- 1 Title: Exposure duration modulates the response of Caribbean corals to global change stressors
- 3 **Running Title:** Exposure duration modulates coral physiology
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18 Abstract

19 Global change is threatening coral reefs, with rising temperatures leading to repeat bleaching 20 events (dysbiosis of coral hosts and their symbiotic algae) and ocean acidification reducing net 21 coral calcification. Although global-scale mass bleaching events are revealing fine-scale patterns 22 of coral resistance and resilience, traits that lead to persistence under environmental stress remain 23 elusive. Here, we conducted a 95-day controlled-laboratory experiment to investigate how 24 duration of exposure to ocean warming (28, 31°C), acidification ($pCO_2 = 400-2800 \mu atm$), and 25 their interaction influence the physiological responses of two Caribbean reef-building coral 26 species (Siderastrea siderea, Pseudodiploria strigosa) from two reef zones of the Belize 27 Mesoamerican Barrier Reef System. Every 30 days, calcification rate, total host protein and 28 carbohydrate, chlorophyll a pigment concentration, and symbiont cell density were quantified for 29 the same coral colony to characterize acclimatory responses of each genotype. Physiologies of 30 the two species were differentially affected by these stressors, with exposure duration 31 modulating responses. Siderastrea siderea was most affected by extreme pCO_2 (~2800 µatm), 32 which resulted in reduced calcification rate, symbiont density, and chlorophyll a concentration. 33 Siderastrea siderea calcification rate initially declined under extreme pCO_2 but recovered by the 34 final time point, and overall demonstrated resistance to next-century pCO_2 and temperature 35 stress. In contrast, *P. strigosa* was more negatively impacted by elevated temperature $(31^{\circ}C)$. 36 Reductions in *P. strigosa* calcification rate and total carbohydrates were consistently observed 37 over time regardless of pCO_2 treatment, with the greatest reductions observed under elevated 38 temperature. However, nearshore colonies of P. strigosa maintained calcification rates under 39 elevated temperature throughout all exposure durations, suggesting individuals from this 40 environment may be locally adapted to the warmer temperatures characterizing their natal reef 41 zone. This experiment highlights how tracking individual coral colony physiology across broad 42 exposure durations can capture acclimatory responses of corals to global change stressors.

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Keywords: coral physiology, temperature stress, ocean acidification, *Siderastrea siderea*,
 Pseudodiploria strigosa

47 **<u>1. Introduction</u>**

48 Since the Industrial Revolution, anthropogenic activities have increased the partial 49 pressure of atmospheric carbon dioxide (pCO_2) , which has warmed the atmosphere by 50 approximately 0.6°C (Pörtner et al., 2019). As atmospheric temperatures increase, so do sea 51 surface temperatures (SSTs), a trend that has continued since 1970 (Pörtner et al., 2019). 52 Increasing atmospheric pCO_2 has also caused surface ocean pH to decrease at a rate of 0.017 to 53 0.027 units per decade since the 1980s, potentially exacerbating the impacts of warming SSTs on 54 marine organisms (e.g. Pörtner et al., 2019). The resulting warming and acidification have 55 impacted organisms across the globe, as thermal niches shift and habitats rapidly change (Morley 56 et al., 2018; Parmesan and Yohe, 2003; Pörtner et al., 2019). The negative effects of global 57 change are predicted to strengthen and, under the Intergovernmental Panel on Climate Change's 58 (IPCC) most extreme emissions scenario (RCP8.5), oceans are expected to take up 5 to 7 times 59 more heat and decrease by 0.3 pH units by 2100 (Pörtner et al., 2019).

60 Coral reefs are valuable economic and ecological resources (Costanza et al., 2014) that 61 are particularly vulnerable to ocean warming and acidification. The high biodiversity of coral 62 reefs depends on the obligate symbiosis between the coral animal and its symbiotic algae of the 63 family Symbiodiniaceae (previously genus Symbiodinium; LaJeunesse et al., 2018). This 64 symbiosis is sensitive to thermal anomalies and, because tropical coral species live within 1°C of 65 their upper thermal limit, even slight increases in SST can result in bleaching (breakdown of the 66 coral-algal symbiosis) and ultimately mortality if the symbionts fail to repopulate the coral host. 67 Global coral bleaching events are occurring with increasing frequency and severity as SST 68 continues to increase (Hughes et al., 2017).

Ocean acidification is the result of increased atmospheric CO₂ dissolving into seawater,
 reducing its pH, and altering its carbonate chemistry (Doney et al., 2009; Orr et al., 2005). This

alteration of carbonate chemistry includes a decrease in carbonate ion concentration ($[CO_3^{2-}]$), 71 which reduces the saturation state of seawater with respect to aragonite (Ω_{araa}) —which can 72 73 make it more challenging for corals to build their aragonite skeletons (Doney et al., 2009; 74 Kleypas et al., 1999). Laboratory experiments have shown that ocean acidification conditions 75 projected for the end of next century can have negative (Comeau et al., 2013; Hoegh-Guldberg et 76 al., 2007; Horvath et al., 2016; Kroeker et al., 2010), neutral (Revnaud et al., 2003; Ries et al., 77 2010), and parabolic (Castillo et al., 2014) impacts on coral calcification, while field experiments 78 have yielded more negative outcomes (Albright et al., 2018; Comeau et al., 2019; Jokiel et al., 79 2008; Kline et al., 2019). The direction and magnitude of calcification responses to acidification 80 are influenced by many factors (Kornder et al., 2018), including ability to regulate calcifying 81 fluid chemistry in support of calcification (e.g. Ries, 2011), species differences (Bove et al., 2019; Okazaki et al., 2017), calcification rate under ambient conditions (Shaw et al., 2016), CO₂-82 83 induced fertilization of photosynthesis (Castillo et al., 2014), coral gender (Holcomb et al., 84 2012), experimental duration (Kline et al., 2019), co-occurring thermal stress (Anthony et al., 85 2011; Kroeker et al., 2013), and heterotrophic capacity (Cohen and Holcomb, 2009; Towle et al., 86 2015). Coral calcification in response to temperature stress is similarly complicated by a number 87 of factors, and a meta-analysis by Kornder et al. (2018) highlighted strong taxonomic variation 88 and that thermal stress is more pronounced in adults and during the summer. Similar to 89 acidification, coral calcification under elevated temperature can be mediated by heterotrophic 90 capacity (Aichelman et al., 2016; Grottoli et al., 2006; Rodolfo-Metalpa et al., 2008; Towle et 91 al., 2015). In addition to considering calcification rate, energetic reserves are critical to coral 92 health and resistance to stressors, and have been associated with bleaching susceptibility

93 (Anthony et al., 2009; Grottoli et al., 2014; Levas et al., 2018) and in determining whether a
94 bleaching event will lead to mortality (Anthony et al., 2009; Grottoli et al., 2006).

95 Fewer studies consider the combined effects of temperature and acidification stress and, 96 similar to studies investigating the effects of independent stressors, such experiments have 97 produced variable results. Several studies have demonstrated that elevated temperatures have 98 stronger negative effects on the coral holobiont (combination of coral animal, algal symbiont, 99 and associated microbiota) when compared to acidification, including negative effects on 100 calcification rates (Anderson et al., 2019; Schoepf et al., 2013), larval development (Chua et al., 101 2013), and survivorship (Anderson et al., 2019). Although no studies have found synergistic 102 effects of temperature and acidification on coral calcification (i.e. combined effects are greater 103 than sum of individual effects), numerous studies have demonstrated that the effects of 104 temperature and acidification stress can be additive, in terms of impacts on coral calcification 105 (Agostini et al., 2013; Edmunds et al., 2012; Horvath et al., 2016; Kornder et al., 2018; Prada et 106 al., 2017; Rodolfo-Metalpa et al., 2011) and metabolism (Agostini et al., 2013). A more 107 complete understanding of the combined effects of global change stressors will require 108 investigations of multiple species and stressors across longer timescales with a focus on multiple 109 physiology metrics.

The effects of global change on the coral holobiont not only vary by the stressors in question, but also by species. Species differences in response to global change stressors have been observed in coral calcification rate (Bove et al., 2019; Edmunds et al., 2012; Edmunds et al., 2019; Okazaki et al., 2017) and recovery of energetic reserves through time (i.e. total soluble lipid; Levas et al., 2018). Additionally, spatial scale appears to play a role in some corals' responses to global change, with differential stress tolerance observed across populations along a

reef system (e.g. the Great Barrier Reef; Dixon et al., 2015) and across reef zones (Castillo et al., 2012; Kenkel et al., 2013a; Kenkel et al., 2013b; Kenkel and Matz, 2016). Differences in coral thermal tolerance can occur on spatial scales as small as between tidal pools in American Samoa (Bay and Palumbi, 2014; Oliver and Palumbi, 2011; Palumbi et al., 2014), illustrating that adaptation and/or acclimation to fine scale environmental differences can play a role in determining coral response to global change stressors.

122 In addition to understanding stress responses across species and spatial scales, 123 considering how duration of stress exposure affects the physiological response of the coral 124 holobiont is critical (McLachlan et al., 2020). This goal is complicated by the difficulty of 125 executing long-term laboratory (ex situ) experiments, and it is therefore relatively rare for studies 126 to track coral physiology under controlled conditions for extended periods of time. However, 127 several studies have been conducted for approximately 90 days or more (e.g., Anderson et al., 128 2019; Bove et al., 2019; Castillo et al., 2014; Comeau et al., 2019; Kline et al., 2019), and each 129 reveal nuanced patterns of stress and resilience in corals. For example, Comeau et al. (2019) 130 observed that acidification (1,050 μ atm pCO₂) caused a rapid, but species-specific, alteration of 131 calcifying fluid chemistry in four coral and two calcifying algae species throughout the entire 132 one-year duration of the experiment (Comeau et al., 2019). Additionally, by measuring S. siderea 133 growth every 30 days throughout a 90-day experiment, Castillo et al. (2014) showed that 134 calcification responses to acidification (604 μ atm pCO₂) varied substantially through time—with 135 calcification rates increasing in response to moderate acidification (604 μ atm pCO₂) between 0 136 and 60 days, and decreasing between 60 and 90 days. Finally, Levas et al. (2018) tracked corals 137 for 11 months following experimental bleaching and found interspecific differences in the timing 138 of recovery. Specifically, Porites divaricata initially catabolized lipids and decreased

139 calcification but largely recovered within 11 months, while *P. astreoides* fully recovered within 140 1.5 months after increasing feeding and symbiont nitrogen uptake (Levas et al., 2018). It is 141 therefore clear that tracking coral physiology through time can provide valuable insights into 142 how corals respond to short-, moderate-, and long-term global change stress.

143 Here, two ecologically important reef-building coral species (Siderastrea siderea and 144 Pseudodiploria strigosa) from two reef zones (forereef and nearshore) of the Belize 145 Mesoamerican Barrier Reef System (MBRS) were maintained under a fully crossed acidification 146 (ca. 400 µatm [present day], ca. 640 µatm [next century], ca. 2800 µatm [extreme]) and 147 temperature (28, 31°C) experiment for 95 days. In order to characterize the responses of these species to projected global change, holobiont physiology of each coral colony was monitored 148 149 every 30 days (exposure duration: 0-30 days = T_0-T_{30} = short-term, 30-60 days = $T_{30}-T_{60}$ = 150 moderate-term, 60-95 days = T_{60} - T_{95} = long-term), including metrics for both the coral host 151 (calcification rate, total protein, total carbohydrates) and symbiont (symbiont cell density, 152 chlorophyll *a* pigment concentration). The work presented here elucidates the impact of exposure 153 duration on corals' acclimatory response to global change stressors.

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155 **<u>2. Materials and methods</u>**

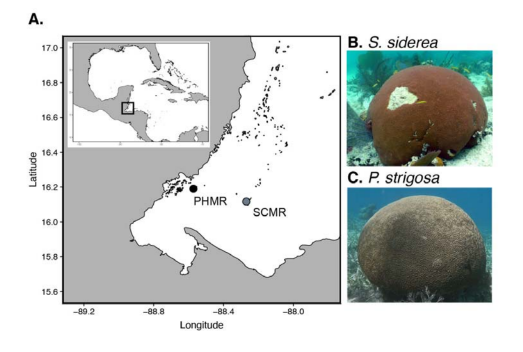
156 (2.1) Coral collection and experimental design

Data presented here are from an experiment run in parallel with an experiment published by Bove et al. (2019). Therefore, experimental design and culturing conditions are similar to those presented therein. However, data presented here pertain to a different suite of coral individuals than explored in Bove et al. (2019), and only two species (instead of four) are explored in the present study. Timing of the two experiments is also staggered by 30 days; for

162 comparison time T_0 in the present experiment corresponds to the "pre-acclimation period" in 163 Bove et al. (2019). Methods specific to this experiment are presented below. However, readers 164 are referred to Bove et al. (2019) for methods that pertain to both studies, such as those relating 165 to the control and monitoring of seawater carbonate chemistry parameters.

166 Three colonies of Siderastrea siderea and Pseudodiploria strigosa were collected from a 167 nearshore (NS; Port Honduras Marine Reserve, PHMR; 16°11'23.5314"N, 88°34'21.9360"W) 168 Sapodilla Cayes Marine Reserve, SCMR; 16°07'00.0114"N, and а forereef (FR; 169 88°15'41.1834"W) site along the southern portion of the Belize Mesoamerican Barrier Reef 170 System (MBRS) in June 2015 (N=3/species/site; Figure 1). Colonies were separated by at least 5 171 m to maximize the likelihood of obtaining genetically distinct individuals. Following collection, 172 all coral colonies (3 colonies x 2 reef zones x 2 species = 12 putative genotypes) were 173 transported to the Northeastern University (NU) Marine Science Center and fragmented into 24 174 genetically identical fragments. One FR P. strigosa colony did not survive fragmentation, 175 leaving a total of 3 genotypes for NS and FR S. siderea, 3 genotypes for NS P. strigosa, and 2 176 genotypes for FR P. strigosa genotypes (total of 6 S. siderea and 5 P. strigosa genotypes). After 177 fragmentation, coral fragments were allowed to recover for 23 days in natural flow-through 178 seawater with salinity and temperature (\pm SD) of 30.7 \pm 0.8 and 28.2 \pm 0.5°C, respectively. After 179 recovery, temperature and pCO_2 were incrementally adjusted over a period of 20 days until 180 target treatment conditions were achieved. During this time, temperatures of the elevated 181 temperature treatments were increased by 0.4°C every 3 days and pCO₂ was adjusted by 0 µatm 182 (present day), +30 µatm (end of century), and +240 µatm (extreme) every 3 days. The six 183 experimental treatments consisted of a full factorial design of two temperatures (target 28, 31°C) 184 and three pCO_2 levels (target 400, 700, 2800 μ atm). Coral fragments were distributed such that

four replicate fragments from each genotype were represented in each of the six experimental treatments. The six experimental treatments were replicated in three aquaria, for a total of 18 42L acrylic aquaria. Each aquaria was illuminated on a 10:14 h light:dark cycle with 300 μ mol photons m⁻² s⁻¹ of photosynthetically active radiation (PAR). The experimental system used natural flow-through seawater and coral fragments were fed every other day with a mixture of frozen adult *Artemia* sp. and freshly hatched *Artemia* sp.



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Figure 1. (A) Map of coral colony collection sites from the forereef (SCMR = Sapodilla Cayes
Marine Reserve) and nearshore (PHMR = Port Honduras Marine Reserve) sites of the Belize
Mesoamerican Barrier Reef System (MBRS). (B) Example of *Siderastrea siderea* colony (photo
credit: K.D. Castillo). (C) Example of *Pseudodiploria strigosa* colony (photo credit: H.E.
Aichelman).

198 Coral fragments were maintained in treatment conditions for a total of 95 days (9 August 199 2015 – 12 November 2015). Average (\pm SE) *p*CO₂ and temperature conditions throughout the 200 experimental period were: 358 (\pm 131) µatm, 28.2 (\pm 0.1) °C; 424 (\pm 40) µatm, 31.3 (\pm 0.1) °C; 201 674 (\pm 21) µatm, 28.0 (\pm 0.1) °C; 606 (\pm 36) µatm, 30.9 (\pm 0.1) °C; 2750 (\pm 161) µatm, 28.4

202 (± 0.1) °C; 2917 (± 174) µatm, 31.1 (± 0.1) °C. Seawater parameters (including temperature, 203 *p*CO₂, salinity) at each time point are shown in Figure S1 and all measured and calculated 204 seawater parameters are reported in Tables S1 and S2, respectively.

205 Approximately every 30 days (exposure duration: T_0-T_{30} days = short-term, $T_{30}-T_{60}$ days 206 = moderate-term, T_{60} - T_{95} days = long-term), a fragment of each coral colony was removed from 207 each of the six experimental conditions, flash frozen in liquid nitrogen, and stored at -80° C for 208 subsequent analysis aimed at tracking genet-level physiology through time. In the event of 209 mortality that yielded insufficient coral fragments for sampling at all time points, corals were 210 preferentially sampled at the end of the experiment (long-term exposure: T₉₅) instead of after 211 moderate-term exposure (T_{60}). As a result, sample sizes are lower for both species at T_{60} 212 compared to the other time points. Following completion of the experiment, all remaining coral 213 fragments were flash frozen in liquid nitrogen and maintained at -80°C until subsequent 214 processing, at which time fragments were airbrushed to remove host tissue and symbiont cells. 215 Tissue slurries were homogenized using a Tissue-Tearor (Dremel; Racine, WI, USA) and 216 centrifuged to pelletize the symbiont cells. Coral host tissue and symbiont fractions were 217 separated for subsequent physiological assays.

218 (2.2) Coral host physiology measurements

Coral growth rates were estimated over the course of the experiment using the buoyant weight technique (Davies, 1989). Buoyant weight measurements were obtained in triplicate for each coral fragment at each of the four time points (T_0 , T_{30} , T_{60} , T_{95}), averaged, and normalized to the coral specimens' surface area (see methods below). As in Bove et al. (2019), a subset of fragments from both species were used to confirm the relationship between buoyant weight and dry weight. These two measurements were correlated for both species (*S. siderea* $R^2 = 0.90$, p < 225 0.001; *P. strigosa* $R^2 = 0.81$, p < 0.001), indicating that change in buoyant weight should reflect 226 a proportionate change in dry weight. Equations used to calculate dry weight from buoyant 227 weight are shown below. Dry weight was converted from g to mg, corrected to surface area of 228 each fragment and to number of days in experimental treatment to calculate calcification rate 229 (mg cm⁻² day⁻¹).

230

S. siderea: Dry weight (g) = 1.95 * BW (g) + 3.60, $R^2 = 0.90$

231

P. strigosa: Dry weight (g) =
$$1.63 * BW (g) + 6.96$$
, $R^2 = 0.81$

Growing surface area was quantified in triplicate from observed live tissue in photos of coral fragments taken at each timepoint using ImageJ software (Rueden et al., 2017). The same surface area values of each coral fragment were used to normalize all host and symbiont physiology parameters within an experimental timepoint.

236 Because corals were frozen on the same day for each time point, there was no need to 237 correct for number of days in experimental treatment for physiological metrics other than 238 calcification rate. Total coral host protein content was quantified from host tissue slurry using a 239 bicinchoninic acid (BCA) protein assay. Host tissue slurry was vortexed with glass beads for 15 240 minutes and then centrifuged for 3 minutes at 4000 RPM. Next, 15 µL of the centrifuged sample 241 was added to 235 µL artificial seawater along with 250 µL of Bradford reagent. After samples 242 were mixed, absorbance was measured in a BioSpectrometer (Eppendorf, Hauppauge, NY, USA) 243 at 562 nm. Coral protein concentrations were calculated using a standard curve of bovine serum albumin ranging from 0 to 1000 μ g mL⁻¹ and normalized to living coral surface area. 244

Total host carbohydrates were quantified using the phenol-sulfuric acid method (as in Masuko et al., 2005), which measures all monosaccharides, including glucose—the major photosynthate translocated from symbiont to coral host (Burriesci et al., 2012). An aliquot of

coral host tissue was diluted to 50 μ L with artificial seawater (Instant Ocean Sea Salt), to which 150 μ L of sulfuric acid and 30 μ L of 5% phenol were added. Following a 5-minute incubation at 90°C and another 5-minute incubation at room temperature, absorbance at 490 nm was measured in a spectrophotometer (Synergy H1 Microplate Reader; BioTek Instruments; VT, USA). Carbohydrate concentrations were calculated using a standard curve of D-glucose solutions ranging from 0.039 to 2 mg mL⁻¹ and normalized to living coral surface area.

254 (2.3) Symbiodiniaceae physiology measurements

Symbiont cells were quantified using the hemocytometer method similar to Rodrigues and Grottoli (2007). After vortexing the symbiont pellet, a 1:1 Lugol's iodine and formalin solution was added for contrast and cell preservation. Triplicate 10 µL subsamples were counted on a hemocytometer using a light microscope, averaged, and normalized to slurry volume and live coral tissue surface area.

260 Symbiont photosynthetic pigments (chlorophyll a, abbreviated Chl a) were quantified 261 spectrophotometrically following the method of Marchetti et al. (2012). Briefly, 40 mL of 90% 262 acetone was added to the symbiont pellet, homogenized, then stored in the dark for 24 hours. 100 263 µL of each sample was then diluted in 7.9 mL of 90% acetone. A 10AU Field and Laboratory 264 Fluorometer (Turner Designs, San Jose, CA) was used to measure the initial concentration (R_b), 265 then 2 drops of 10% HCl was added to the sample tube, after which a second fluorometer reading was taken (R_a). Total Chl *a* content ($\mu g L^{-1}$) was calculated using the equation below, where 266 267 0.548 is a calibration constant specific to the fluorometer used, 40 mL is the volume of acetone 268 left overnight, and 80 is the dilution factor. Total Chl a was then normalized to live coral surface 269 area to get units of μ g Chl *a* cm⁻².

Chl
$$a (ug L^{-1}) = 0.548 \times (R_b - R_a) \times 40 mL \times 80$$

270 (2.4) Statistical analyses

271 All statistical analyses were completed using R version 3.5.2 (R Core Team, 2017). A 272 series of linear models (package *lmer*) were fitted for each species and each individual 273 physiology parameter using a forward model selection method. The best fit model was derived 274 by starting with the intercept-only model and then using forward-selection to incorporate 275 additional parameters, starting with the most significant parameter, until further addition of 276 parameters did not significantly improve the model fit. Additional parameters were retained in 277 the model if they were significant (p < 0.05) and produced smaller AIC values (Akaike, 1974). 278 These parameters included fixed effects of time, temperature, pCO_2 , and reef zone, which were 279 all coded as factors. Parameter interactions were only considered if those two parameters were 280 already significant and included in the model. A random effect of genotype was included in all 281 models to account for physiological variation across genotypes. Note that for calcification rate 282 data, multiple fragments of each genotype were represented at each time point. Because 283 genotype is included in the model as a random effect, multiple fragment numbers do not 284 artificially increase the sample size and instead only increase the precision of the rate 285 measurement for that colony. The linear models used for each individual physiology parameter 286 are included along with summary statistics for S. siderea and P. strigosa in Table S3. Post-hoc 287 pairwise comparisons of significant main effects were assessed using a Tukey's HSD test, 288 implemented in the *lsmeans* function with the option "adjust = tukey". Summary statistics for all 289 post-hoc comparisons are reported in Table S4.

A Principal Components Analysis (PCA) was constructed using the *FactoMineR* package (Lê et al., 2008) to assess how overall physiologies were modulated through time for each species. All physiology parameters were log-transformed, and calcification rates were x+2 log-

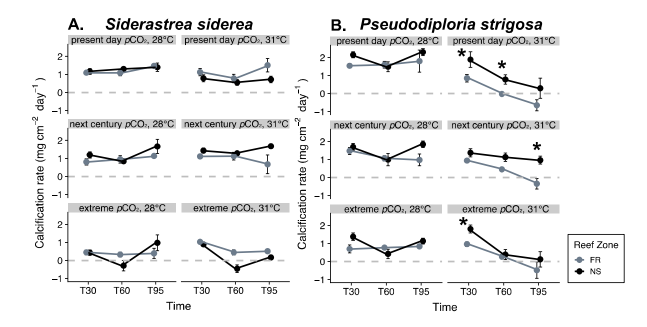
293	transformed. Only individual coral fragments for which all physiology parameters were present
294	(calcification rate, total protein, total carbohydrate, symbiont density, Chl a) were included in
295	this analysis. Significance of each factor in the PCA was assessed using the <i>adonis()</i> function in
296	the vegan package (Oksanen, 2011), which was run with 10,000 permutations using the model
297	below. Summary statistics for all Adonis tests are reported in Table S5.
298	Adonis(scores ~ reef zone * pCO_2 * temperature + genotype)
299	Correlation matrices of all host and symbiont physiology parameters for both species
300	through time were built using the <i>corrplot()</i> function with a significance threshold of $p = 0.05$.
301	The impacts of temperature and pCO_2 on all host and symbiont physiology parameters of only <i>P</i> .
302	strigosa were assessed via linear regression modeling, as no noteworthy correlations were found
303	for S. siderea. To estimate significance of the predictors and their interactions, increasingly
304	parsimonious, nested linear models (using <i>lmer()</i>) were compared with likelihood ratio tests.
305	Conditional R-squared values (accounting for both fixed and random effects) of the regressions
306	were determined using the <i>r.squaredGLMM()</i> function in the <i>MuMIn</i> package. Summary
307	statistics for all linear regressions are reported in Table S6.
308	All data and code used for the analyses presented herein can be found on the GitHub
309	repository associated with this publication
310	(https://github.com/hannahaichelman/TimeCourse_Physiology).
311	
312	<u>3. Results</u>
313	(3.1) Effects of thermal and acidification stress on S. siderea and P. strigosa calcification rates
314	Experiment duration (p < 0.001), pCO_2 (p < 0.001), and the interaction of experiment
315	duration and pCO_2 (p = 0.007), significantly influenced calcification rates of S. siderea (Figure

316 2A). Under extreme pCO_2 conditions, *S. siderea* calcification rates were significantly reduced 317 relative to both present day (Tukey p < 0.001) and next century (Tukey p = 0.017) pCO_2 318 treatments. Through time, calcification rates of *S. siderea* in extreme pCO_2 conditions declined 319 between T₃₀ and T₆₀ (Tukey p < 0.001), and also over the entire duration of the experiment (T₃₀ 320 to T₉₅; Tukey p < 0.001). Temperature was not included in the best-fit model, and therefore did 321 not have a significant effect on *S. siderea* calcification.

322 *P. strigosa* calcification rates were significantly affected by experiment duration (p < p) 323 0.001), pCO_2 (p < 0.001), temperature (p < 0.001) and reef zone (p = 0.036) (Figure 2B). 324 Calcification rates were reduced at elevated temperature (31°C) relative to control temperature 325 (Tukey p < 0.001) and nearshore corals calcified faster than forereef corals (Tukey p = 0.039). 326 When compared to the present day pCO_2 treatment, calcification rates were also reduced under 327 extreme (Tukey p < 0.001) and next century pCO₂ conditions (Tukey p = 0.017). A significant 328 interaction of temperature and experimental duration was also detected for P. strigosa 329 calcification rates (p < 0.001), with P. strigosa calcification rates decreasing between T_{30} and T_{60} 330 under both control and elevated temperatures (Tukey p < 0.05); however, these reductions were 331 no longer detectable after moderate- and long-term exposure (T_{60} and T_{95}). When considering the 332 full duration of the experiment (T_0 to T_{95}), *P. strigosa* calcification rates decreased under 333 elevated temperature, but not under control temperature (Figure 2B). Additionally, a significant 334 interaction between reef zone and temperature on *P. strigosa* calcification rate was detected (p = 335 0.031), with elevated temperatures more negatively influencing calcification of forereef corals 336 than nearshore corals (Tukey p = 0.024). Lastly, the interaction between temperature and pCO₂ 337 significantly affected *P. strigosa* calcification rates (p = 0.0009). There were no significant 338 differences in *P. strigosa* calcification rates amongst pCO_2 treatments under elevated

339 temperature treatments; however, calcification rates in control temperatures were significantly

reduced under extreme pCO_2 compared to the present day pCO_2 conditions (Tukey p < 0.001).



341

342 **Figure 2.** Siderastrea siderea (A) and Pseudodiploria strigosa (B) calcification rate (mg cm⁻²) 343 day⁻¹) at each experimental time point (short-term = T_{30} ; moderate-term = T_{60} ; long-term = T_{95}). 344 Facets represent each of the six treatments (pCO₂: present day [~400 µatm], next century [~640 345 µatm], extreme [~2800 µatm]; temperature: 28°C, 31°C) and, within a facet, data are separated by reef zone ("FR" = forereef; "NS" = nearshore). Points represent mean calcification rates since 346 347 the previous time point (i.e. T_{30} represents calcification between T_0 and T_{30}). Asterisks (*) 348 indicate significant (Tukey p < 0.05) differences in calcification rates between reef zones within 349 a time point. Error bars represent standard error. For S. siderea (A), each data point represents 350 three colonies. For *P. strigosa* (B), each forereef point represents 2 colonies and each nearshore point represents 3 colonies, except at the extreme $pCO_2/28^{\circ}C$ treatment at T₆₀ and T₉₅, where 351 352 only one forereef colony is represented due to mortality. 353

354

355 (3.2) Effects of thermal and acidification stress on *S. siderea* and *P. strigosa* host energy reserves

356 (3.2.1) *Siderastrea siderea* total protein and total carbohydrates

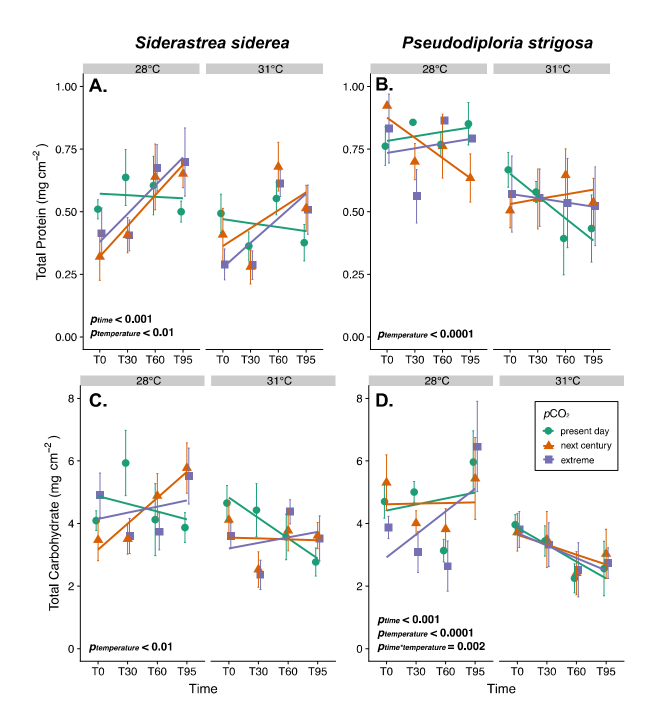
Both experiment duration (p < 0.001) and temperature (p < 0.01) had significant effects on *S. siderea* total protein content (Figure 3A), with elevated temperatures reducing total protein

359 concentrations relative to corals in control temperatures (Tukey p = 0.01). Regardless of pCO_2

and temperature treatments, *S. siderea* total protein content increased through time, with T_{95} exhibiting higher mean protein concentrations than T_0 (Tukey p = 0.03). Temperature was the only factor that influenced total carbohydrates of *S. siderea* (Figure 3C; p < 0.01), with significantly less carbohydrate at elevated temperature relative to the control temperature treatment (Tukey p = 0.003).

365 (3.2.2) *Pseudodiploria strigosa* total protein and total carbohydrates

366 For *P. strigosa*, temperature was the only factor that significantly influenced total host 367 protein (p < 0.001), with reduced protein concentrations under elevated temperatures compared 368 to control temperature treatments (Tukey p < 0.001). However, total carbohydrate concentrations 369 were significantly affected by experiment duration (p < 0.001) and temperature (p < 0.001). 370 Similar to total protein, P. strigosa total carbohydrate was reduced under elevated temperatures 371 compared to control conditions (Tukey p < 0.001). Additionally, the interaction between 372 experimental duration and temperature had a significant effect on *P. strigosa* carbohydrates (p = 373 0.002), with P. strigosa exhibiting increasing carbohydrate concentrations between T_{30} and T_{95} 374 under control temperatures (Tukey p = 0.004). In contrast, under elevated temperatures, P. 375 strigosa exhibited no change in carbohydrate concentrations over time (all Tukey p > 0.05), 376 although a trend of decreasing carbohydrate concentrations was observed from T₀ to T₉₅ (Figure 377 3D).





379 Figure 3. Host energy reserves (total protein [A,B] and total carbohydrate [C,D]) of Siderastrea 380 siderea (A,C) and Pseudodiploria strigosa (B,D) across four experimental durations (day $0 = T_0$; 381 day 30 [short-term] = T_{30} ; day 60 [moderate-term] = T_{60} ; day 95 [long-term] = T_{95}). Within each panel, results are faceted by temperature treatment (28, 31° C) and colored by pCO₂ treatment 382 (present day [~400 μ atm] = green, next century [~640 μ atm] = orange, extreme [~2800 μ atm] = 383 purple). Each point is an average of corals from nearshore and forereef reef zones with n = 4-6384 (S. siderea) and n = 3-6 (P. strigosa) distinct fragments (n = 1 / genotype). Significant factors are 385 indicated in each panel. Lines represent linear fits (using ggplot2 stat smooth() method to 386

visualize differences regardless of model) for each treatment through time, and error bars
 represent standard error at each time point.

- 390 (3.3) Effects of thermal and acidification stress on symbiont physiology of *S. siderea* and *P*.
- 391 strigosa

392 (3.3.1) Symbiont physiology of *S. siderea* (Chl *a* concentration, symbiont cell density)

393 Experiment duration (p < 0.001), temperature (p < 0.01), and pCO_2 (p = 0.04) had 394 significant effects on S. siderea symbiont cell density (Figure 4A), with cell densities decreasing 395 from T_0 to T_{95} (Tukey p < 0.001). Additionally, S. siderea cell densities were reduced under 396 elevated temperatures compared to control conditions (Tukey p = 0.017) and under extreme 397 pCO_2 compared to present day conditions (Tukey p = 0.043). The interaction between 398 experiment duration and pCO_2 also affected S. siderea symbiont cell density (p = 0.005); 399 however, within pCO_2 treatments the only significant change in cell densities was a decrease at 400 low pCO₂ between T_{30} and T_{95} (Tukey p = 0.003). Both experiment duration (p < 0.001) and 401 pCO_2 (p < 0.001) had significant effects on S. siderea Chl a concentration (Figure 4C). In 402 contrast to symbiont cell density, Chl a concentration increased from T_0 to T_{95} (Tukey p < 403 0.001). Although corals under present day and next century pCO_2 treatments exhibited similar 404 Chl a concentrations, corals under extreme pCO_2 had significantly less Chl a compared to those 405 under both present day (Tukey p = 0.002) and next century (Tukey p = 0.01) pCO₂ conditions.

406 (3.3.2) Symbiont physiology of *P. strigosa* (Chl *a* concentration and symbiont cell 407 density)

Experiment duration (p < 0.001) and temperature (p < 0.001) both had significant effects on *P. strigosa* symbiont cell density. Regardless of pCO_2 treatment, *P. strigosa* under elevated temperatures had reduced symbiont densities compared to corals in control temperatures (Tukey p < 0.001). Additionally, *P. strigosa* symbiont density was reduced at T₉₅ relative to T₀ (Tukey p

- 412 < 0.001). Temperature (p < 0.001) and pCO_2 treatments (p = 0.028) had significant effects on P.
- 413 strigosa Chl a concentration; however, experimental duration did not. Similar to symbiont
- 414 density, P. strigosa exhibited reduced Chl a concentrations under elevated temperatures
- 415 compared to control temperatures, regardless of pCO_2 treatment (Tukey p < 0.001). Additionally,
- 416 *P. strigosa* Chl *a* concentrations were reduced under extreme pCO_2 compared to present day
- 417 pCO_2 treatment regardless of temperature treatment (Tukey p = 0.02).

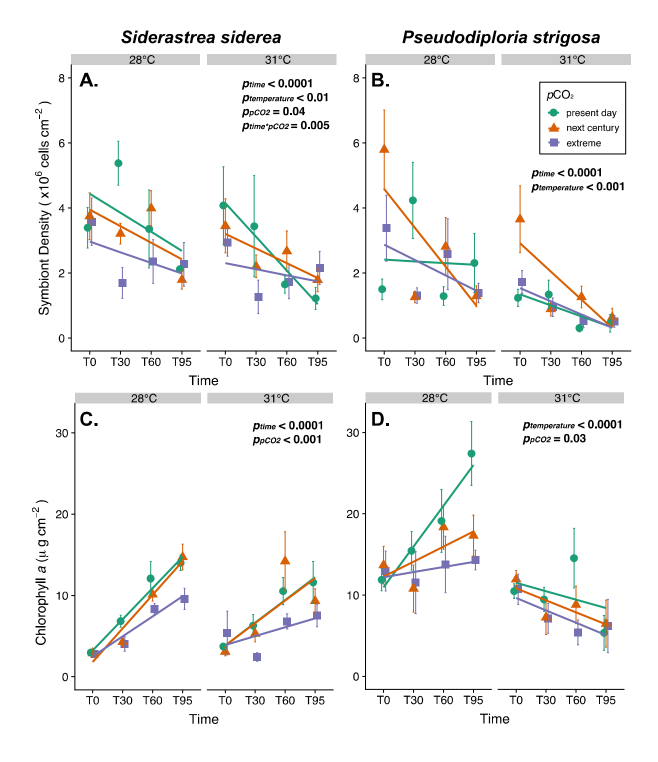


Figure 4. Symbiodiniaceae physiology (symbiont cell density [A,B] and Chl *a* concentration [C,D]) of *Siderastrea siderea* (A,C) and *Pseudodiploria strigosa* (B,D) across four experimental durations (day $0 = T_0$; day 30 [short-term] = T_{30} ; day 60 [moderate-term] = T_{60} ; day 95 [longterm] = T_{95}). Within each panel, results are faceted by temperature treatment (28°C and 31°C) and colored by *p*CO₂ treatment (present day [~400 µatm] = green, next century [~640 µatm] =

orange, extreme [~2800 μ atm] = purple). Points represent averages of n = 4-6 (*S. siderea*) and n = 3-6 (*P. strigosa*) fragments from nearshore and forereef reef zones (n = 1 / genotype). Significant factors are indicated in each panel. Lines represent linear fits (using ggplot2's stat_smooth() method to visualize differences regardless of model) for each treatment through time, and error bars are standard error at each time point.

- 430 431
- 432 (3.3) Holobiont physiology through time
- 433 (3.3.1) *Siderastrea siderea* holobiont physiology

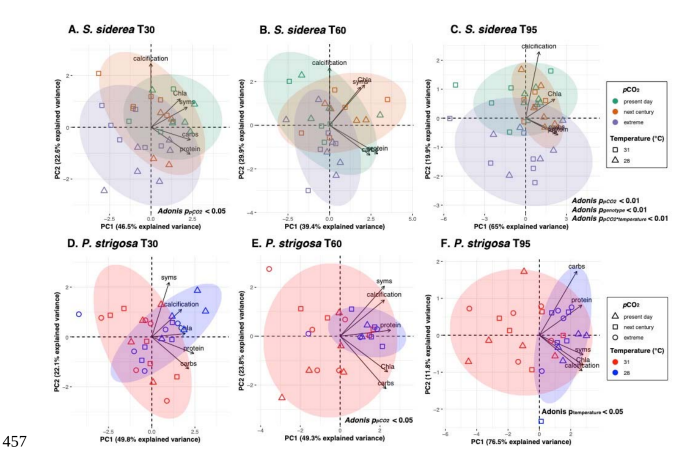
434 Overall S. siderea holobiont physiology clustered more strongly by pCO_2 than by 435 temperature (Figure 5A-C). There was a nearly significant effect of pCO_2 on holobiont 436 physiology after short-term exposure (T_{30} , Adonis p = 0.054; Figure 5A), and a significant effect 437 at T_{95} (long-term: Adonis p = 0.002; Figure 5C). At T_{95} , the interaction of pCO_2 and temperature 438 was also significant (Adonis p = 0.001; Figure 5C). Comparing PCAs in Figure 5A-C with 439 individual physiology results (Figures 3A,C and 4A,C) shows that pCO_2 significantly reduced S. siderea calcification, symbiont density, and Chl a concentration, but did not have a significant 440 441 effect on total carbohydrates or protein. These results are consistent with the PCA loadings for 442 calcification, symbiont density, and Chl a concentration discriminating between clusters of 443 fragments in extreme pCO_2 treatment and the other acidification treatments (Figure 5A-C). Reef 444 zone did not have a significant main or interactive effect on S. siderea holobiont physiology for 445 any exposure duration.

446

(3.3.2) Pseudodiploria strigosa holobiont physiology

Holobiont physiology of *P. strigosa* clustered more strongly by temperature than by pCO_2 treatment, especially after long-term exposure (T₉₅; Figure 5D-F). At T₆₀ (moderate-term), there was a significant effect of pCO_2 on holobiont physiology (Adonis p = 0.029; Figure 5E). However, at T₉₅ (long-term) the effect of pCO_2 was no longer significant, and only temperature

had a significant effect (Adonis p = 0.045; Figure 5F). Additionally, the interaction of reef zone and temperature had a marginally significant effect on holobiont physiology after long-term exposure (T₉₅; Adonis p = 0.053; Figure 5F). Comparing PCAs in Figure 5D-F with results from individual physiology parameters (Figures 3B,D and 4B,D) shows that elevated temperature had consistent negative effects on all physiology parameters.



458 Figure 5. Influence of global change stressors and exposure duration on holobiont physiology. Principal Components Analysis (PCA) of log-transformed holobiont physiology 459 data, including total carbohydrate (carbs; mg cm⁻²), total protein (protein; mg cm⁻²), symbiont 460 461 density (syms; cells cm⁻²), chlorophyll a concentration (Chla; ug cm⁻²), and calcification rate (mg $cm^{-2} day^{-1}$) for Siderastrea siderea (A-C) and Pseudodiploria strigosa (D-F). Colors represent 462 pCO_2 treatment for S. siderea (A-C: green = present day [~400 µatm], orange = next century 463 464 $[\sim 640 \text{ µatm}]$, purple = extreme $[\sim 2800 \text{ µatm}]$) and temperature treatment for *P. strigosa* (**D-F**: red = 31°C, blue = 28°C). Shapes represent temperature treatment for S. siderea (A-C: square = 465 466 31°C, triangle = 28°C) and pCO₂ treatment for P. strigosa (**D-F**: triangle = present day [~400] 467 μ [~2800 μ atm], square = next century [~640 μ atm], circle = extreme [~2800 μ atm]). Points represent an

individual coral fragment's combined physiology at each time point (\mathbf{A} , \mathbf{D} = short-term [\mathbf{T}_{30}], \mathbf{B} , \mathbf{E} = moderate-term [\mathbf{T}_{60}], \mathbf{C} , \mathbf{F} = long-term [\mathbf{T}_{95}]). Individuals were only included if they had data for each of the five parameters at each time point. The x- and y-axes indicate the variance explained (%) by the first and second principle component, respectively.

472

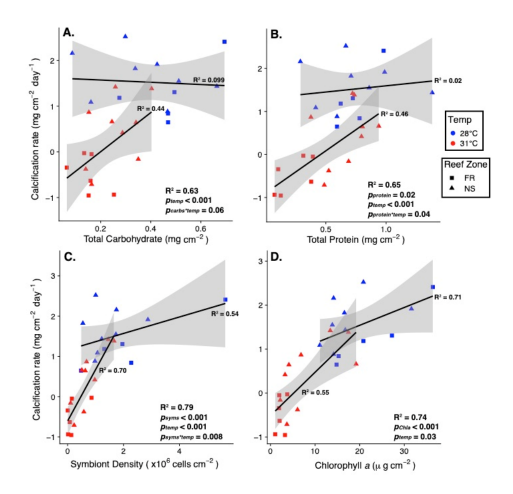
473 (3.3.3) Combined species holobiont physiology

474 Siderastrea siderea and P. strigosa had distinct holobiont physiologies at all 475 experimental durations (Adonis $p_{species} < 0.001$ for short-term [T₃₀], moderate-term [T₆₀], and 476 long-term [T₉₅]; Figure S2; Table S5). Although species had a significant main effect on 477 combined physiology through time, S. siderea and P. strigosa exhibit the most divergent 478 physiologies at T_{30} , and then converge to be entirely overlapping at T_{60} and T_{95} (Figure S2). 479 There were also significant independent effects of temperature and pCO_2 on the combined 480 physiology for both species at each time point (Adonis p < 0.05 for all time points; Figure S2, 481 Table S5).

482 (3.4) *Pseudodiploria strigosa* trait correlations

483 *Pseudodiploria strigosa* correlation matrices (Figure S3) revealed that calcification rate 484 was significantly correlated with all other physiology parameters (p < 0.05) after 95 days of 485 experimental treatment (long-term; T₉₅). As illustrated in *P. strigosa* results above, temperature 486 had a main effect on the relationship between calcification and all other predictor variables (total 487 carbohydrate and protein, symbiont density, and Chl a concentration; Figure 6). Correlations 488 with calcification rate were significantly different across temperature treatment for total protein 489 (p = 0.04) and symbiont density (p = 0.001) in *P. strigosa* at T₉₅ (long-term exposure). Under 490 elevated temperatures at T_{95} , *P. strigosa* fragments with higher total protein (p = 0.04) and 491 symbiont densities (p = 0.001) were able to maintain faster calcification rates (Figure 6 B,C). A 492 similar trend was observed for total carbohydrates; however, this relationship was not significant

493 (p = 0.06; Figure 6A). The interactive effect of temperature and the predictor variables on *P*. 494 *strigosa* calcification rate was not significant until the final observation point of the experiment 495 (T_{95} ; Figure S4). *Siderastrea siderea* correlation matrix and linear regression analyses did not 496 reveal any significant interactions with treatment.



497

498 Figure 6. Correlations of *Pseudodiploria strigosa* calcification rate with total host carbohydrate 499 (A), total host protein (B), symbiont density (C), and chlorophyll a concentration (D) after longterm exposure to experimental treatments (T_{95}) . Colors represent temperature treatment (red = 500 501 31° C, blue = 28° C) and shapes represent reef zone (square = forereef [FR], triangle = nearshore 502 [NS]). Points represent individual coral fragments. Significant factors are indicated within each 503 panel. Lines represent linear models of measured parameters within treatment through time, fit 504 using ggplot2's *stat smooth()* method with gray shading representing 95% confidence intervals for each temperature. Conditional R-squared (R^2) values (Nakagawa and Schielzeth, 2013) are 505 reported for the whole model (bottom right corner of each facet) and for each temperature (next 506 507 to the line of best fit).

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510

511 **<u>4. Discussion</u>**

512 (4.1) *Siderastrea siderea* and *P. strigosa* exhibit divergent responses to warming and 513 acidification

514 Our results demonstrate that S. siderea and P. strigosa exhibit divergent responses to two 515 co-occurring global change stressors—warming and acidification— and that these responses are 516 modulated by exposure duration (i.e. short-term = T_0-T_{30} , moderate-term = $T_{30}-T_{60}$, and long-517 term = T_{60} - T_{95}). Overall physiological performance of S. siderea was more negatively affected 518 by acidification through time (Figure 5A-C), while temperature stress had a more negative effect 519 on P. strigosa physiology through time (Figure 5D-F). Such species-specific responses to 520 temperature and acidification are not uncommon in reef-building corals. For example, when 521 testing how twelve Caribbean coral species responded to crossed temperature and acidification 522 conditions for six weeks, Okazaki et al. (2017) observed some species exhibited no growth 523 response to either temperature or acidification (including S. siderea and P. strigosa), while other, 524 more abundant Florida species (e.g. O. faveolata and P. astreoides) exhibited decreased 525 calcification rates under both stressors. In contrast to Okazaki et al. (2017), we find that S. 526 siderea and P. strigosa holobiont physiology was significantly affected by acidification and 527 elevated temperature, respectively. Although this difference may be due to the duration of 528 experimental stress (>30 days longer than Okazaki et al. (2017)), it could also be a result of 529 Okazaki et al. (2017) using less extreme treatments (30.3°C and 1300 μ atm pCO₂) than those 530 used here (31°C and 2800 µatm). It is also possible that populations of S. siderea and P. strigosa 531 in the Florida Keys, the source of corals used in the Okazaki et al. (2017) study, could be less

532 susceptible to stress than populations from the southern MBRS used here. According to the 533 climate variability hypothesis (CVH; Stevens, 1989), higher latitude populations (e.g. Florida 534 Keys) that experience more variable thermal regimes are predicted to be more phenotypically 535 flexible and exhibit a wider range of thermal tolerances compared to populations existing closer 536 to the equator (e.g. MBRS; Bozinovic et al., 2011). In a meta-analysis of Caribbean coral 537 calcification responses to acidification, elevated temperature, and the combination of the two 538 stressors, Bove et al. (2020) found similar regional differences in stress responses between corals 539 from the Florida Keys and Belize. While calcification of Florida Keys corals did not clearly 540 respond to acidification, elevated temperature, or their combination, elevated temperature 541 reduced calcification rates in corals from Belize (Bove et al., 2020). While acknowledging 542 differences in annual temperature variability, Bove et al. (2020) highlight differences in 543 experimental treatment extremes as the main driver of calcification. While consideration of 544 treatment level is critical, such population-level differences in stress tolerance have been 545 previously observed in corals, for example in Acropora millepora across 5° of latitude on the 546 Great Barrier Reef (Dixon et al., 2015). Regardless, results of the present study contribute to a 547 growing body of literature supporting S. siderea's resistance to conditions of elevated 548 temperature and acidification (Banks and Foster, 2016; Bove et al., 2019; Castillo et al., 2014; 549 Castillo et al., 2011; Davies et al., 2016).

High resistance of *S. siderea* to global change stressors was previously reported by Castillo et al. (2014), which found that only the most extreme temperature (32°C) and acidification (2553 μ atm *p*CO₂) treatments resulted in reduced calcification rates. In the context of global change scenarios projected by the IPCC, Castillo et al. (2014) concluded that *S. siderea* will be more negatively impacted by elevated temperatures over the coming century, given that

555 the IPCC's next-century acidification projections did not reduce calcification rates. The findings 556 of the present study are consistent with this previous work, as only the extreme pCO_2 treatment-557 but not next century pCO_2 treatment-reduced calcification rate in S. siderea (Figure 2A). Gene 558 expression profiling of S. siderea from the Castillo et al. (2014) coral fragments revealed that 559 thermal stress caused large-scale down regulation of gene expression, while acidification stress 560 elicited upregulation of proton transport genes (Davies et al., 2016), potentially offsetting the 561 effects of acidification at the site of calcification (e.g. Ries, 2011; Schoepf et al., 2017). These 562 findings provide further support for *S. siderea*'s ability to acclimate to acidification stress.

563 Bove et al. (2019) investigated the combined effects of similar temperature and 564 acidification treatments on four species of reef building corals: S. siderea, P. strigosa, Porites 565 astreoides, and Undaria tenuifolia. After 93 days, all species exhibited calcification declines 566 under increased pCO_2 . However, P. strigosa was the only species that exhibited reduced 567 calcification under elevated temperature, which is consistent with results present here and 568 highlights that thermal stress more negatively impacts *P. strigosa* than *S. siderea* (Figure 5D-F). 569 Bove et al. (2019) also found that S. siderea was the most resistant of the four species studied, as 570 it was able to maintain positive calcification rates even in the most extreme acidification 571 treatment (~3300 μ atm pCO₂)—findings that are also corroborated here (Figure 2A). 572 Interestingly, by quantifying net calcification rates at 30-day increments, we show that S. siderea 573 net calcification was negative under extreme pCO_2 at T_{60} and then these rates recovered by T_{95} 574 (Figure 2A). This suggests that these corals are acclimating to stressful conditions over time, 575 perhaps through transcriptome plasticity, as previously proposed by Davies et al. (2016).

576 (4.2) Stress differentially modulates coral physiology across species

577 Under thermal and acidification stress, corals can draw on energy reserves, including 578 lipids, proteins, and carbohydrates, to maintain and/or produce tissue and skeleton (Anthony et 579 al., 2009; Schoepf et al., 2013). In addition to using energetic reserves, heterotrophic feeding 580 (Aichelman et al., 2016; Drenkard et al., 2013; Edmunds, 2011; Towle et al., 2015) or enhanced 581 productivity of Symbiodiniaceae owing to CO₂ fertilization of symbiont photosynthesis (Brading 582 et al., 2011) can augment energetic resources in zooxanthellate corals. Coral energetic reserves 583 can therefore influence resistance to and recovery from stress events (Edmunds et al., 2016; 584 Grottoli et al., 2006; Grottoli et al., 2014; Schoepf et al., 2015).

585 Results of the present study show that the host energy reserves (protein, carbohydrates) of 586 S. siderea and P. strigosa responded to global change stress (warming, acidification) in different 587 ways. Between T_0 and T_{60} , *P. strigosa* exhibited reduced carbohydrates regardless of treatment, 588 indicating catabolism of this energy reserve (Figure 3D). This was followed by the restoration of 589 carbohydrates (i.e. acclimation) at control temperatures at T_{95} (Figure 3D), which likely 590 supported the positive calcification rates also observed under these conditions (Figure 6A, Figure 591 S4). Protein reserves do not show the same trend as carbohydrates through time (Figure 3B), 592 potentially owing to *P. strigosa* catabolizing carbohydrates before proteins, which has been 593 observed over shorter time scales for other scleractinian coral species (Grottoli et al., 2004). 594 However, elevated protein reserves did predict faster calcification rates in *P. strigosa* under 595 elevated temperatures, but not until the final observation point of the experiment (T_{95} ; Figure 6B, 596 Figure S4). As photosynthate translocated from symbionts is a major source of carbohydrates to 597 the coral host (Burriesci et al., 2012; Muscatine, 1990), reductions in symbiont cell density, Chl 598 a concentration, and carbohydrates of P. strigosa at elevated temperature suggest that symbionts 599 were translocating fewer resources to the host, which likely contributed to observed reductions in

calcification under these elevated temperature conditions, particularly after long-term exposure
(T₉₅, Figure 6). Total protein and carbohydrate abundances of *S. siderea*, similar to those of *P. strigosa*, declined under elevated temperatures (Figure 3A,C)—consistent with previous work
highlighting upregulation of protein catabolism pathways in *S. siderea* exposed to long-term
thermal stress (Davies et al., 2016).

605 An overall trend in reduced symbiont density and increased Chl a concentration through 606 time were observed under most pCO_2 and temperature conditions, except for *P. strigosa* under 607 elevated temperature (Figure 4). Given that both species under most treatments exhibited this 608 pattern, it cannot be ruled out that these changes in symbiont physiology were influenced by 609 other factors, including incomplete symbiont acclimation to experimental light environment 610 (Roth, 2014) and seasonal patterns in symbiont density and pigment concentration (Fitt et al., 611 2000)—which may have masked the symbiont response to thermal stress within S. siderea. In 612 contrast, P. strigosa exhibited reduced symbiont density and Chl a concentrations under elevated 613 temperature (Figure 4B,D), a pattern that is more consistent with thermally induced bleaching 614 (Brown, 1997; Fitt et al., 2001; Glynn, 1993; Warner et al., 1999; Weis, 2008) and further 615 illustrates the susceptibility of this species to thermal stress.

616 (4.3) Nearshore *P. strigosa* are more resistant than forereef conspecifics

Natal reef zone was a significant predictor of host physiology, particularly for *P. strigosa*, as nearshore corals calcified faster overall (Figure 2B). Although reef zone differences in calcification were observed for *S. siderea* (particularly through time), colonies from one reef zone did not clearly outperform colonies from the other reef zone (Figure 2A). In contrast, the reef-zone-specific calcification response of *P. strigosa* may arise from local adaptation of the host to distinct temperature regimes. On the MBRS, nearshore habitats are characterized by 623 higher maximum temperatures, greater annual temperature range, and more days above the 624 regional thermal bleaching threshold compared to forereef sites (Baumann et al., 2016). Local 625 adaptation to distinct reef zones is not uncommon in corals, and has been previously shown to 626 affect responses of corals to thermal stress. For example, P. astreoides was previously found to 627 be locally adapted to distinct thermal regimes in the Florida Keys, with inshore corals 628 demonstrating higher thermal tolerance, constitutively higher expression of specific metabolic 629 genes, and greater gene expression plasticity compared to offshore conspecifics (Kenkel et al., 630 2013a; Kenkel et al., 2013b; Kenkel and Matz, 2016). Additionally, P. strigosa is a 631 hermaphroditic broadcast spawning species, and previous work on populations from the Flower 632 Garden Banks demonstrated that larvae have short pelagic larval durations (PLD; Davies et al., 633 2017), which could facilitate local adaptation as larvae are more likely to locally recruit (Davies 634 et al., 2017; Mayorga-Adame et al., 2017). However, it is unknown if *P. strigosa* on the MBRS 635 have similarly short PLDs. We hypothesize that nearshore P. strigosa are locally adapted and/or 636 acclimated to more variable and stressful nearshore conditions, allowing maintenance of higher 637 calcification rates under thermal stress compared to their forereef counterparts.

It is important to note that responses to stress based on reef zone could be obscured by uneven sampling across reef zones, as forereef genotypes of *P. strigosa* were slightly underrepresented in the experiment (2 genotypes present vs. the standard 3) due to mortality before the experiment began. It is possible that full replication of genotype would have yielded different effects of reef zone in other *P. strigosa* physiology parameters.

Although the *S. siderea* PLD is unknown, previous work has shown high population connectivity across 2,000 km of the Brazil coast (Nunes et al., 2011). If MBRS *S. siderea* populations are similarly connected, it is possible that no genetic differences exist across

646 nearshore and forereef environments-potentially explaining the lack of reef-zone-specific 647 responses observed here. Yet this result is inconsistent with previous findings that forereef 648 colonies of MBRS S. siderea exhibited greater physiological stress than inshore colonies when 649 exposed to higher temperatures (Castillo and Helmuth, 2005), reduced skeletal extension rates 650 relative to inshore colonies over multi-decadal warming of the reef system (Castillo et al., 2012), 651 and reduced symbiont photophysiology relative to inshore colonies under higher temperatures 652 (Davies et al., 2018). However, consistent with results presented here, Bove et al. (2019) also 653 found no evidence for reef-zone differences in physiology for S. siderea, or for any of the other 654 species tested. Nevertheless, the observation that both nearshore and forereef S. siderea 655 performed well under global change stressors provides further support for resistance of this 656 species.

(4.4) Time-course experiments reveal acclimation to thermal stress in two common Caribbeancorals

659 This study contributes to a growing body of literature demonstrating the value of 660 assessing time-course physiology of corals exposed to global change stressors. Although studies 661 investigating independent effects of temperature and acidification on scleractinian corals have 662 yielded great insight into the effects of future global change on coral systems (e.g. Albright et al., 663 2018; Anthony et al., 2011; Carricart-Ganivet et al., 2012; Comeau et al., 2013; Jokiel and Coles, 664 1990; Jury et al., 2010), the combined effects of these stressors remain less explored-665 particularly in the context of how the coral stress response is modulated by the duration of those 666 stressors. By characterizing coral host and symbiont physiology of a colony through time, 667 acclimatory responses were identified in two common Caribbean reef-building coral species. The 668 results of this study provide further evidence of the species-specific nature of this acclimation.

669 For example, under extreme pCO_2 and elevated temperature, S. siderea calcification rate appears 670 to recover by the end of the experiment while *P. strigosa* calcification rate declines into negative 671 net calcification. In addition to furthering our understanding of how corals could respond to 672 projected future ocean conditions, the exposure duration component of this study suggests that 673 species will exhibit differential persistence through ephemeral stress events. It is clear that local 674 heat waves that raise SST and upwelling events that reduce pH, which already threaten coral 675 populations, may threaten coral species in different ways in the future depending on the 676 timescales of these events.

677 Acclimation is an important mechanism by which corals can withstand changing 678 environmental conditions, and transcriptome plasticity is one way by which corals can acclimate 679 to stress (Davies et al., 2016; Kenkel and Matz, 2016). A coral reciprocal transplant study 680 between reef zones in the Florida Keys demonstrated that adaptive gene expression plasticity, 681 specifically plasticity of stress response genes, was associated with reduced susceptibility to a 682 summer bleaching event (Kenkel and Matz, 2016). In addition to plasticity providing a 683 mechanism for acclimation within a generation, rapid evolutionary adaptation of corals to 684 warmer oceans has also been observed on the Great Barrier Reef (Dixon et al., 2015; Matz et al., 685 2018). However, recent widespread declines in coral abundance, diversity, and health suggest 686 that rates of intra- and trans-generational adaptation to global change stressors within most coral 687 populations are insufficient for mitigating the deleterious impacts of recent and future CO₂-688 induced global change (Thomas et al., 2018). Understanding the interplay of acclimation and 689 adaptation in scleractinian corals is therefore essential for projecting how coral reef ecosystems 690 will fare in the higher-CO₂ future (Chevin et al., 2010; Thomas et al., 2018). This study furthers 691 understanding of how exposure duration modulates coral physiology across reef zones in two

692 prevalent Caribbean reef-building species. Future studies focusing on long-term acclimation
693 capacities of corals will further elucidate mechanisms of resistance and resilience in corals'
694 response to global stressors.

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707

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716 **<u>7. References</u>**

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