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Dark Production of Extracellular Superoxide by the Coral *Porites astreoides* and Representative Symbionts

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Zhang T, Diaz JM, Brighi C, Parsons RJ, McNally S, Apprill A and Hansel CM (2016) Dark Production of Extracellular Superoxide by the Coral Porites astreoides and Representative Symbionts. Front. Mar. Sci. 3:232. doi: 10.3389/fmars.2016.00232 The reactive oxygen species (ROS) superoxide has been implicated in both beneficial and detrimental processes in coral biology, ranging from pathogenic disease resistance to coral bleaching. Despite the critical role of ROS in coral health, there is a distinct lack of ROS measurements and thus an incomplete understanding of underpinning ROS sources and production mechanisms within coral systems. Here, we quantified in situ extracellular superoxide concentrations at the surfaces of aquaria-hosted Porites astreoides during a diel cycle. High concentrations of superoxide (~10's of nM) were present at coral surfaces, and these levels did not change significantly as a function of time of day. These results indicate that the coral holobiont produces extracellular superoxide in the dark, independent of photosynthesis. As a short-lived anion at physiological pH, superoxide has a limited ability to cross intact biological membranes. Further, removing surface mucus layers from the P. astreoides colonies did not impact external superoxide concentrations. We therefore attribute external superoxide derived from the coral holobiont under these conditions to the activity of the coral host epithelium, rather than mucus-derived epibionts or internal sources such as endosymbionts (e.g., Symbiodinium). However, endosymbionts likely contribute to internal ROS levels via extracellular superoxide production. Indeed, common coral symbionts, including multiple strains of Symbiodinium (clades A to D) and the bacterium Endozoicomonas montiporae LMG 24815, produced extracellular superoxide in the dark and at low light levels. Further, representative P. astreoides symbionts, Symbiodinium CCMP2456 (clade A) and E. montiporae, produced similar concentrations of superoxide alone and in combination with each other, in the dark and low light, and regardless of time of day. Overall, these results indicate that healthy, non-stressed P. astreoides and representative symbionts produce superoxide externally, which is decoupled from photosynthetic activity and circadian control. Corals may therefore produce extracellular superoxide constitutively, highlighting an unclear yet potentially beneficial role for superoxide in coral physiology and health.

Keywords: coral, superoxide, reactive oxygen species, photosynthesis, Symbiodinium, stress

INTRODUCTION

Cellular redox homeostasis is required for the proper biological functioning and health of all living systems. Reactive oxygen species (ROS)—including the superoxide anion $(O_2^{\bullet-})$, its conjugate acid the hydroperoxyl radical (HO₂), hydrogen peroxide (H₂O₂), and the hydroxyl radical (HO[•])—can threaten this homeostasis, based on their ability to degrade vital cell components such as lipids and DNA (Lesser, 2006), as well as their capacity to activate programmed cell death pathways (Cai and Jones, 1998). ROS are an inevitable consequence of aerobic life and commonly arise as metabolic byproducts in oxygenated cells. In contrast to such hazardous intracellular ROS production, however, recent evidence highlights the benefits of extracellular ROS to the physiological health and functioning of macro- and micro-organisms, including bacteria (Saran, 2003; Buetler et al., 2004), fungi (Aguirre et al., 2005), algae (Kustka et al., 2005; Rose et al., 2005, 2008b; Weinberger, 2007; Roe and Barbeau, 2014), plants (Minibayeva et al., 2009), and animals (Babior, 1999). For instance, ROS can serve in autocrine growth (Oda et al., 1995; Saran, 2003; Buetler et al., 2004), immune defense (Babior, 1999; Weinberger, 2007), metal nutrient acquisition (Rose et al., 2005; Fujii et al., 2010; Roe and Barbeau, 2014), wound repair (Minibayeva et al., 2009), and cell differentiation (Aguirre et al., 2005).

In corals, intracellular ROS production, typically stimulated by excessive heat exposure, has a negative role in coral health by causing oxidative stress and bleaching, or the loss of essential photosynthetic endosymbiotic Symbiodinium algae (Lesser, 2011). The key source of intracellular ROS in stressed corals is thought to be disrupted photosynthetic electron transport in Symbiodinium and the coral mitochondria, which leads to the reduction of endogenous photosyntheticallyderived O₂ (Jones et al., 1998; Tchernov et al., 2004; Lesser, 2006; Venn et al., 2008; Weis, 2008). In contrast, corals also produce ROS externally (Saragosti et al., 2010; Shaked and Armoza-Zvuloni, 2013), which involves the reduction of exogenous O₂ in seawater at the coral surface. Extracellular and intracellular ROS production therefore occurs via independent pathways in corals, as depicted in Figure 1. Specifically, photosynthetic superoxide production should not contribute directly to external superoxide concentrations (Figure 1). Photosynthetically derived intracellular ROS in corals occurs within Symbiodinium, which localize in membrane-bound symbiosomes inside the coral host's inner gastrodermal cell layer. Despite the ability of hydroperoxyl radical (the protonated form of superoxide, $\mathrm{HO}_2^{\bullet})$ to diffuse through membranes at a rate approaching that of water, superoxide anion $(O_2^{\bullet-})$ dominates in pH conditions above 4.8 (Bielski et al., 1985) and has a limited ability to diffuse across membranes (Korshunov and Imlay, 2002). Recent evidence that the symbiosome has a pH of \sim 4 (Barott et al., 2015) may indicate that superoxide could diffuse from the symbiosome into the gastrodermal coral cells. Yet, unprotonated superoxide would dominate once in coral gastrodermal cells, which have a pH of \sim 7 (Venn et al., 2009; Barott et al., 2015). Thus, as an anion at the physiological pH conditions within the coral tissues, superoxide is likely incapable of crossing the many biological membranes that separate the photosynthetic electron transport chain of *Symbiodinium* from seawater (Rose, 2012) (see **Figure 1**). In addition, the typical intracellular lifetime ($\sim \mu s$) and diffusive distance of superoxide (~ 100 's of nm) (Lesser, 2006) are also too short to allow extensive transport from its site of production within *Symbiodinium*.

Instead, extracellular superoxide production by corals likely originates from the coral host's epithelial cells or epibionts, including bacteria within the coral mucus layer (Figure 1). This external superoxide production probably occurs through the activity of extracellular oxidoreductase enzymes, such as NAD(P)H oxidases (Babior, 1999; Saran, 2003; Aguirre et al., 2005; Kustka et al., 2005; Rose et al., 2005; Weinberger, 2007; Andeer et al., 2015). Indeed, the inhibition of coral-derived extracellular superoxide production by the broad spectrum NADPH oxidoreductase inhibitor diphenyleneiodonium (DPI) suggested that extracellular superoxide production by corals is mediated by this large and diverse family of NAD(P)H-oxidizing enzymes (Saragosti et al., 2010), similar to other organisms, such as mammalian leukocytes (Babior, 1999), fungi (Aguirre et al., 2005), macroalgae (Weinberger, 2007), phytoplankton (Kustka et al., 2005; Rose et al., 2005), bacteria (Andeer et al., 2015), and plants (Minibayeva et al., 2009). Photosynthesis may play an indirect role in the production of external superoxide by corals by supplying NADPH that stimulates the activity of superoxide-producing oxidoreductases on the coral surface (Saragosti et al., 2010). This indirect involvement of photosynthesis in extracellular superoxide generation may also explain previous observations of light-stimulated extracellular superoxide production by phytoplankton, including the colonial diazotroph Trichodesmium (Hansel et al., 2016) and Symbiodinium (Saragosti et al., 2010), which also likely generate extracellular superoxide via cell surface associated NAD(P)H oxidoreductases.

In addition to their production through independent physiological pathways, intra- and extracellular superoxide may also have fundamentally different physiological consequences to coral health. For instance, dark production of extracellular superoxide has been observed in unstressed corals (Saragosti et al., 2010), *Symbiodinium* (Saragosti et al., 2010), and bacterial genera commonly associated with corals (Diaz et al., 2013), suggesting a constitutive role for external superoxide in healthy biological systems. Although the underlying reasons for this production remain unresolved, recent observations suggest that external ROS production, and NADPH oxidoreductases putatively involved in superoxide production, may promote coral thermotolerance (Dixon et al., 2015), immune defense (Banin et al., 2003; Libro et al., 2013), and prey acquisition (Armoza-Zvuloni et al., 2016).

The goal of the current study was to investigate the dark production of external superoxide by unstressed corals and cultured representatives of their microbial symbionts. While direct measurements of ROS within coral tissues are difficult and artifact-prone, recent advances in chemiluminescent techniques have made non-invasive measurements of external superoxide production by corals and their symbionts more tractable. Here, we measured external superoxide production



FIGURE 1 | Simplified schematic to illustrate the points of origin (denoted by a dot) for superoxide (O₂) (in red) and hydrogen peroxide (H₂O₂) (in blue) within coral tissue. The outer membranes of host cells, mitochondria (blue oval), *Symbiodinium* (green circle), and bacteria (pink rod) are represented as potential sites of intracellular and extracellular ROS production (not to scale). While this schematic may not depict all the possible sources of superoxide and hydrogen peroxide production in corals, it is meant to provide a simple conceptual framework. In particular, the superoxide anion only has a limited ability to cross membranes at physiological pH, and thus intracellular formation of superoxide does not significantly impact extracellular pools. In contrast, hydrogen peroxide can travel throughout the coral tissue (transport denoted by dotted lines), and thus intracellular hydrogen peroxide sources can contribute to extracellular pools. Superoxide produced outside of cells via extracellular or transmembrane proteins can interact with surrounding cells. For instance, extracellular superoxide production by *Symbiodinium* or bacteria could directly interact with coral cells (see top gastrodermal cell). Direct measurements of external superoxide would likely include superoxide formed at the plasma membrane of coral epithelial cells as well as extracellular superoxide generated by bacteria in the mucus layer.

from aquaria-hosted *Porites astreoides* corals in the dark and low light levels at two times during the day. *P. astreoides* was chosen as a representative coral because it is found in a variety of reef habitats around the world and has been associated with Caribbean reefs for millennia (Pandolfi and Jackson, 2006). The high recruitment success of *P. astreoides* has helped to increase the relative abundance of this species on Caribbean reefs (Green et al., 2008) and its microbial community has been well-described (Rohwer et al., 2002; Morrow et al., 2012; Rodriguez-Lanetty et al., 2013). In addition to *P. astreoides*, we identified and then examined representative cultures of its microbial symbionts for the ability to produce extracellular superoxide within corals. As coral-*Symbiodinium* relationships can be flexible, we further examined the ability of diverse *Symbiodinium* clades to produce extracellular superoxide.

MATERIALS AND METHODS

Coral Collection and Aquaria Incubations

Porites astreoides colonies were collected from 3–9 m water depth at 3 sites in Bermuda in July 2013: Hog Breaker Reef (N 32° 27.5′ W 64° 49.8′), an unnamed patch reef (N 32° 26.042′ W 64° 49.248′), and Three Hill Shoals (N 32° 41′ W 64° 73.3′). The mid-day photosynthetically active radiation (PAR) at coral depth in these reefs was \sim 300–500 µmol photon m⁻² s⁻¹. Colonies ranged from 15–20 cm diameter, and colonies of similar size were specifically selected to avoid the need for surface area normalization of superoxide measurements. *P. astreoides* colonies were placed in collection bags, which were sealed underwater. Upon surfacing, the corals were transferred into a cooler containing reef water from the site and transported (<1 h) to the laboratory at the Bermuda Institute of Ocean Sciences (BIOS). Ten colonies were taken from each site (n = 30) in compliance with the BIOS Collection and Experimental Ethics Policy (CEEP). This was considered Limited Impact Research, and as such a collection permit was not required.

At BIOS, specimens were placed in outdoor fiberglass holding aquaria for 2 days to allow them to acclimate to ex situ conditions. Following acclimation in the holding aquaria, P. astreoides colonies were transferred to indoor aquaria outfitted with a lighting system configured with full spectrum LED lamps and a light controller that simulates a daily light cycle $(PAR = \sim 100 \,\mu mol \text{ photon } m^{-2} \text{ s}^{-1})$. PAR at the surface of the corals in the tanks was $61 \,\mu$ mol photon m⁻² s⁻¹, which is lower than the mid-day light conditions on the reef (see above). While these levels are low, they are in the range of known compensation ranges for photosynthetic activity [3-233 μ mol photon m⁻² s⁻¹, (Mass et al., 2007)]; yet, where P. astreoides falls within this range is currently unknown, and thus the level of photosynthesis occurring under the low light conditions used for this study is unknown. Three coral colonies were placed in each of three replicate aquaria and acclimated for an additional 2 days before starting superoxide measurements. Control aquaria, in which corals were absent, were incubated under the same conditions. Each aquarium was fed with local seawater from Ferry Reach. Briefly, water from 20 m off shore and one meter in depth was pumped through a BIOS flow-through sample system that consisted of a coarse mesh filter followed by a step filtration system of 50 $\mu m,$ then 5 $\mu m,$ to remove larger organisms including some planktonic grazers. The flow-through system was equipped with air wand bubblers to provide constant aeration and mixing.

For a subset of corals, mucus was removed from the coral surfaces by inverting coral heads for 2 h, which was previously determined to be the optimal time for removing the majority of surface mucus layer while not compromising coral health.

Extracellular Superoxide Measurements of Aquaria-Hosted Corals

Superoxide concentrations were measured with a flow-through FeLume Mini system (Waterville Analytical, Waterville ME) via the specific reaction between superoxide and the chemiluminescent probe methyl Cypridina luciferin analog (MCLA, Santa Cruz Biotechnology). The FeLume system is composed of two separate fluid lines, one of which is dedicated to the analyte solution and the other to the MCLA reagent. The reagent was amended with 100 µM diethylenetriaminepentaacetic acid (DTPA) in order to sequester trace metal contaminants that would otherwise significantly reduce the lifetime of superoxide. To measure superoxide, both the analyte solution and the MCLA reagent were independently flushed through the FeLume system at an identical flow rate (i.e., 2 mL min⁻¹) using a peristaltic pump. The solutions converge in a spiral flow cell immediately adjacent to a photomultiplier tube, which continuously acquires data that is displayed in real time using a PC interface. The MCLA reagent consisted of $4.0 \,\mu M$ MCLA with 100 µM DTPA in 0.10 M MES buffer adjusted to pH 6.0. In previous studies, concentrations of the MCLA reagent ranged from 1 to 5 µM (Rose et al., 2008a, 2010; Heller and Croot, 2010; Roe et al., 2016; Zhang et al., 2016), depending on the sensitivity required.

For measurements of superoxide in aquaria, a cart adjacent to the aquaria was used to host the FeLume in order to reduce the length of tubing (travel time) required to transport the water from the aquaria to the instrument. To eliminate abiotic photochemical processes that may produce superoxide during in situ measurements, opaque tubing was used, and the entire analytical system was shielded from light. Seawater was directly pumped into the FeLume by placing the analyte tubing at discrete positions within the aquarium in relation to the corals. For each aquarium, a number of superoxide measurements were conducted. First, a baseline was acquired from a reagent blank to account for the chemiluminescence signal resulting from the autooxidation of MCLA. The reagent blank consisted of aged filtered seawater (AFSW) amended with DTPA (AFSW+DTPA). To prepare AFSW+DTPA, Ferry Reach water was collected from the flow-through system and filtered (0.2- μ m SterivexTM, Millipore). The filtered seawater was aged in the dark (>12 h) to deplete superoxide, amended with 50 µM DTPA, and left in the dark for an additional 12 h to allow DTPA to complex trace metals.

After running the reagent blank, the seawater signal was measured at two background locations in the aquarium: First, a position in the water at the top of the tank and second, at the same depth as the corals but 2–3 cm away (horizontal

distance). Data were acquired until a stable, steady-state reading was achieved ($< \sim 4\%$ coefficient of variation). The tubing was then moved to the seawater immediately adjacent to the coral (<1 cm away from the coral, without touching the surface of the coral). Seven measurements were made at various locations along the coral surfaces. The tubing was then moved back to the background seawater at positions that were equivalent to the initial background measurements. Data were acquired for at least 2 min to ensure that the signals returned to the original background values. Finally, superoxide dismutase (SOD), an enzyme that rapidly degrades superoxide, was added (800 U L^{-1} SOD) to an aliquot of water taken from the coral surface in order to confirm that the signal was in fact attributable to superoxide. SOD routinely lowers the signal derived from the reagent blank, likely reflecting a small concentration of superoxide present in the AFSW+DTPA or more likely produced during the autooxidation of MCLA (Hansard et al., 2010).

Before converting superoxide chemiluminescence signals to concentrations, all measurements were corrected by subtracting the signal obtained from the reagent blank. Superoxide concentrations were not corrected for decay that occurred within the tubing during the brief time required to pump the sample from the point in the aquaria into the FeLume, and thus the values represent conservative estimates. Chemiluminescence signals were calibrated by generating stock solutions of superoxide from potassium dioxide (KO₂). Considering the short lifetime of superoxide, standards were prepared immediately before analysis. Primary stock solutions were made by dissolving a small quantity of KO₂ in a basic matrix (0.03 N NaOH, 50–100 μ M DTPA, pH = 12.5). Superoxide concentrations in primary standards were quantified by measuring the difference in absorbance at 240 nm before and after the addition of SOD (\sim 8 U mL⁻¹, final) and then converting to molar units based on the molar absorptivity of superoxide $(2183 \text{ L mol}^{-1} \text{ cm}^{-1})$ at 240 nm, pH = 12.5, and corrected for the absorption of hydrogen peroxide formed during decay) (Bielski et al., 1985). Primary stocks had to be substantially diluted in order to generate representative concentrations for analysis on the FeLume. To generate secondary stocks (4-55 nM), the primary stock solution was diluted into AFSW+DTPA. For each calibration point, a separate FeLume run was conducted as follows: First, the reagent blank was run until a stable baseline (<4% coefficient of variation) was achieved for ~ 1 min. Then the secondary standards were pumped directly into the FeLume, and the decay of superoxide was monitored for at least 1 min. Finally, SOD was added to the secondary standard ($\sim 0.8 \text{ U mL}^{-1}$, final) in order to confirm that the signal was attributable to superoxide.

Chemiluminescence signals collected during the decay of superoxide standards were extrapolated backwards in time to the point when the primary standard was quantified spectrophotometrically. Extrapolations assumed first order decay kinetics because decay data were log-linear. Multipoint calibration curves were constructed based on the linear regression of several pairs of extrapolated luminescence signals vs. superoxide concentrations. Calibrations yielded linear curves (e.g., $R^2 > 0.94$), with a sensitivity that ranged from 0.09 to 0.11 luminescence units per pM superoxide. The half-life of

superoxide in the calibration matrices ranged from 0.89 to 2.11 min and the extrapolation time was 0.93-1.27 min. The detection limit of this method, calculated as three times the standard deviation of a series of blank measurements, was 0.13 nM. Superoxide measurements were compared using unpaired two-sample *t*-tests.

Ancillary Data from Aquaria Water

Water temperature, pH, salinity and dissolved oxygen (DO), were measured using a YSI 556MPS handheld multiparameter system. The YSI pH probe consists of a glass bulb filled with a stable pH = 7 reference solution. The potential across the glass surface created between the water sample and reference solution is measured and converted to pH using a 3-point calibration (pH 4, 7, 10). For the analysis of metals, including total dissolved iron (Fe), manganese (Mn), and copper (Cu), triplicate samples were taken from aquaria surface water on days 1, 4, and 7. Samples were filtered (0.2- μ m), amended with 2% (v/v) nitric acid (HNO₃, trace metal grade), and analyzed using inductively coupled plasma mass spectroscopy (ICP-MS).

Identification of P. astreoides Symbionts

Bacteria, archaea, and Symbiodinium were identified from tissue and mucus homogenates from six P. astreoides colonies and mucus from 10-12 colonies. Samples for symbiont identification were taken after all superoxide measurements. Tissue was frozen (-80°C), airbrushed, and extracted using the Power Plant DNA extraction kit (Mo Bio laboratories), using a modified protocol described previously (Apprill et al., 2013). Briefly, the modifications to the extraction protocol included two enzyme digestions including a 10 units/µL addition of lysozyme (Ready-Lyse Lysozyme Solution, #R1802M) for 10 min at room temperature and an addition of 20 mg/mL of proteinase K (Qiagen) incubated at 65°C for 10 min at 1000 RPM. Processing of mucus samples followed a method modified from Giovannoni et al. (1990, 1996). Briefly, mucus (1 mL) was filtered $(0.2 \,\mu\text{m})$ under gentle vacuum (~100 mm Hg), and stored in lysis buffer (20 mM EDTA, 400 mM NaCl, 0.75 M sucrose, 50 mM Tris.HCl) at -80°C. For DNA extraction, sodium dodecyl sulfate (1% final) and proteinase K (200 μ g mL⁻¹ final) were added to the sample and incubated at 37 $^\circ C$ for 30 min and then at 55 $^\circ C$ for 30 min. The lysates were extracted with an equal volume of phenol:isoamylalcohol:chloroform (25:1:24) followed by two subsequent equal volumes of isoamylalcohol:chloroform (1:24). Purification of the DNA was conducted using sodium acetate (3 M) and isopropanol for at least 1 h at -20° C and centrifuged $(20,000 \times g)$ at room temperature for 30 min. The resulting pellet was washed with 80% ethanol, vortexed for 30 s and centrifuged at $16,000 \times g$ for 10 min.

Bacterial and archaeal community composition was assessed by targeting the V4 region of the SSU rRNA gene using modified primers, 515F and 806RB (Apprill et al., 2015). Triplicate 25 μ L PCR reactions were conducted per sample, which consisted of 1.25 U GoTaq Flexi DNA Polymerase (Promega), 5X Colorless GoTaq Flexi Buffer, 2.5 mM MgCl₂, 200 μ M dNTP mix, 200 nM of each barcoded primer, and 1–4 ng of genomic template. The reactions were carried out on a thermocycler

(Bio-Rad Laboratories) with the following conditions: An initial denaturation step at 95°C for 2 min; 27-32 cycles of 95°C for 20 s, 55°C for 15 s, and 72°C for 5 min; concluding with an extension at 72°C for 10 min. Reaction products (5 µL) were screened on a 1% agarose/TBE gel using a HyperLadder 50 bp standard (generally 5 ng μ L⁻¹) (Bioline). Replicate reactions were purified using the QIAquick Purification Kit (Qiagen), and quantified using the Qubit fluorescent broad range dsDNA assay (Life Technologies). Amplicons were pooled in equimolar ratios and shipped to the University of Illinois W.M. Keck Center for Comparative and Functional Genomics for preparation and sequencing using 250 bp paired-end MiSeq (Illumina), as detailed previously (Kozich et al., 2013). Sequence analyses were conducted using mothur v.3.3.3 (Schloss et al., 2009) and included assembly of the paired ends, amplicon size selection (253 bp median size) and alignment to the SSU rRNA gene. Chimera detection was also conducted via UCHIME (Edgar et al., 2011) in mothur, and chimeric sequences were subsequently removed. Taxonomic classification of sequences was conducted using the k-nearest neighbor algorithm with the SILVA SSU Ref database (release 119). Sequence data are available at NCBI's SRA under accession PRJNA 312300.

Symbiodinium taxonomy was resolved using a portion of the chloroplast 23S rRNA gene using primers 23S1 (5'-GGCTGTAACTATAACGGTCC-3') and 23S2 (5'-CCATCGTA TTGAACCCAGC-3') (Zhang et al., 2000). Each sample was amplified in triplicate reactions with 1.25 U of GoTaq Flexi DNA Polymerase, 5X Colorless GoTaq Flexi Buffer, 2.5 mM MgCl₂, 200 µM dNTP mix, 200 nM of each barcoded primer, and 4-9 ng of genomic template. The reactions were carried out on a Bio-Rad thermocycler with the following conditions: 95°C for 7 min followed by 39 cycles of 95°C for 45 s, 55°C for 45 s, and 72°C for 45 s. An additional extension step that was made at 72°C for 10 min concluded the reaction. Reaction products were visualized, gel excised and purified using the QIAquick gel extraction kit (Qiagen). Purified products were cloned (32 clones per colony) using the pGem-T Easy system (Promega) and shipped to Beckman Coulter Genomics for single pass Sanger sequencing. Clade affiliation was determined by aligning to reference sequences previously identified by Pochon and colleagues (Pochon et al., 2012) using ClustalW and visualizing with a weighted guide tree all within MacVector (v. 14.5.3, Apex). Sequence data are available at NCBI under accessions KX885228-KX885405.

Symbiont Cultures

Endozoicomonas montiporae LMG24815 and a member of *Symbiodinium* clade A4, *Symbiodinium* CCMP2456, were chosen as representative *P. astreoides* symbionts. *E. montiporae* LMG 24815, which was originally isolated from the coral *Montipora aequituberculata* (Yang et al., 2010) was obtained from the Belgian Co-ordinated Collections of Micro-organisms culture repositories. *E. montiporae* was grown in liquid marine broth at 23°C, and growth was quantified with cell counts after staining with 2-(4-amidinophenyl)-1H-indole-6-carboxamidine (DAPI) (Porter and Feig, 1980). *Symbiodinium* cultures (CCMP2456, CCMP2470, CCMP3364, CCMP2466,

CCMP3408, and CCMP2556) were obtained from the National Center for Marine Algae and Microbiota (NCMA, and formerly the CCMP), Bigelow Laboratories, East Boothbay, ME. *Symbiodinium* cultures were grown in L1 media (Guillard and Hargraves, 1993) using filtered (0.2 μ m) natural seawater (Vineyard Sound, MA) as a base. All *Symbiodinium* cultures were incubated at 23°C using 140 μ mol photons m⁻² s⁻¹ on a 14:10 h light:dark cycle, and growth was tracked by monitoring the relative fluorescence of chlorophyll with a Turner fluorometer (AquaFluor[®]). *Symbiodinium* cell counts were performed under transmitted light using a counting chamber, following preservation with Lugol's solution (2%, final concentration).

Extracellular Superoxide Production by Symbiont Cultures

The ability of coral symbionts to produce extracellular superoxide in the dark and in the presence of ambient light (PAR = 45-60 μ mol photon m⁻² s⁻¹) was examined during the day $(\sim9am-1pm)$ and, in select cases, at night $(\sim8-11pm)$. These ambient light levels were likely, though not confirmed, near the compensation point (Savage et al., 2002) for photosynthesis, particularly since the Symbiodinium light microenvironment in hospite is greatly reduced compared to surface irradiance levels (Lichtenberg et al., 2016). Representative P. astreoides symbionts (Symbiodinium CCMP2456 and E. montiporae) were screened for extracellular superoxide production separately and together in a full-factorial day/night, low light/dark experiment. For experiments involving the combination of Symbiodinium CCMP2456 and E. montiporae, cultures were mixed immediately prior to analysis. In low light experiments, ambient PAR originated mainly from the laboratory lights and natural sunlight through the lab windows. For experiments conducted in the dark, lab lights were switched off, and the entire FeLume system was shielded from light using a professional-grade dark bag designed for changing photography film. In all experiments, opaque tubing was utilized in order to eliminate abiotic photo-production of superoxide in the sample tubing.

Extracellular superoxide produced by laboratory cultures was measured using a previously described MCLA/FeLume method (Diaz et al., 2013) with a few modifications (see above for general description of the FeLume system). Briefly, carrier solutions consisting of artificial seawater (Van Waasbergen et al., 1993) buffered with 20 mM phosphate (pH = 7.6) and amended with DTPA (75 μ M) were gently pumped (2 mL min⁻¹) across a sterile syringe filter placed in the FeLume's analyte line for approximately 2 min in order to generate stable baseline signals (<4% coefficient of variation). Next, the pump was stopped, the syringe filter was removed, and using a syringe, organisms were gently deposited on the filter (Symbiodinium = 0.2 or $5 \,\mu$ m, *E. montiporae* = $0.2 \,\mu$ m, *Symbiodinium* + *E. montiporae* = $0.2 \,\mu$ m). Then the organism-loaded filter was placed back inline, and the pump was restarted (2 mL min^{-1}) . In principle, superoxide produced extracellularly is entrained by the carrier solution and detected upon mixing with MCLA in the flow cell downstream of the organisms. Biological signals were collected until a stable, steady-state reading was achieved ($< \sim 4\%$ coefficient of variation) and maintained for at least 1 min. Finally, SOD was added to the carrier solution (0.8 U mL⁻¹, final) in order to confirm that superoxide was responsible for the signal observed.

Stable biological signals were averaged and corrected for background luminescence by subtracting the average initial baseline (i.e., obtained with the clean syringe filter inline, immediately before the addition of organisms and without the addition of SOD). Corrected biological signals were converted to superoxide concentrations via calibration with multi-point KO₂ standard curves under identical conditions as biological experiments, similar to the protocol described elsewhere (Diaz et al., 2013) and outlined above for the aquaria superoxide measurements. The typical detection limit [defined as three times the average standard deviation (SD) of replicate baseline signals] was $0.24 \pm 0.11 \text{ nM}$ (avg $\pm SD$). As determined from calibration experiments, superoxide half-lives varied inversely with superoxide concentration and ranged from 1.1-38.5 min, and extrapolation times were typically 64.6 \pm 10.6 s (avg \pm SD) under our experimental conditions. Biogenic superoxide concentrations were not corrected for superoxide decay and thus represent conservative estimates. As above, calibrations yielded linear curves (e.g., $R^2 > 0.92$), with a sensitivity that averaged 2.2 \pm 0.6 pM per luminescence count.

Net superoxide production rates were calculated as the product of the steady-state superoxide concentration and flow rate (final units of pmol h⁻¹). The production rate of superoxide by each replicate was normalized to the total number of cells loaded on the filter (final units of amol cell⁻¹ h⁻¹). Cell normalized production rates were converted to surface-area normalized rates (amol μ m⁻² h⁻¹). Surface area (SA) for each organism was calculated based on its typical length (*l*) and width (*w*). For Symbiodinium, SA was calculated as a sphere, $S = 4\pi r^2$, where $r = \frac{l}{2} = \frac{w}{2}$; or prolate spheroid $SA = 2\pi B^2 \left(1 + \frac{A \sin^{-1}(m)}{mB}\right)$, where $A = \frac{l}{2}, B = \frac{w}{2}$ and $m = \sqrt{1 - (\frac{B}{A})^2}$. For *E. montiporae*, SA was calculated as a rod (SA = $4\pi s^2 + 2\pi sl$), where $s = \frac{w}{2}$. Superoxide measurements were compared using unpaired two-sample *t*-tests.

RESULTS

P. astreoides Aquaria Incubations

Physiochemical measurements confirmed that corals had acclimated to indoor aquaria conditions within 2 days, based on similar temperature, pH, dissolved oxygen, conductivity, and salinity between control and coral-containing tanks (**Table 1**). The concentrations of dissolved Fe, Mn, and Cu in control aquaria were consistently higher than those in coral aquaria, and the metal concentrations decreased with time in all aquaria (**Table 2**).

Superoxide was measured on days 3-7 following the placement of corals in the indoor aquaria, During these days, superoxide in surface seawater from the three control aquaria (no corals) ranged from 0.5 to 4.4 nM (Figure 2), with an

Day [#]							
0	1	2	3	4	5	6	7
26.3 (0.3)	24.1 (0.2)	24.0 (0.2)	24.3 (0.2)	24.2 (0.1)	24.6 (0.3)	24.9 (0.2)	24.5 (0.0)
104 (1)	104 (1)	104 (1)	104 (2)	104 (1)	104 (2)	108 (1)	105 (0)
55.3 (0.1)	56.1 (0.1)	56.7 (0.4)	57.1 (0.5)	57.7 (0.6)	58.1 (0.6)	57.4 (0.6)	58.2 (0.7)
36.7 (0.0)	37.3 (0.1)	37.7 (0.4)	38.1 (0.5)	38.5 (0.5)	38.8 (0.5)	38.2 (0.4)	39.0 (0.6)
8.04 (0.02)	8.08 (0.01)	8.08 (0.02)	8.12 (0.02)	8.16 (0.02)	8.20 (0.01)	8.21 (0.01)	8.14 (0.02)
26.2 (0.2)	24.1 (0.2)	24.1 (0.2)	24.3 (0.0)	24.2 (0.1)	24.6 (0.2)	24.8 (0.2)	24.5 (0.0)
104 (1)	104 (3)	102 (1)	104 (3)	103 (1)	104 (0)	108 (0)	104 (2)
55.4 (0.1)	56.3 (0.3)	57.0 (0.5)	57.6 (0.5)	58.1 (0.5)	58.7 (0.5)	57.5 (0.4)	58.7 (0.4)
36.7 (0.0)	37.1 (0.4)	38.0 (0.4)	38.4 (0.4)	38.8 (0.4)	39.2 (0.3)	38.3 (0.3)	39.2 (0.4)
8.03 (0.02)	8.03 (0.03)	8.04 (0.02)	8.07 (0.01)	8.10 (0.01)	8.14 (0.00)	8.13 (0.01)	8.04 (0.02)
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TABLE 1 | Geochemical measurements in tanks.*

*Mean value for three replicate tanks; standard deviation in parentheses.

*Superoxide measurements made on days 3 through 7.

TABLE 2	Metal concentrations [#] in aquaria water	
TABLE 2	Metal concentrations" in aquaria water.	

Tank	Day	Fe (mg/L)	Mn (mg/L)	Cu (mg/L)
Control	1	141.0±20.1	2.3 ± 0.4	38.1 ± 4.4
	4	170.0 ± 97.4	1.0 ± 0.3	25.1 ± 2.3
	7	56.9 ± 14.4	0.7 ± 0.2	21.7 ± 2.5
Coral	1	58.0 ± 16.6	0.8±0.1	23.1±3.6
	4	47.1 ± 8.2	0.7 ± 0.1	17.1 ± 6.8
	7	31.6 ± 9.7	0.4 ± 0.0	17.6 ± 2.0

[#]Values represent mean \pm standard deviation of triplicate samples.

average value of 2.3 ± 1.1 nM (mean \pm *SD*, n = 30). These concentrations did not change significantly throughout the day. For example, in the morning (9 am) with low lights on, the average superoxide concentration in control seawater was 2.3 ± 0.9 nM. In the dark at night (9 pm), the average superoxide concentration was 2.2 ± 1.2 nM, which was statistically similar to morning concentrations (p = 0.8). Only on day 4 was a significant (p < 0.05) difference observed between morning and night superoxide concentrations within the control aquaria.

The concentration of superoxide at the top of coralcontaining aquaria ranged from 0.1 to 1.4 nM. These superoxide concentrations were typically lower (average 0.6 ± 0.3 nM; n = 30) than those observed in the control aquaria without corals (average 2.3 ± 1.1 nM), but this difference was only significant (p < 0.05) on days 4 (am and pm), 5 (am and pm), and 7 (am only) (**Figure 2**). The superoxide concentration in the seawater approximately 3 cm from the corals ranged from 0.2 to 1.7 nM, with an average value of 0.9 ± 0.4 nM. These concentrations were not significantly different than the surface water measurements, except for two time points (day 4 and day 7 at 9 pm) when the concentrations were significantly (p < 0.05) higher. Similar to control aquaria without corals, background superoxide concentrations in coral-containing aquaria were similar from day to night (p > 0.05).

Superoxide concentrations directly at the surface of the corals were significantly (p < 0.05) higher than superoxide measured in control aquaria and at background locations within the coral-containing aquaria. The average superoxide concentration at coral surfaces was 35.0 ± 8.1 nM (n = 30), and the range of superoxide concentrations at coral surfaces was 21.3 to 50.3 nM (**Figure 2**). Coral-derived superoxide concentrations did not vary significantly over the 5-day incubation period. Although average superoxide concentrations at coral surfaces were higher in the morning in the presence of low light compared to the dark night values, these differences were only significant for one time point (day 5). Even for this time point, however, dark production accounted for 79% of the superoxide produced in the presence of low light in the morning.

For one aquarium incubation, mucus was removed from the coral surface and the corals were placed back in the aquarium. Superoxide measurements on these low-mucus corals were made within an hour. The tank-surface water and water 3 cm from the coral surface were 0.2 nM (n = 1) and $0.5 \pm 0.1 \text{ nM}$ (n = 2), respectively. The average concentration of superoxide at the coral surface was $47.7 \pm 12.3 \text{ nM}$ (n = 7), which was not significantly different than corals that did not undergo mucus removal (**Figure 2**).

Identification of *P. astreoides* Symbionts

Cloning and sequencing of chloroplast 23S rRNA genes demonstrated that the experimental corals all contained *Symbiodinium* belonging to clade A (32 clones each from 6 colonies). In both the mucus and the combined mucus and tissue samples, bacterial and archaeal SSU rRNA gene sequencing revealed that communities were dominated by sequences affiliated with the *Endozoicomonas* genus of Gammaproteobacteria (*Oceanospirillales* family) (**Table 3**). Relative abundances of *Endozoicomonas* sequences were higher



(n = 3). For each coral tank, measurements were taken at one position in the surface water (**B**; n = 3), 2 positions ~3 cm away from the corals (**C**; n = 6), and 7 measurements at various locations along the coral surface (concentration modeled from data that were recorded for ~30 s; **D**; n = 21). Superoxide concentrations (mean \pm *SD*) were measured in the light during the day (9 am; PAR = ~50 μ mol photon m⁻² s⁻¹; white bars) and in the dark at night (9 pm; PAR = 0 μ mol photon m⁻² s⁻¹; black bars). Note the difference in scale for the y-axes. Significant differences (p < 0.05) between day and night superoxide concentrations indicated by an asterisk.

in the combined mucus and tissue samples ($89.88 \pm 16.68\%$) compared to just the mucus (30.92 ± 16.16 and $57.47 \pm 24.11\%$ mean abundances for days 0 and 4). *Rhodobacteraceae* were also consistent associates of all mucus and combined mucus and tissue samples, with mean abundances ranging from 3.44-4.97%.

Superoxide Production by Representative *Porites* Symbionts

Representative cultures of the most abundant *P. astreoides* symbionts, clade A4 *Symbiodinium* strain CCMP2456 and *E. montiporae* LMG 24815, were examined for their ability to produce extracellular superoxide. Both organisms produced extracellular superoxide in the morning and night in the presence and absence of low light, with steady-state superoxide concentrations ranging from 306.73 ± 59.88 to $795.26 \pm 81.55 \, \text{pM}$ (Figure 3; Table 4). Although steady state superoxide concentrations were similar between these organisms, cell-normalized rates of superoxide production were

consistently several orders of magnitude lower for *E. montiporae* than *Symbiodinium* CCMP2456, with rates ranging from 0.54 \pm 0.06 to 1.00 \pm 0.29 amol cell⁻¹ h⁻¹ and 887.8 \pm 202.7 to 1353.1 \pm 531.5 amol cell⁻¹ h⁻¹, respectively. When normalized to cell surface area, superoxide production rates by *Symbiodinium* CCMP2456 (4.22 \pm 2.06 to 5.32 \pm 2.09 amol μ m⁻² h⁻¹) were still considerably higher than those by *E. montiporae* (0.06 \pm 0.01 to 0.11 \pm 0.03 amol μ m⁻² h⁻¹; **Table 4**).

In addition to monocultures, extracellular superoxide production was measured in combined samples of *Symbiodinium* CCMP2456 and *E. montiporae*. These were normalized to the total number of *Symbiodinium* CCMP2456 and *E. montiporae* cells and ranged from 0.94 \pm 0.14 to 1.24 \pm 0.71 amol cell⁻¹ h⁻¹. Expected levels of superoxide production by combined cultures were calculated by extrapolating individual superoxide production rates by *Symbiodinium* CCMP2456 and *E. montiporae* under the same temperature conditions

Major taxa	Mucus, I	0ay 0 (<i>n</i> = 12)	Mucus,	Day 4 (<i>n</i> = 10)	Mucus and Tissue ($n = 6$)	
	Range (%)	Mean (%) (SD)	Range (%)	Mean (%) (SD)	Range (%)	Mean (%) (SD)
Bacteroidetes	1.59–5.74	3.16 (1.39)	0.61–5.49	2.57 (1.40)	0.51-6.76	2.22 (2.24)
Synechococcus	0.11-0.52	0.22 (0.15)	0.03–0.38	0.14 (0.11)	0-0.03	0.01 (0.01)
Rhodobacteraceae	1.38-11.21	4.97 (3.00)	1.91-15.92	4.46 (4.10)	0.81-10.43	3.44 (3.61)
SAR11 clade	1.06-5.80	2.16 (1.26)	0.31-0.95	0.59 (0.22)	0.02-0.17	0.10 (0.05)
Sphingomonadales	1.63-7.10	3.95 (1.47)	1.25-6.34	3.72 (1.64)	0.02-0.53	0.14 (0.19)
Rhizobiales	1.47-21.07	13.19 (6.61)	0.61-5.36	2.02 (1.47)	0.17-1.90	0.67 (0.63)
Other Alphaproteobacteria	2.38-7.09	3.45 (1.29)	0.9–8.83	2.86 (2.35)	0.99-4.84	2.06 (1.45)
Endozoicomonas	3.89-55.84	30.92 (16.12)	4.04-84.21	57.47 (24.11)	47.88-93.35	80.88 (16.68)
Alteromonadaceae	1.05-15.42	5.97 (3.96)	0.73-12.16	2.67 (3.40)	0.05-0.95	0.34 (0.32)
Vibrionaceae	0.15-11.82	3.22 (3.22)	0.02-7.34	0.85 (2.28)	0-0.13	0.04 (0.05)
Pseudomonas	0.57-11.39	3.43 (3.48)	0.45-10.07	3.04 (2.73)	n.r.	n.r.
Pseudoalteromonadaceae	0.64-26.67	10.07 (8.61)	0.00-2.25	0.41 (0.67)	n.r.	n.r.
Other Gammaproteobacteria	2.48-13.64	6.08 (3.61)	0.97-8.45	3.04 (2.10)	0.44-4.28	1.47 (1.44)
Oxalobacteraceae	0.30-3.88	1.67 (0.96)	0.50-30.23	11.62 (8.71)	0-0.09	0.03 (0.03)
Other Betaproteobacteria	0.60-5.66	2.87 (1.83)	0.26–3.20	1.64 (0.93)	0.09-1.70	0.42 (0.63)
Other Proteobacteria	0.51-7.27	1.71 (1.82)	0.15-1.76	0.76 (0.58)	0.73-4.42	1.84 (1.43)
Planctomycetes	0.01-0.77	0.39 (0.25)	0.02-1.67	0.42 (0.50)	0.48-9.09	2.66 (3.21)
Other Bacteria	0.81-5.02	2.30 (1.33)	0.43-4.08	1.56 (1.28)	1.48-6.90	3.60 (2.04)
Archaea	0.10-0.47	0.28 (0.12)	0.01–0.53	0.15 (0.15)	0.02-0.16	0.09 (0.06)
n.r. = not represented.						
shading key:	M M	ean 0–5% ean 5–10%		Mean 10–20% Mean >20%		

TABLE 3 | Percent composition of major bacterial and archaeal taxa represented in coral colonies (SD = standard deviation).

and at the same time of day. For example, mixed cultures contained approximately 4.4×10^4 and 8.6×10^7 cells of *Symbiodinium* CCMP2456 and *E. montiporae*, respectively, and predicted superoxide production rates for the mixed consortia ranged from 1.10 ± 0.10 to 1.81 ± 0.60 amol cell⁻¹ h⁻¹ (**Figure 3**). Based on *t*-tests conducted on the difference between measured and calculated production rates under the same light/time conditions, combining the *Symbiodinium* and *E. montiporae* cultures did not have a significant effect on extracellular superoxide production (p > 0.05).

The potential effect of low light and time of day on extracellular superoxide production by *Symbiodinium* CCMP2456, *E. montiporae*, and combined cultures were examined. Although the presence of ambient low light levels (PAR = $45-60 \mu$ mol photon m⁻² s⁻¹) generally increased average superoxide production rates by *Symbiodinium* CCMP2456, *E. montiporae*, and combined cultures (**Figure 3**), these increases were not significant (p > 0.05) (**Figure 3**; **Table 4**). In fact, the only significant difference was a significant decrease in superoxide production by *E. montiporae* under low light conditions (0.54 ± 0.06 amol cell⁻¹ h⁻¹) as opposed to in the dark (0.81 ± 0.04 amol cell⁻¹ h⁻¹) during the evening (p = 0.03; **Figure 3**).

Other Symbiodinium spp. Incubations

To obtain a broader sense of extracellular superoxide production by a variety of *Symbiodinium* strains, we also examined cultures representative of phylogenic clades B2, C1, D, and D1. A wide range of extracellular superoxide production rates was measured for these five *Symbiodinium* strains, from 55.0 \pm 2.8 to 1265.4 \pm 294.1 amol cell⁻¹ h⁻¹ (**Figure 4**). The lowest production rates were measured for *Symbiodinium* CCMP3364 (clade B2) and the highest from *Symbiodinium* CCMP2466 and CCMP3408 (clades C1 and D1, respectively) (**Figure 4**). For all cultures, surface-area normalized rates ranged from 0.12 \pm 0.01 to 4.17 \pm 0.97 amol μ m⁻² h⁻¹ (**Table 4**). Due to differences in initial cell densities, these inter-strain comparisons of superoxide production rates must be interpreted with caution, however (see Discussion below).

Extracellular superoxide production rates for each *Symbiodinium* strain were generally higher in the presence of low light than in the dark, but this difference was only statistically significant for *Symbiodinium* CCMP3364 (clade B2), in which the presence of low light stimulated extracellular superoxide production by 50% (p = 0.04; **Table 4**). On the other hand, *Symbiodinium* CCMP2466 (clade C1) produced significantly more superoxide in the dark (1183.8 ± 26.1 amol cell⁻¹ h⁻¹) than in the presence of low light (830.4 ± 75.4 amol cell⁻¹ h⁻¹; p = 0.03; **Figure 4**).

DISCUSSION

ROS play a critical yet enigmatic role in coral physiology and health, from bleaching to potential immune defense. Thus,



an improved understanding of the factors that regulate ROS homeostasis in corals is necessary in order to understand the health and functioning of coral reef ecosystems. Here, we examined extracellular superoxide production by aquaria-hosted *P. astreoides* colonies. In these experiments, superoxide was measured in control (coral-free) aquaria, at various background locations within coral-containing aquaria, and at coral surfaces. Background concentrations of superoxide in the experimental and coral-free control aquaria (0.2–1.4 nM; **Figure 2**) were similar to concentrations of superoxide previously observed in productive brackish and marine waters (Rusak et al., 2011; Zhang et al., 2016). Superoxide concentrations at coral

surfaces (29.0 \pm 7.6 to 41.9 \pm 0.9 nM) were 27 to 96 times higher than background seawater concentrations in the same aquarium, which was a significant difference (p < 0.05), indicating that corals are a source of superoxide. Consistent with this point source production and with the short lifetime of superoxide in these waters (average half-life in DTPA treated reef water = ~1.3 min), coral-derived superoxide was limited to the corals' immediate surroundings. For example, within coral-containing aquaria, superoxide concentrations significantly decreased at a distance of only 3 cm from the coral. Superoxide concentrations 3 cm away from the coral surface and at the aquarium surface were similar with the exception of two time

Organism	Cell length (μm)	Cell width (μm)	Cell surface area (μm ²)	Time of day	Light condition	Steady-state concentration (pM)	Production rate (pmol hr ⁻¹)	Cell normalized [*] production rate (amol cell ⁻¹ hr ⁻¹)	Surface area normalized production rate [#] (amol μ m ⁻² h ⁻¹)
DIEL EXPERIMENTS									
Symbiodinium	9	9	254.5 ^a	AM	Light	462.92 ± 49.80	55.55 ± 5.98	1328.5 ± 150.11	5.22 ± 0.59
CCMP2456 (clade A4)				AM	Dark	314.98 ± 89.08	37.80 ± 10.69	887.81 ± 202.67	3.49 ± 0.80
				PM	Light	408.29 ± 78.15	48.99 ± 9.38	1353.14 ± 531.52	5.32 ± 2.09
				PM	Dark	306.73 ± 59.88	36.81 ± 7.19	1072.59 ± 525.82	4.22 ± 2.07
Endozoicomonas	2	1	9.4 ^b	AM	Light	596.75 ± 42.76	71.61 ± 5.13	1.00 ± 0.29	0.11 ± 0.03
montiporae LMG24815				AM	Dark	382.65 ± 70.80	45.92 ± 8.49	0.69 ± 0.31	0.07 ± 0.03
				PM	Light	411.27 ± 59.24	49.35 ± 7.11	0.54 ± 0.06	0.06 ± 0.01
				PM	Dark	615.77 ± 63.48	73.89 ± 7.62	0.81 ± 0.04	0.09 ± 0.00
Symbiodinium			-	AM	Light	633.74 ± 209.67	76.05 ± 25.16	1.24 ± 0.71	_
CCMP2456 (clade				AM	Dark	614.87 ± 205.84	73.78 ± 24.70	1.20 ± 0.69	-
A4) + E. montiporae $IMG24815$				PM	Light	795.26 ± 81.55	95.43 ± 9.79	1.05 ± 0.05	-
EWI024013				PM	Dark	722.31 ± 154.10	86.68 ± 18.49	0.94 ± 0.14	-
SYMBIODINIUM STRA	AINS								
Symbiodinium	9	9	254.5 ^a	AM	Light	517.72 ± 90.05	62.13 ± 10.81	740.08 ± 35.38	2.91 ± 0.14
CCMP2456 (clade A4)				AM	Dark	349.20 ± 36.15	41.90 ± 4.34	517.70 ± 77.92	2.03 ± 0.31
Symbiodinium	13.5	11.5	464.4 ^c	AM	Light	864.93 ± 74.47	103.79 ± 8.94	84.38 ± 7.26	0.18 ± 0.01
CCMP3364 (clade B2)				AM	Dark	564.06 ± 28.69	67.69 ± 3.44	55.03 ± 2.80	0.12 ± 0.01
Symbiodinium	10	10	314.2 ^a	AM	Light	791.01 ± 47.18	94.92 ± 5.66	830.37 ± 75.38	2.64 ± 0.24
CCMP2466 (clade C1)				AM	Dark	1139.34 ± 94.33	136.72 ± 11.32	1183.85 ± 26.08	3.77 ± 0.08
Symbiodinium	10.5	9.5	303.6 ^c	AM	Light	861.33 ± 218.12	103.36 ± 26.17	1265.40 ± 294.16	4.17 ± 0.97
CCMP3408 (clade D1)				AM	Dark	760.61 ± 118.45	91.27 ± 14.21	1128.17 ± 182.37	3.71 ± 0.60
Symbiodinium	10	10	314.2 ^a	AM	Light	274.39 ± 57.20	32.93 ± 6.86	393.36 ± 165.23	1.25 ± 0.52
CCMP2556 (clade D) [#]				AM	Dark	213.78 ± 53.33	25.65 ± 6.40	310.50 ± 138.30	0.99 ± 0.44

TABLE 4 Summary of superoxide measurements as the mean :	SEM for biological replicates (n =	= 3) analyzed during mid-exponential growth.
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^{*}Cell numbers were $\sim 10^5$ to 10^6 cells for Symbiodinium spp. and 5×10^7 to 1×10^8 cells for E. montiporae.

[#]Surface area calculated as described in methods for ^asphere, ^brod, or ^cprolate spheroid.

points (day 4 and 7 at 9 pm), when superoxide in surface seawater was higher (p < 0.05; Figure 2). Further, compared to control aquaria that did not contain corals, concentrations of superoxide in seawater at the surface of coral-containing aquaria were typically 52-90% lower. However, this result was only statistically supported (p < 0.05) in half the measurements, indicating that the presence of corals did not have a significant influence on superoxide concentrations at the surface of the aquaria. The degradation of coral-derived superoxide is, at least partly, due to the reaction between superoxide and redox active metals, such as iron (Fe) and copper (Cu) (Voelker et al., 2000; Rose, 2012), which were present at relatively high levels in the aquaria seawater (Table 2). The higher metal concentrations in control aquaria compared to coral-containing tanks likely reflects direct attenuation and potentially uptake of metals by the corals, because reaction with superoxide would not lead to a decrease in total metal concentrations. Similarly, the decrease in metal concentrations over time in all aquaria, including the coral-free control aquaria, is likely explained by adsorption of the metals to the glass walls of the aquaria.

The superoxide concentrations measured at P. astreoides surfaces were not significantly different over the course of the 5-day incubation and thus represent consistent concentrations controlled by simultaneous production and decay reactions at the coral surface (Figure 2). In addition, variability in extracellular superoxide production as a function of time of day and presence of low light levels was not observed for P. astreoides (Figure 2). Although coral colonies produced higher average concentrations of superoxide in the presence of low light in the morning on most days, this trend was not statistically significant (p > 0.05), except on day 5 (Figure 2). This result may be due to the low light levels used (PAR = $\sim 100 \,\mu$ mol photon $m^{-2} s^{-1}$) that may not have been sufficient for significant photosynthetic activity. Nevertheless, this result rules out abiotic photo-oxidation mechanisms as a major source of the superoxide observed, further indicating that the coral itself is producing extracellular superoxide. Moreover, the production



of extracellular superoxide by *P. astreoides* in the dark indicates that this pathway is independent of photosynthesis. In addition to the results from *P. astreoides* described herein, dark production of both extracellular superoxide and hydrogen peroxide has been shown previously for aquaria-hosted specimens of the coral *Stylophora pistillata* (Saragosti et al., 2010; Armoza-Zvuloni and Shaked, 2014). Thus, non-photosynthetic sources of extracellular ROS clearly exist in multiple coral species.

The superoxide concentrations measured at the surface of P. astreoides in this study are among the highest concentrations of superoxide reported for marine systems. For example, coastal and open ocean environments typically exhibit superoxide concentrations in the pM range (Rose, 2012). However, concentrations up to $\sim 1 \text{ nM}$ were previously reported in the surface waters of the Great Barrier Reef (Rose et al., 2010), which is similar to background levels of superoxide in seawater from this study (0.2–1.4 nM). Furthermore, up to \sim 33 nM superoxide was reported in association with the deep chlorophyll maximum in one site off the coast of New Zealand, indicating that high biological activity can substantially elevate seawater superoxide concentrations (Rusak et al., 2011). Indeed, in a previous aquaria study, the coral S. pistillata increased seawater superoxide concentrations from $\sim 2 \text{ nM}$ (background level) to $\sim 20 \text{ nM}$ in the dark and \sim 35 nM in the light (Saragosti et al., 2010), which is similar but slightly lower than the average concentrations reported herein for *P. astreoides* (max 41.9 \pm 0.9 nM). Finally, perhaps the highest producer of extracellular superoxide is the ichthytoxic raphidophyte Chatonella marina, which is capable of producing up to 140 nM of superoxide at bloom-level cell densities (Garg et al., 2007). Thus, while the concentrations of P. astreoides-derived superoxide are relatively high, they are not entirely unprecedented.

Production of superoxide at the surface of P. astreoides may also result in substantial generation of hydrogen peroxide. Hydrogen peroxide concentrations of ~500 nM have been reported previously at the surface of Porites sp. (Shaked and Armoza-Zvuloni, 2013). Whether this hydrogen peroxide derives from direct production at the coral surface, through the reduction of external coral-derived superoxide, and/or through diffusion of internal hydrogen peroxide into the external environment remains unclear. Regardless, external hydrogen peroxide may serve a beneficial role in corals, by acting in prey acquisition and the defense against pathogens regulated by physical and chemical stimuli (Armoza-Zvuloni et al., 2016). Corals also have a strong ability to degrade external hydrogen peroxide (Armoza-Zvuloni and Shaked, 2014) and superoxide (Saragosti et al., 2010), which may presumably be up-regulated if external ROS levels become hazardous to the coral and allow for tight regulation of ROS levels in the coral vicinity.

Superoxide is produced at multiple locations and by different mechanisms within the coral holobiont, including photosynthetic production via the reduction of endogenous O₂ within Symbiodinium cells (Figure 1). However, as mentioned above (see Introduction), this internal superoxide is unlikely to be detected at coral surfaces. Rather, the most likely sources of external superoxide include epibiotic microbes residing in the coral mucus layer, as well as the coral host's epithelial cells. To test the potential contribution of mucus-associated microbes to external superoxide levels, select P. astreoides colonies were dripped to remove the majority of their mucus layer. Mucus-dripped corals produced similar concentrations of superoxide as non-dripped colonies, suggesting that extracellular superoxide fluxes are likely attributable, at least in large part, to the coral host, rather than epibionts. Similarly, previous studies have recognized a prominent role for the coral host in the production of superoxide (Dykens et al., 1992; Nii and Muscatine, 1997; Saragosti et al., 2010) and hydrogen peroxide (Mydlarz and Jacobs, 2006; Shaked and Armoza-Zvuloni, 2013; Armoza-Zvuloni and Shaked, 2014; Armoza-Zvuloni et al., 2016).

Endosymbionts including Symbiodinium and tissue-hosted bacteria can be ruled out as contributors to extracellular superoxide observed at the coral surface, but extracellular production by these organisms can contribute to internal (tissue) ROS concentrations, which contribute to internal redox homeostasis and may lead to oxidative stress within the coral host under suboptimal conditions. Similar to previous observations of Symbiodinium (CCMP2466, CCMP831) (Saragosti et al., 2010) and heterotrophic bacteria (Learman et al., 2011; Diaz et al., 2013), we observed dark production of extracellular superoxide by non-stressed Symbiodinium (CCMP2456, CCMP3364, CCMP2466, CCMP3408, CCMP2556) (Figure 4; Table 4), as well as non-stressed E. montiporae (Figure 3; Table 4), a widespread and abundant bacterial symbiont of P. astreoides and other corals. Significant strain-specific differences in cell-normalized extracellular superoxide production by Symbiodinium were observed. However, these results should be interpreted with caution. Cell-normalized extracellular superoxide production has been shown to vary inversely with

cell density (i.e., the number of cells loaded for analysis onto the FeLume) (Marshall et al., 2005; Hansel et al., 2016). Thus, even though production rates are cell-normalized, accurate comparisons between Symbiodinium strains can only be made at the same cell density. Because inter-strain comparisons were not the goal of this study, we did not control for the possible effect of cell density between different strains. In other words, the number of cells analyzed was not consistent among the different strains, which prevents our ability to discern whether inter-strain differences were a result of differences in our analytical procedure (i.e., cell load on the FeLume) or actual physiological variability between strains. Consistent with cell density playing a dominant inverse role in extracellular superoxide production, the Symbiodinium strain with the lowest superoxide production rate (CCMP3364; Clade B2) was analyzed at a ~10-fold higher cell density compared to all the other Symbiodinium cultures (Figure 4). If cell density is inversely related to extracellular superoxide production by Symbiodinium in coral tissue, then corals with higher Symbiodinium abundances may be less susceptible to superoxide accumulation and oxidative stress, which runs contrary to the previous finding that higher densities of Symbiodinium may increase the vulnerability of corals to bleaching (Cunning and Baker, 2013).

Within a single microbial strain (and also for the mixed cultures), identical cell loads were used to generate superoxide measurements in the dark and under low light conditions. Low illumination (PAR = $45-60 \,\mu$ mol photon m⁻² s⁻¹) did not significantly increase extracellular superoxide production by E. montiporae or any Symbiodinium strain in our study, with the exception of Symbiodinium CCMP3364, in which light enhanced extracellular superoxide production by $\sim 50\%$ (Figure 4). In a previous study that examined extracellular superoxide production by two Symbiodinium strains in the presence and absence of heat stress, light significantly stimulated extracellular superoxide production by Symbiodinium CCMP2466 (both temperature conditions) and Symbiodinium CCMP831 (heat stress only) (Saragosti et al., 2010). However, we also examined Symbiodinium CCMP2466 and found the opposite effect: Low illumination significantly attenuated extracellular superoxide production by \sim 30% in this organism. This discrepancy is likely due to the low light levels utilized here (45-60 µmol photon $m^{-2} s^{-1}$) and the higher levels (300 µmol photon $m^{-2} s^{-1}$) in the previous study (Saragosti et al., 2010). Interestingly, low light levels also significantly decreased extracellular superoxide production by \sim 30% in *E. montiporae*, although this only occurred at night and not during the day (Figure 3). Overall, our results point to dark pathways of extracellular superoxide production by multiple Symbiodinium strains and the coral bacterium E. montiporae under non-stressful conditions. Furthermore, extracellular superoxide production by Symbiodinium clearly exhibits strain-specific differences in terms of each strain's responsiveness to even low light levels.

In addition to monocultures, we examined extracellular superoxide production by *Symbiodinium* CCMP2456 (clade A4) and *E. montiporae* in the presence of each other under dark

and low light conditions at different times of day (**Figure 3**). Although it is unknown whether and to what degree these symbionts may interact within the coral holobiont, our goal was to assess extracellular superoxide production at a level of biological complexity that may be more representative of natural conditions within a coral than microbial monocultures would be. These two organisms did not have an effect on each other, however, both in terms of overall extracellular superoxide production rates and the lack of response of these rates to low light and time of day (**Figure 3**). These results suggest that *Symbiodinium* and *Endozoicomonas* may not interact within corals in any way that would affect internal redox homeostasis.

The lack of day-to-night differences in dark extracellular superoxide production by P. astreoides and coral symbionts not only excludes photosynthetic mechanisms of external superoxide generation but also rules out circadian control under these conditions. A variety of physiological processes in corals are thought to be regulated by circadian rhythms, such as calcification, reproduction, and tentacle expansion (for nighttime feeding) (Sorek et al., 2014). A recent study suggests that external release of hydrogen peroxide by S. pistillata is associated with zooplankton grazing (Armoza-Zvuloni et al., 2016). However, the day-night uniformity of superoxide concentrations produced by *P. astreoides* here suggests that external superoxide production is not associated with circadian processes such as tentacle expansion and therefore may not play a role in feeding, at least in these corals under the conditions of this study.

Overall, our data indicate that a dark pathway of extracellular superoxide production exists within healthy, pigmented P. astreoides and representative coral symbionts, illustrating a lack of dependence of this process on photosynthesis and stress. Therefore, a constant dark pool of superoxide, which is unrelated to stress, is present at the coral surface and within coral tissues, since extracellular superoxide production by non-stressed endosymbionts contributes to internal ROS levels. We speculate that constitutive extracellular sources of dark superoxide may be beneficial to coral health. For example, putative superoxide-generating NADPH oxidases have been implicated in coral thermotolerance (Dixon et al., 2015) and in the resistance of corals to pathogenic white band disease (Libro et al., 2013). Furthermore, superoxide may serve as a defensive mechanism against heat-dependent pathogenic bleaching by the bacterium Vibrio shiloi, which expresses an extracellular SOD as a virulence factor (Banin et al., 2003). Extracellular superoxide may also serve a role in coral growth, as proposed for other organisms including bacteria, phytoplankton, and harmful bloom-forming algae, where extracellular superoxide production is a cellular mechanism for controlling cell proliferation (Oda et al., 1995; Saran, 2003; Buetler et al., 2004; Marshall et al., 2005; Hansel et al., 2016).

The findings of this study do not discount photosyntheticallyderived ROS production within *Symbiodinium* as an important source of ROS inside coral tissue (Weis, 2008) but instead improve our understanding of ROS homeostasis in corals by supporting a separate pathway of superoxide generation. Furthermore, our results point to potentially novel aspects of ROS in coral physiology and health, which may have implications for the future of coral reefs. Thus, the physiological role, controls, and consequences of external, dark superoxide production by corals should be investigated further.

AUTHOR CONTRIBUTIONS

CH and AA conceived of the study and designed the experiments. TZ, JD, CB, RP, and SM collected and analyzed the data; JD, CH, and AA interpreted the data and wrote the manuscript with input from TZ, CB, RP, and SM.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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