gene-based characterization of the prokaryotic community composition, demultiplexed sequence reads (~54,000 per sample on average) were processed in “DADA2” v.1.18.0 [50] running in “R” v.4.0.3 [51]. Following removal of primers and sequencing artifacts, sequences in the DADA2 pipeline were quality filtered with a maximum expected error (max EE) of 2, poor quality read ends were trimmed (truncQ = 2), and forward and reverse reads were truncated at 240 and 160 bp, respectively. The error rates were estimated based on all samples and used for inference of true sequence variants. Following the merging of paired reads and removal of chimeras, amplicon sequence variants (ASVs) were classified against the SILVA database version 138 [52] and all non-bacterial ASVs, as well as those with more than 1% relative abundance in negative extraction, were removed from the dataset in "phyloseq" v.1.34.0 [53]. This yielded an average of ~35,000 retained reads per sample distributed over 1,797 ASVs. The effect of the temperature treatment on the bacterial community composition was assessed at the ASV level. For this, the dataset was cleaned from sparsely distributed ASVs using “pime” v.0.1.0 with a prevalence threshold of 40% (~23,000 reads per sample after filtering) and communities were compared using analysis of similarities (ANOSIM) with Bray–Curtis dissimilarities calculated from relative abundances of ASVs as implemented in “vegan” 2.5-7 [54, 55]. Differential abundance of bacterial taxa between temperature treatments was assessed using linear discriminant analysis effect size (LEfSe) as implemented in the “microbial” R package v.0.0.19 [56].

For the **nifH**-based characterization of the diazotroph community composition, primers were removed from demultiplexed sequence reads (~202,000 per sample) with “Cutadapt” v.2.10 [57]. Using “DADA2” v.1.18.0 [50] running in “R” v.4.0.3 [51], reads were quality filtered with a max EE of 2, poor quality read ends were trimmed (truncQ = 2), and forward and reverse reads were truncated at 220 and 160 bp, respectively. The error rates were estimated based on all samples and used for inference of true sequence variants. Following the merging of paired reads and removal of chimeras, ASVs were translated to the protein level with FrameBot v.1.2.0 using the included nifH reference file [58]. Translated sequences were assigned to **nifH** gene clusters following Meunier et al. [59], using a consensus assignment based on three criteria adapted after Angel et al. [60]: first, sequences were aligned in “MAFFT” v.7.475 [61] and analyzed using a Classification and Regression Tree model following Frank et al. [62]. Second, an Evolutionary Placement Algorithm tree [63] was constructed using the corresponding RaxML implementation and sequences, and were placed on the reference tree after alignment against a reference dataset containing 13 **nifH** gene clusters, as implemented in “DADOAP pipeline” [60]. Third, sequences were queried against the curated **nifH** gene database provided by the Zehr lab (https://wwwzehr.pmc.ucsc.edu/nifH_Database_Public) for gene clustering as well as taxonomic assignment. Based on this approach, ASVs assigned to **nifH** paralogs not involved in nitrogen fixation (clusters IV and V) as well as all ASVs with more than 1% relative abundance in negative extraction were excluded from the dataset in “phyloseq” v.1.34.0 [53]. This yielded an average of ~65,000 retained reads per sample (93,000–186,000 reads including **nifH** paralogs) distributed over 238 ASVs (876 ASVs including **nifH** paralogs). The effect of the temperature treatment on the diazotroph community composition was assessed at the ASV level. To do this, the dataset was cleaned from sparsely distributed ASVs using “pime” v.0.1.0 with a prevalence threshold of 40% (~155,000 reads per sample after filtering) and communities were compared using ANOSIM with Bray–Curtis dissimilarities calculated from relative abundances of ASVs as implemented in “vegan” 2.5-7 [46, 47]. Differential abundance of diazotroph taxa between temperature treatments was assessed using LEfSe as implemented in the “microbial” R package v.0.0.19 [56].

**Acetylene reduction assays**

Nitrogen fixation rates of coral holobionts were quantified indirectly using the acetylene (C2H2) reduction assay [64]. For this, corals were transferred into 1 L glass incubation chambers (Weck, Wehr-Öningen, Germany) and submerged in filtered seawater (0.22 µm) to a final volume of 720 mL. After this, 80 mL of C2H2-saturated seawater (continuously bubbled with C2H2 for 30 min prior to incubations) was added and the chambers were sealed gas-tight with a modified glass lid containing a syringe injection port. Ten percent (20 mL) of the gas headspace was replaced with pure acetylene. Incubation chambers were transferred into temperature-regulated water baths equipped with magnetic stirrers and Radion lights (Ecotech Marine, Inc.) to maintain temperature and light conditions identical to treatment and aquaria conditions. At the beginning and after 24 h of incubation, 2.5 mL gas samples were collected from the chamber headspace with a gas-tight syringe equipped with a push-pull valve (Trajan, Ringwood, Australia) and transferred into vacuum blood collection tubes for subsequent analysis. Ethylene (C2H4) concentrations in the gas samples were determined using gas chromatography with a helium pulsed discharge detector (Agilent 7890B GC system with HP-PLOT/Q column; lower detection limit for C2H4: 0.3 p.p.m.). Rates of C2H4 evolution during the incubations were calculated based on changes in C2H4 concentrations in the headspace taking into account temperature-dependent solubility of C2H4 in seawater [65] as well as background fluxes of seawater controls without corals to account for possible planktonic background C2H4 production (ca. an order of magnitude lower than in coral incubations) [65]. C2H4 evolution rates were converted to their equivalent nitrogen fixation rates using the theoretical conversion factor of 4 and normalized to the surface area of coral fragments [66]. Importantly, the choice of the correct conversion factor is widely debated and may depend on the species and environmental context, as such absolute rates should be interpreted with caution [67]. For rate normalization, surface areas of coral fragments were estimated by 3D computer modeling [68]. Forty to 50 photos were taken of the coral fragments from all angles. The Autodesk Photo-to-3D cloud service (Autodesk, Mill Valley, CA, USA) was used to generate 3D models, which were used for surface area analysis with ReCap Photo v.4.2.0.2 (Autodesk).

**Isotope labeling and bulk C analysis**

To quantify the assimilation of microbially fixed nitrogen in the coral holobiont, an isotope labeling approach with 15N2 isotope-enriched seawater was adapted after the dissolution technique outlined in Wilson et al. [69]. First, filtered seawater (0.22 µm) was degassed by constant stirring at low pressure (<200 mbar) for 30 min and degassed seawater was transferred into 1 L borosilicate bottles with gas-tight rubber septa. To allow for higher 15N2 concentrations during incubations without depleting oxygen (O2) levels, bottles were injected with 25 mL of a gas mixture containing 80% 15N2 (98 atm%) (Cambridge Isotope Laboratories, Tewksbury, MA, USA) and 20% O2 using a gas-tight syringe equipped with a push-pull valve (Trajan). Bottles were vigorously shaken until the bubble volume remained stable and the bottles were stored under constant agitation at 4 °C overnight. 15N2-enriched seawater and freshly filtered seawater (0.22 µm) were mixed at equal parts for the final incubations. Notably, this approach did not compensate for the reduction of pCO2 during the degassing. As such, we cannot rule out that alterations in seawater carbonate chemistry (albeit consistent across all treatments) may have had minor effects on coral physiology during the incubations.

For the isotope labeling, corals were incubated in bubble-free, sealed 1 L glass chambers (Weck) with gas-tight glass lids. Incubation chambers were transferred into temperature-regulated water baths equipped with magnetic stirrers and Radion lights (Ecotech Marine, Inc.) to maintain temperature and light conditions identical to treatment and aquaria conditions. After 24 h, the seawater in the incubations was replaced with equal parts of 15N2-enriched seawater and freshly filtered seawater (0.22 µm), and the chambers were sealed again for another 24 h of incubation. After a total of 48 h, incubations were terminated and corals were sampled for isotope analysis. The tip of coral fragments (~0.5 cm) was removed using a clipper and processed for NanoSIMS analyses as outlined below. The remaining fragment was transferred into a ziplock bag, covered in 2 mL of deionized water, and its tissue was removed from the coral skeleton using air pressure. The resulting tissue slurry was freeze-dried and homogenized subsamples were transferred into tin capsules for bulk N isotope analysis. Total nitrogen content, as well as 15N/14N ratios, were measured using an elemental analyzer (Thermo Flash EA 1112) coupled to a stable isotope mass spectrometer (IRMS, DELTA V Advantage). Notably, we did not quantify concentrations of 14N and 15N in the incubation water. Hence, 15N/14N ratios do not allow for absolute quantification of nitrogen assimilation rates in the present study. However, as all corals were incubated in the same incubation water with identical 15N2 concentrations, levels of 15N enrichments provide a direct reflection of relative nitrogen assimilation rates in these corals. For this, levels of 15N enrichment were expressed using the delta (δ) notation in units per mil of stable isotope ratios calculated as: δ15N (%) = (15N15N_{sample} / 15N15N_{standard} − 1) × 1,000.
NanoSIMS imaging
Following $^{15}$N$_2$ incubations, tips of coral fragments were immediately transferred into a fixative solution (1.25% glutaraldehyde and 0.5% paraformaldehyde in 0.1 M phosphate buffer) at 4°C for 24 h. Fixed samples were washed once in phosphate-buffered saline (1×), decalcified in 0.1 M ethylenediaminetetraacetic acid (exchanged daily) for 2 weeks after which the tissue was dissected into a strip of two to three individual polyps. The tissue was dehydrated in a series of increasing ethanol concentrations (50%, 70%, 90%, and 100%), transferred to acetone, and gradually infiltrated with increasing concentrations of SPURR resin (25%, 50%, 75%, and 100%). Subsequently, tissues were embedded in SPURR resin and cut into 170 nm sections using an Ultracut E microtome (Leica Microsystems, Wetzlar, Germany) and mounted on pulse-discharge silicon wafers. Wafers with sample sections were gold-coated (ca. 15 nm) and analyzed with the NanoSIMS 50 ion probe [70] at the Center for Microscopy, Characterization, and Analysis at the University of Western Australia. Surfaces of samples were bombarded with a 16 keV primary Cs$^+$ beam focused to a spot size of about 100 nm with a current of ~2 pA. Secondary molecular ions $^{12}$C$^{12}$C$^-$, $^{12}$C$^{13}$C$^-$, $^{12}$C$^{14}$N$^-$, and $^{12}$C$^{15}$N$^-$ were simultaneously collected in electron multipliers at a mass resolution ($M/ΔM$) of about 8,000. Images for all targeted secondary molecular ions were collected by rastering the primary beam across the sample with a dwell time of 9 ms per pixel; six planes were recorded for each area. Charge compensation was not necessary. Although samples from all coral colonies were analyzed on the NanoSIMS, only data and images from the colony replicates using a sign test. The relationship between nitrogen fixation rates and $^{13}$N$_2$ assimilation was assessed via a correlation analysis using Pearson’s product–moment correlation coefficient. NanoSIMS measurements of $δ^{15}$N in the host and symbiont tissue/cells were analyzed using a two-way analysis of variance using holobiont compartment and treatment as explanatory variables.

RESULTS AND DISCUSSION
Diazotroph community composition varies with temperature
To assess the role of diazotrophs in the early response of the coral holobiont to heat stress, we sampled the colonies of S. pistillata on day 10 of the experiment (7 days at a maximum temperature of 32.9°C). At this time point, corals from heat stress and control treatments maintained a healthy appearance, similar levels of algal symbiont densities, and showed no visual signs of bleaching (for an extended discussion of processes and timeframes of bleaching during the experiment, please refer to Rädecker et al. [15]). The early heat-stress response did not affect overall bacterial community composition (ANOSIM, $p = 0.145$; Fig. S1). Members of the order Oceanospirillales, which are prevalent in healthy corals and S. pistillata in particular [48, 71, 72], represented the largest component of the coral microbiomes from both treatments (~20% of 16S rRNA sequences).

In contrast to the stable overall bacterial microbiome composition, nifH sequencing revealed that heat stress caused a distinct shift in the community composition of diazotrophs (ANOSIM, $p = 0.049$). This shift was not driven by the loss or the recruitment of novel taxa but rather by variations in the relative abundance of taxa already present in the holobiont (Fig. 1A, B). The diazotroph community was largely dominated by the orders Alteromonadales and Chroococcales accounting for 65% and 27% of nifH sequences in colonies from the ambient control, respectively. Heat stress, however, caused a significant decline in the relative abundance of Chroococcales (LEfSe, $\chi^2 = 3.94$, $p = 0.047$) accompanied by the proportional (albeit not significant) increase of Alteromonadales in the diazotroph community (LEfSe, $\chi^2 = 2.46$, $p = 0.117$) (Fig. 1B). Importantly, Chroococcales were previously identified as endosymbionts in the epithelium of the Caribbean coral Montastrea cavernosa [73] and have been proposed to be an important source of fixed nitrogen for algal symbionts in these corals [74]. Consequently, the observed shifts in diazotroph community composition could directly affect nitrogen availability for other holobiont members during heat stress.

**Fig. 1** Characterization of coral-associated diazotroph communities after 10 days of heat stress. A Relative diazotroph community composition of individual colonies under control (left) and heat stress (right) conditions based on nifH amplicon sequencing. Notably, no archaeal diazotrophs could be detected in the present study. B Change in the relative abundance of dominant diazotroph orders during heat stress relative to control conditions (mean ± SE). Asterisks indicate a significant change ($p < 0.05$) under heat stress compared to control corals.
Using the acetylene reduction technique, we estimated daily nitrogen fixation activity in S. pistillata holobionts. Under ambient control conditions (i.e., 29.1 °C), coral holobionts showed detectable rates of nitrogen fixation around 1.0 ± 0.8 nmol N₂ cm⁻² day⁻¹ in line with rates previously reported for corals from this region (Fig. 2A) [31, 75, 76]. Further, ¹⁵N₂ isotope labeling resulted in enriched ¹⁵N values in the soft tissue of coral holobionts (mean ± SE = 16.0 ± 4.7‰), which positively correlated with nitrogen fixation rates across coral colonies (Pearson's correlation, r = 0.99, p = 0.002, Fig. 2B, C). Although these nitrogen fixation rates may be relatively low compared to other nutrient sources, they are a non-negligible component of the overall nutrient cycling in the nitrogen-limited S. pistillata holobiont.

Algal symbionts assimilate diazotroph-derived nitrogen in a stable coral holobiont

Using the acetylene reduction technique, we quantified daily nitrogen fixation activity in the coral colony with the highest nitrogen fixation activity (Fig. 3A–D). Overall, the algal symbionts showed significantly higher levels of ¹⁵N enrichment than the coral host tissue (118.7 ± 9.4‰ (symbionts), 9.5 ± 1.6‰ (host); Tukey's honestly significant difference (HSD), p < 0.001, Fig. 3F). These observations are consistent with previous reports for S. pistillata from the Great Barrier Reef and show that a large fraction of diazotroph-derived nitrogen is eventually assimilated by the algal symbionts [33]. In addition, NanoSIMS images revealed the presence of small ¹⁵N enrichment hotspots in the host epithelium that displayed two distinct morphologies: oval compartments/cells 2–3 µm in length and clusters of smaller circular and rod-shaped compartments/cells with 1–3 µm in length (Fig. 3B, C). Although the NanoSIMS images do not allow us to elucidate the identity of these hotspots, their shape and location are broadly consistent with previous reports of the presence of endosymbiotic bacteria in the coral holobiont [73, 77–79]. Specifically, the larger oval hotspots resemble observations of Chroococcales in the epithelium of M. cavernosa, whereas the rod-shaped clusters resemble previously described bacterial aggregates in the coral tissue in their structure and localization [73, 77–79]. It is thus plausible that the observed epithelial ¹⁵N hotspots represent endosymbiotic diazotrophs in the tissue of S. pistillata. However, if these ¹⁵N hotspots are indeed diazotrophs, they are not situated near the algal symbionts that are the primary sink of diazotroph-derived nitrogen. In many other phototroph–diazotroph symbioses, diazotrophs predominantly release fixed nitrogen via the passive diffusion of ammonium [80]. Although the ways of nitrogen release by coral-associated diazotrophs are currently unknown, it is plausible that passive ammonium release and/or the catabolic breakdown of diazotrophs during host digestion may contribute to the inorganic nitrogen pool in the holobiont. If this is the case in S. pistillata, the uneven partitioning of diazotroph-derived nitrogen between the coral host and its algal symbionts likely reflects the respective metabolic demands of symbiotic partners for external nitrogen sources. In this scenario, the nitrogen-limited state of algal symbionts might support strong concentration gradients in the host tissue, which would enable efficient assimilation of exogenous nitrogen (derived from either diazotrophs or from the surrounding seawater) [81]. In a stable holobiont state, diazotrophs thus provide a nitrogen source to support algal symbiont growth and allow for net productivity of the coral holobiont (Fig. 4A).

Reduced assimilation of fixed nitrogen by algal symbionts despite increased nitrogen fixation activity during heat stress

On day 10 of the experiment, heat stress caused a 74% increase in nitrogen fixation rates (indirectly quantified as acetylene reduction) compared to coral holobiont rates from ambient control conditions (sign test, z = 2.24, p = 0.025, Fig. 2A). Similar increases in coral-associated nitrogen fixation during heat stress have been previously reported for other coral species [40, 41], suggesting that the here reported processes are unlikely restricted to the S. pistillata holobiont, but may be a common feature of coral-associated diazotroph communities. ¹⁵N₂ isotope labeling in this study revealed that the additional fixed nitrogen was not assimilated in the soft tissue of the holobiont, in which ¹⁵N enrichment decreased by 30% compared to ambient controls (sign test, z = 2.24, p = 0.025, Fig. 2B). NanoSIMS analysis corroborated this observation, revealing that the ¹⁵N enrichment remained stable in the host tissue (Tukey’s HSD, p = 0.976) but declined by 53% in the algal symbionts compared to ambient controls (Tukey’s HSD, p < 0.001, Fig. 3E, F). Furthermore, we did not detect any epithelial ¹⁵N enrichment hotspots in corals exposed to heat stress, suggesting lower abundances of
heat stress (Fig. 2A), hence, was likely not driven by endosymbiotic microbes better adapted to exploit the altered nutrient regime during heat stress [82]. Mass spectrometry and NanoSIMS analyses only quantify the assimilation of anabolically incorporated $^{15}$N into the coral and symbiont cells. As such, the assimilation of diazotroph-derived nitrogen in other holobiont compartments (i.e., the coral skeleton and the surface mucus layer) is not accounted for in our analyses. Indeed, a study by Moynihan et al. [34] recently suggested that endolithic microbes in the coral skeleton were the main source and sink of diazotroph-derived nitrogen in the closely related Pocillopora acuta holobiont. Further, El-Khaled et al. [85] showed that moderate increases in ammonium and phosphate by the coral host during heat stress would directly affect nutrient availability in other compartments of the coral holobiont such as the skeleton (Fig. S2 [15]). It is thus plausible that the increase in nitrogen fixation activity described here was predominantly driven by an increase in the activity of endolithic microbes associated with the holobiont. While some studies suggested that endolithic microbial communities may eventually release some of their nutrients to the coral tissue [34, 42, 86, 87], our results clearly show that diazotroph-derived nitrogen (regardless of its origin within the holobiont) is an insignificant source of nitrogen for the coral-algal symbiosis during heat stress. As such, the increase in stimulated nitrogen fixation may primarily be absorbed within the endolithic community itself, thereby supporting the frequently documented rapid proliferation of endolithic microbes during heat stress [86, 88, 89]. Further, nitrogen fixation rates correlate with denitrification activities in some Red Sea corals [75]. Although we did not quantify denitrification rates in the present study, increases in the abundance and/or activity of denitrifying microbes may have contributed to the removal of diazotroph-derived nitrogen from the coral holobiont during heat stress. Taken together, we conclude that the excess availability and release of ammonium through host catabolic processes, combined with increased 

![Fig. 3 NanoSIMS imaging of $^{15}$N$_2$ assimilation in the coral holobiont after 10 days of heat stress. A Mosaic of hue saturation images of $^{15}$N/$^{14}$N ratios illustrating the spatial distribution of $^{15}$N enrichment in the coral tissue under control conditions. B, C Enlarged $^{15}$N/$^{14}$N NanoSIMS images of epithelial $^{15}$N$_2$ hotspots highlighted in A. Epithelial hotspots were primarily characterized by two distinct morphologies: oval compartments/cells with 2–3 µm length (yellow circles) and clusters of smaller rod-shaped compartments/cells with 1–3 µm in length (white circles). D, E Hue saturation images of $^{15}$N/$^{14}$N ratios illustrating $^{15}$N assimilation in the coral host and algal symbionts under control (D) and heat stress (E) conditions. F $^{15}$N$_2$ assimilation in the coral tissue and algal symbionts under control (blue) and heat stress (red) conditions based on NanoSIMS images ($n = 15$ per treatment). Differing letters above the boxplot indicate significant differences between groups ($\rho < 0.05$). All scale bars are 10 µm.](https://example.com/fig3.png)
microbial utilization of fixed nitrogen, likely reduce the relative contribution of diazotroph-derived nitrogen to holobiont nitrogen cycling during heat stress.

**The functional importance of diazotrophs in the coral holobiont**

Nitrogen fixation rates in coral holobionts are highly variable—they can differ between host species, locations, and fluctuate depending on local environmental conditions [34, 37, 38, 41, 42, 82, 87, 90]. Indeed, the molecular, physiological, and ultrastructural characterization of nitrogen fixation presented here paints a complex picture of the role of diazotrophs in coral holobiont functioning. Our results suggest that endosymbiotic diazotrophs in the coral epithelium actively fix nitrogen in the coral holobiont. The characterization of tissue-associated diazotroph communities suggests that the observed 15N hotspots may resemble individual endosymbiotic diazotrophs of the order Chroococcales and bacterial aggregates potentially including diazotrophs of the order Alteromonadales. Further, the disappearance of these 15N hotspots during heat stress coincided with a decline in Chroococcales nifH relative sequence abundance and the reduced 15N assimilation in algal symbiont cells. Taken together, our findings indicate that endosymbiotic diazotrophs in the coral tissue, especially Chroococcales, represent an important source of nitrogen for algal symbionts under nitrogen-limited conditions. Hence, the association with diazotrophs may supplement holobiont nutrition and enable net nitrogen-limited conditions. However, anthropogenic impacts such as eutrophication and ocean warming likely reduce the benefits of hosting diazotrophs in the coral holobiont.

**DATA AVAILABILITY**

All raw sequencing data have been deposited under BioProject ID PRJNA741490 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA741490). Further, ASV abundance tables (16S rRNA and nifH gene amplicons) as well as physiological and NanoSIMS data have been deposited at Zenodo.org and are freely available at https://doi.org/10.5281/zenodo.5552980.

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AUTHOR CONTRIBUTIONS
NR, CP, MP, J-BR, and CRV conceived the experiment. NR, CP, HG, and FR conducted the experiment. NR, CP, AG, LP, PG, and JB performed sample and data analysis. All authors contributed to writing and revising the manuscript.

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