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### Short-term interactive effects of increased temperatures and acidification on the calcifying macroalgae Lithothamnion crispatum and Sonderophycus capensis

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| 8        | calcifying macroalgae Lithothamnion crispatum and Sonderophycus capensis   |
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| 32       |  |
| 33       | Abstract   |
| 34       | Combined effect of acidification and sea warming derived from future conditions  |
| 35       | of Climate Change have been little investigated in marine photoautotrophs, especially on   |
| 36       | sensitive organisms such as calcifying macroalgae. The aim of this investigation was to  |
| 37       | evaluate the interactive effects of acidification and increased temperatures on the two  |
| 38       | Brazilian calcifying macroalgae Lithothamnion crispatum and Sonderophycus capensis.  |
| 39       | Both species were cultured for 48 h under: (1) current pH (7.8 $\pm$ 0.2) and temperature (18  |
| 40       | $\pm$ 2°C) during winter; (2) future pH (7.4 $\pm$ 0.2) and temperature (30 $\pm$ 2°C) during  |
| 41       | summer; (3) current temperature and future pH; and (4) future temperature and current  |

42 pH. We evaluated photosynthetic performance (measured  $\Delta F/F'_m$ ), growth rates (weight), 43 relative carbonate content, and total phenolic compounds. Our results showed similar 44 negative effects under decreased pH and increased temperatures in both species, although carbonate content S. capensis was less affected than in L. crispatum. Total phenolic 45 46 compounds measured in S. capensis showed the highest levels at potential future pH and 47 temperature conditions. Given that stress conditions associated with decreased pH and 48 increased temperatures are important inductors of an oxidative response, it is likely that 49 phenolic compounds are synthetized to fulfil an antioxidant purpose. Even though 50 physiological performance was affected in both calcifying macroalgae under the most 51 likely negative future pH and temperature conditions, their biological viability indicates 52 they may be able to thrive under coming Climate Change scenarios.

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54 **Keywords:** Climate Change; acidification; warming; macroalgae.

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#### 56 **INTRODUCTION**

57 Atmospheric carbon dioxide (CO<sub>2</sub>) concentrations have been increasing from 280 58 ppm since the pre-industrial period to above 400 pm nowadays; moreover, CO<sub>2</sub> levels are 59 expected to increase in up to 970 ppm by the end of the 21<sup>st</sup> Century (Harley et al. 2006; 60 Harley et al. 2012). Predicted CO<sub>2</sub> rise and accompanied increase in global temperatures 61 is expected to induce greater sea surface temperatures, which could fluctuate between 2 62 and 6°C depending on the strictness of the extrapolation (IPCC 2014). On the other hand, 63 approximately 25% of atmospheric CO<sub>2</sub> is absorbed by the ocean, which in reaction with 64 seawater increases the levels of carbonic acid ( $H_2CO_3$ ), subsequently diminishing the pH; 65 this process is expected to induce a 0.2-0.4 pH decline by the year 2100 (Feely et al. 66 2004). Ocean acidification produces changes in total dissolved inorganic carbon (DIC), manifested in higher concentration of carbonate ions  $(CO_3^{2-})$  and bicarbonate  $(HCO_3^{-})$ ; 67 important molecules that may affect physiological processes, mostly in marine calcifying 68 69 organisms (Martin and Hall-Spencer 2017; Orr et al. 2005; Koch et al. 2013). As signal 70 of Climate Change, monitoring of coastal marine habitats has revealed a concerning 71 increase in the incidence of mass mortalities in flora and fauna due to marine heat waves 72 as observed in Australia (Wernberg et al. 2011) or in the south Atlantic (Ferreira et al. 73 2015). Elevated temperature, even over a short period, can be lethal due to biochemical 74 damages (Gouvêa et al. 2017). In these coastal and shallow regions, high pH variability is 75 observed, eventually exposing organisms to conditions frequently predicted to the end of 76 this century considering ocean acidification (Duarte et al. 2013). Changes in ocean 77 circulation also provide localized and short variability in ocean pH due 78 upwelling/downwelling zones (Gruber et al. 2012; Findlay et al. 2013), CO<sub>2</sub> enrichment 79 from volcanic vents (Hall-Spencer et al. 2008), and as a function of the metabolism of 80 some marine communities (Anthony et al. 2011) or even in anoxic zones (Melzner et al. 2013). Therefore, extreme values of temperature and pH/pCO<sub>2</sub> present in different 81 regions of the world can accentuate or even represent nowadays ocean acidification 82 83 impacts on the biology of marine biodiversity.

Calcifying macroalgae are key ecological components of the Atlantic coast. They 84 are considered ecosystem bioengineers, as create habitat, niche for settlement and nursery 85 for other algae, invertebrates, and fish (Figueiredo et al. 2012; Riul et al. 2009; Amado-86 Filho et al. 2007). Furthermore, these organisms are major producers of carbonate 87 88 sediment in the marine environment (Basso 2012). Previous studies have shown that 89 unbalanced DIC mediated by decreased pH reduces the saturation of calcium carbonate 90 (CaCO<sub>3</sub>) in calcifying marine organisms, such calcifying macroalgae, corals, foraminifera 91 and coccolithophores; this process can affect calcification rates when producing new 92 skeleton (Basso 2012; Martin and Hall-Spencer 2017; Orr et al. 2005). CaCO<sub>3</sub> occurs in skeletons of calcifying macroalgae as calcite or aragonite; however, magnesium  $(Mg^{2+})$ 93 94 can replace calcium in calcite macroalgae, which has been observed to be more prone to dissolution under acidification than macroalgae with aragonite skeleton (Basso 2012; 95 96 Jury et al. 2010; Smith et al. 2012).

97 Calcifying macroalgae are capable of performing calcification and photosynthesis 98 at different rates depending on  $CO_2$  concentrations (Hofmann et al. 2012; Hurd et al. 99 2009; Koch et al. 2013). Despite that acidification can produce adverse effects on 100 calcification rates, changing carbonate content, high concentrations of  $CO_2$  have been 101 detected to favor photosynthesis in some cases (Hurd et al. 2009; Koch et al. 2013). For 102 instance, Semesi et al. (2009) showed that a progressive decline in pH induced a 13% 103 increase in photosynthesis rates in coralline algae, although compromising their structure due to 20% less calcification. Tropical and subtropical organisms survive near suboptimal
physiological temperature; however, Climate Change-mediated increase in sea
temperatures threatens their thermal tolerance thresholds (Latham 2008; Eggert 2012;
Harley et al. 2006). Within tolerance ranges, rising temperature can induce an increase in
photosynthesis, growth and carbonate deposition in calcifying macroalgae, but when
these exceed tolerance thresholds, it can cause bleaching and chlorophyll degradation
(Martin and Hall-Spencer 2017; Glynn 1996).

111 Drastic environmental disturbances can generate biological stress in macroalgae through the over-production of reactive oxygen species (ROS) (Bischof and Rautenberger 112 113 2012). ROS excess can produce damage by oxidizing lipids, proteins and nucleic acids; 114 however, the cell can maintain ROS within homeostatic levels through the production of antioxidant compounds (Moenne et al. 2016). In this context, phenolic compounds have 115 116 been observed to have strong ROS scavenging properties (Sáez et al. 2015; Flores-117 Molina et al. 2016). ROS-induced increase in phenolic compounds in macroalgae have 118 been observed subject to several environmental stressors, among which can be mentioned 119 herbivory, changes in temperature, salinity, irradiance and metal excess (Bischof and 120 Rautenberger 2012; Moenne et al. 2016; Sáez et al. 2015; Flores-Molina et al. 2016).

121 Records on the interaction of different stressors associated with Climate Change 122 on marine calcifying macroalgae are scarce; thus, more information on aspect such as the 123 combined effects of acidification and increased temperatures would significantly improve 124 our understanding on their current and future responses to progressive Climate Change. 125 The aim of this research was evaluate the interactive effects of a short acidification and warming on physiological aspects of the calcifying red coralline Lithothamnion 126 127 crispatum Hauck and the red aragonite Sonderophycus capensis (M) M.J. Wynne. L. L. 128 crispatum has a wide range of latitudinal distribution; considered as a dominant species 129 in the entire Atlantic coast of Brazil, its temperature tolerance ranges between 15 to 30°C 130 (Pascelli et al. 2013; Riul et al. 2009). On the other hand, S. capensis has a limited 131 distribution (southeastern Atlantic of Brazil) with tolerance ranges defined between 17 132 and 20°C (Zavialov et al. 1999). Taking into account limits of temperature tolerance and CaCO<sub>3</sub> types of skeleton in these species, it is expected that L. crispatum and S. capensis 133

have differential responses to single and combined effects of increased temperatures andacidification.

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#### 139 MATERIALS AND METHODS

140 *Collection and experimental design.* Entire individuals of two calcifying red 141 macroalgae species, the coralline L. crispatum and the aragonite-based S. capensis, were 142 collected at 10 m depth from Reserva Biologica do Arvoredo, in Santa Catarina (27° 15' S, 48° 20' W) and in Xavier Island (27° 36' S, 48° 23' O), respectively, in 143 144 Brazil. After collection, individuals were stored in plastic bags with seawater, and immediately transported inside coolers to the Laboratory of Phycology at Universidad de 145 146 Santa Catarina. Prior to experiments, the specimens were acclimated for 24 h with 147 filtered seawater (0.45  $\mu$ m) under constant aeration, temperature (15 ± 2°C) and light (50 umol photons m<sup>-2</sup>s<sup>-1</sup>). The experiment was carried out in a microcosm system over a 148 149 short-term period (48 h). The experimental system consisted in two tanks with 50 L 150 seawater that delivered seawater with a pump to 12 chambers of 1 L. Both macroalgae 151 species were randomly distributed and maintained in four conditions: (1) average current 152 pH (7.8  $\pm$  0.2) and temperature (18  $\pm$  2°C) registered in Florianopolis bay, Brazil, during 153 winter (2); future values of pH (7.4  $\pm$  0.2) and temperature (30  $\pm$  2°C) that can be 154 extrapolated for Florianopolis bay during summer by the end of the 21<sup>st</sup> Century 155 according to predictions by IPCC (2014); (3) Current temperature  $(18 \pm 2^{\circ}C)$  and future pH (7.4  $\pm$  0.2); and (4) Future temperature (30  $\pm$  2°C) and current pH (7.8  $\pm$  0.2). For 156 157 each treatment, three independent replicates were used. pH levels considered for control 158 treatments respond to local levels registered in seawater nearby macroalgae collection 159 sites. Certainly, in these locations there are several estuaries, such as Barra de Lagoa, that 160 supply freshwater and inland material that induces a pH drop in coastal waters; in this 161 context, several reports in this area show average pH levels around 7.8 (Martins-Pereira 162 2004; Dalinghaus 2016; Cancellier-Cechinel 2013). Considering the latter, acidification treatments with a 0.4 pH decrease were made according IPCC predictions upon currentlocal pH levels.

To achieve low pH conditions, bubbles of CO<sub>2</sub> were injected with a pump the 165 166 experimental seawater. The added CO<sub>2</sub> were managed by a pH controller, model pH2010 167 (Weipro, China), connected to solenoid valves that regulated the amount of gas pumped 168 into the seawater (Russell et al. 2009). Experimental temperatures were achieved by placing tanks on a homemade temperature-gradient table; this is a horizontal metal-made 169 table, which surface temperature gradually increases from 15 °C to 35 °C, from one 170 extreme to the other. In addition, the table constantly agitates to maintain temperature 171 172 homogeneity in the seawater.

Water carbonate chemistry parameters.  $HCO_3^{-2}$  and  $CO_3^{-2}$  concentrations,  $pCO_2$ , calcite and aragonite saturation (Ω), were obtained via  $CO_2Sys_v2.1xls$  program, using K1, K2, as in Millero et al. (1998). These values are summarized in Table 1. Estimations were based in measurements of pH, temperature, salinity and total alkalinity in seawater. Gran titration method was used to measure total alkalinity, using 0.0025 N hydrochloric acid (HCl; Merck) (Carmouze 1994); these measurements were taken after 48 h of experiments.

180 Photosynthetic performance. Photosynthetic measurements were performed 181 throughout the course of the experiments in all treatments, at 6, 24 and 48 h. Effective quantum yield of photosystem II (PII;  $\Delta F/F'_m$ ) was measured using a portable pulse-182 183 amplitude modulated (PAM) chlorophyll a fluorometer (DIVING PAM; Walz GmbH, 184 Effeltrich, Germany). This parameter was measured according to Schreiber and Neubauer (Schreiber and Neubauer 1990):  $\Delta F/F_m = (F_m - F_t)/F_m$ . In the latter,  $F_m$  is the maximal 185 fluorescence of light-acclimated sample induced by a saturating actinic light pulse (9000 186 umol photons m<sup>-2</sup> s<sup>-1</sup>, 800 ms), and  $F_t$  is the intrinsic steady-state fluorescence emission 187 188 in light-acclimated sample (Baker 2008).

189 *Growth rate.* To estimate the growth, the change in fresh weight (FW) biomass of 190 the species was measured, prior and after 48 h of experiments. The mean relative growth 191 was expressed as FW per day (FW  $d^{-1}$ ), and calculated according to Packard and 192 Boardman (1999): RG=  $W_t$ - $W_0/t$ .  $W_0$  is the initial weight,  $W_t$  is final weight, and t is time 193 in days (d).

194 *Relative carbonate content (RCC).* Biomass was dried at 60°C for 24 h and
195 weighted. Initial weight (DWi) was obtained and then immersed in 10% HCl (Merck) for
196 24 h to initiate CaCO3 dissolution, in order to maintain the same weight/volume ratio.
197 After the experiments, samples were dried (60°C) again and were re-weighted (DWf)
198 (Figueroa et al. 2014). RCC as a proxy of calcification was calculated to the formula: %
199 RCC ((DWi-DWf)/DWf)\*100).

200 *Content of total phenolic compound.* Due to biomass availability, total phenolic 201 compounds were only determined in S. capensis, according to Randir et al. (2002). After 202 48 h experiments, samples were immediately frozen in liquid nitrogen and stored at -203 80°C. Between 40-50 mg biomass were grounded to powder in a mortar with liquid 204 nitrogen, at 4°C chamber in the darkness. One mL of 80% methanol (Merck) and 5 ml of 205 95% methanol (Merck) were added to each sample in centrifuge tubes. Samples were 206 centrifuged at 10,000 rpm for 5 min at room temperature. One mL of the supernatant was 207 placed in a clean tube and mixed with 5 mL of distilled water, 1 mL of 95% ethanol and 208 500 µL of Folin-Ciocalteu; the mixture was incubated for 5 min in the darkness. Then, 1 209 mL of NaCO<sub>3</sub> (5% w/v) was added and incubated in the dark for 1 h at room temperature. 210 Finally, absorbance was measured in a spectrophotometer, model P-220 (Biospectro, 211 Brazil), at 725 nm. To calculate the content of total phenolic compounds from crude 212 extracts, a standard curve of known concentrations of gallic acid was used (50 to 800  $\mu$ g  $mL^{-1} - r^2 = 0.99; y = 1.254$ ). 213

214 Statistical analyses. Effects of temperature and pH on both species ( $\Delta F/Fm'$ , 215 RCC, growth rate and phenolics content), and chemical changes in seawater, were 216 assessed statistically by using a factorial analysis of variance (Factorial-ANOVA). A post 217 hoc analysis of Student Newman-Keuls tests (SNK) was applied when Cochran's C test 218 and visual inspection of residuals were conducted to test homogeneity of variances 219 (Underwood 1997). A level of significance of 99% confidence interval (p = 0.01) was 220 applied. These analyses were performed using the software Statistica version 7 (StatSoft 221 Inc., Tulsa, OK, USA).

#### 222 **RESULTS**

#### 223 Photosynthetic responses

224 For both species, a significant interaction was found between temperature (18 and  $30^{\circ}$ C) 225 and pH (7.8 and 7.4) on photosynthetic performance (measured as  $\Delta$  F/Fm') (Factorial 226 ANOVA, p < 0.01; Figure 1; Table 2).  $\Delta F/Fm'$  significantly increased with respect to 227 basal values in L. crispatum after 6, 24 and 48 h experiments at 18°C under both 7.4 and 228 7.8 pH conditions (Figure 1A). A similar trend was observed in L. crispatum at 30°C 229 under pH 7.4, although without significant differences at 24 h experiments (Figure 1A). 230 F/Fm' significantly decreased under pH 7.8 at 30°C in all experimental times, with lower 231 values at 24 and 48 h experiments. In relation to S. capensis, no specific trends compared 232 to basal values were observed at 18°C under both pH conditions and at 30°C under pH 233 7.4 (Figure 1B). However, significantly lower  $\Delta$  F/Fm' levels were recorded at 30°C under pH 7.8, especially at 24 and 48 h experiments (Figure 1; Table 2). 234

#### 235 **Responses in calcification**

For both species a significant interaction was observed between temperature (18 and 30°C) and pH (7.8 and 7.4) on RCC (Factorial ANOVA, p < 0.01; Figure 2; Table 2). Similar trends were observed in RCC between studied species. When decreasing the pH from 7.8 to 7.4, RCC decreased significantly under both 18 and 30°C (Figure 2). The exception was at 30°C in *L* . *crispatum*, where no significant differences were detected between both experimental pH conditions (Figure 2). }

#### 242 **Relative growth rate**

No changes were recorded in growth rates in *L. crispatum*, with exception of the treatment at 30°C and pH 7.8, where negative values were observed (Figure 3A). A similar trend of negative growth rates were detected in *S. capensis* at 30°C and pH 7.8, although the rest of the treatments were similarly positive (Factorial ANOVA, p < 0.01; Figure 3B; Table 2).

#### 248 Phenolic compound

A significantly interaction was observed between temperature (18 and 30°C) and pH (7.8 and 7.4) on total phenolic content in *S. capensis* (Factorial ANOVA, p<0.01; Figure 4A). Under both experimental temperatures, total phenolic content increased at pH 7.4 with respect to 7.8.

#### 253 **DISCUSSION**

254 In this investigation, it was observed that the two calcareous macroalgae L. 255 crispatum and S. capensis suffered similar negative effects subject to the effects of 256 increased temperatures, from 18 (similar to nowadays winter temperatures) to 30°C 257 (comparable to summer end of the 21<sup>st</sup> Century temperatures), or decreased pH, from 258 current 7.8 to expected future 7.4 (by the year 2100); this was evidenced in terms of 259 photosynthetic performance ( $\Delta F/Fm'$ ), relative carbon content (RCC) and growth rates. 260 However, when increased temperatures (30°C) were combined with acidification (pH 7.4), at least in terms of photosynthesis and growth, both species seemed to be less 261 262 affected than when both stressors were applied independently. In terms of calcification, 263 acidification at 18°C caused a dramatic decay in RCC in both L. crispatum and S. 264 Moreover, although RCC decreased under increased temperatures and capensis. 265 acidification, compared to 18°C and pH 7.8, levels remained always higher than at 18°C 266 and 7.4 pH; this for both calcareous macroalgae species. Indeed, increased temperatures 267 appeared to diminish the negative effects induced by acidification, especially in relation to calcification, since at increased temperatures of 30°C and low pH of 7.4 both 268 269 macroalgae species displayed higher RCC if compared to treatments at 18°C and pH 7.4. 270 Total phenolic content measured in S. capensis showed the lowest levels at nowadays 271 temperatures and pH. These increased similarly at 18°C and pH 7.4, and at 30°C and pH 272 7.8, and with the highest values at 30°C and pH 7.4.

The detrimental effects of increased temperatures on *L. crispatum* and *S. capensis* could be attributed exceeding the tolerance thresholds of both species. Certainly, it has been observed that increased temperatures beyond tolerance ranges can induce an excess in the levels of reactive oxygen species (ROS) and subsequently oxidative stress in other photoautotrophs. For instance, it has been observed that the reaction catalyzed by the 278 enzyme ribulose-1,5-bisphosphate carboxylase oxygenase (RuBisCO), located in the 279 chloroplast and involved in carbon fixation and photorespiration, can cause greater levels 280 of the ROS  $H_2O_2$  under high temperatures due to increased oxygenase reactions (Kim and 281 Portis 2004). Moreover, it has been observed that increased temperatures induce 282 incomplete H<sub>2</sub>O oxidation in the PSII electron donor side, forming H<sub>2</sub>O<sub>2</sub>, which is then 283 reduced by manganese to the highly oxidizing HO• through Fenton reaction (Pospíšil 2016). In this context, Müller et al. (2012) found a systematic increase in the ROS  $\cdot O^{2-}$  in 284 parallel with greater temperatures of up to 18°C in the gametophytes of the brown 285 286 macroalgae Alaria esculenta, Laminaria digitata, and Saccharina latissima. An extended 287 oxidative stress in macroalgae is capable of causing damage and degradation to 288 photosynthetic pigments; thus, affecting photosynthesis and development (Moenne et al. 289 2016). The effects of high temperatures have been reported in the rhodolith *Lithophylum* 290 *margaritae*, which from 10 to  $30^{\circ}$ C showed optimum oxygenic photosynthetic threshold 291 at  $25^{\circ}$ C; however, when values exceeded these temperatures photosynthesis began to 292 decline (Steller et al. 2007). Reports by Vásquez-Elizondo and Enríquez (2016) demonstrated that from 30°C to 32°C, photosynthesis ( $F_{\nu}/F_m$ ) suffered nearly a 45% 293 294 decline after 3 d of experiments in rhodolith Neogoniolithon sp., articulate Amphiroa 295 tribulus and crustuse *Lithothamnion* sp. Despite the latter, similar increased-temperature 296 experiments (from optimal) on other tropical calcareous macroalgae, such as 297 Tricleocarpa cylindrica, Padina gymnospora, and Lithothamnion corallioides have not 298 evidenced negative effects in terms of photosynthetic performance (Scherner et al. 2016; 299 Noisette et al. 2013). Therefore, photosynthetic performance under increased 300 temperatures seems to be a species-specific feature in calcifying macroalgae.

301 In relation to the effects of acidification, our results demonstrated that negative 302 effects of increasing temperatures from 18 to 30°C were softened when decreasing the 303 pH from 7.8 to 7.4. Our records are similar to those reported by Semesi et al. (2009), 304 which observed that a progressive decrease in pH (from 9.8 to 7.6) were accompanied by 305 an exponential increase in photosynthetic  $O_2$  exchange in the rodolith *Hydrolithon* sp. 306 Recent studies by Scherner et al. (2016) demonstrated the differential effects of low pH 307 on photosynthesis in calcifying macroalgae; while from pH 8.1 to 7.2  $F_{\nu}/F_{m}$  decreased in 308 Lithophyllum stictaeforme, Pneophyllum conicum and Porolithon pachydermum, in 309 Tricleocarpa cylindrica it increased. Additionally, Kram et al. (2015) observed no change 310 in photosynthesis in the articulated Jania adhaerens and in the incrusting Lithothamnion 311 californicum from pH 8.1 to 7.6, although growth rates decreased. Even though photosynthesis enhancement has been observed in different macroalgae species subject to 312 313 acidification, published research suggests that this phenomenon may be more related to 314 CO<sub>2</sub> availability rather than reduction of pH itself. According to Martin and Hall-Spencer 315 (2017), higher concentration of  $CO_2$  in seawater induces macroalgae to rely exclusively 316 of  $CO_2$  diffusion, own-regulating their  $CO_2$ -concentrating mechanisms (CCM) to save 317 energy. Certainly, high availability of diffused CO<sub>2</sub> may provide greater substrate for 318 RUBISCO (ribulose-1,5-bisphosphate carboxylase/oxygenase) and carbon fixation, 319 inducing photosynthesis and growth (Martin and Hall-Spencer (2017).

320 The decalcification of calcifying macroalgae induced by acidification has been 321 already well described (Basso 2012; Feely et al. 2004). Likewise, we observed similar 322 decalcification (about 80%) under lower pH from 7.8 to 7.4 at 18°C in both L. crispatum 323 and S. capensis. Semesi et al. (2009) showed that a progressive decrease in pH values 324 from a 10 to 7.5 led to nearly 100% decalcification in Hydrolithon sp. Similarly, Noisette 325 et al. (2013) observed decrease in calcification rates from pH 8.1 to 7.7 in *Lithothamnion* 326 corallioides. Interestingly, these authors observed that at an increase in environmental 327 temperatures reduced the impact of pH 7.7 in calcification rates. In this context, other 328 studies on coralline algae have shown higher levels of calcification during summer, when 329 average sea surface temperatures are the highest (Steller et al. 2007; Martin and Gattuso 330 2009). Despite the latter, there are also investigations that account for the combined 331 negative effects of increased temperatures and acidification on calcification rates, as 332 described for the coralline *Porolithon onkodes* (Anthony et al. 2008; Diaz- Pulido et al. 333 2012). The information suggests that although temperature softened the detrimental 334 effects of acidification (and increased CO<sub>2</sub>) in L. crispatum and S. capensis, this seems to 335 be an intraspecific feature that cannot be applied to all calcifying macroalgae.

Another feature assessed in *S. capensis* was its phenolic content upon exposure to increased temperatures and decreased pH. Total phenolics increased at nowadays temperatures under acidification, but also at increased temperatures and subject to lower pH. Different investigations have revealed that increased temperatures and acidification are capable of inducing an oxidative stress condition due to de over-production of ROS
by different metabolic pathways (e.g. Celis-Plá et al. 2017; Flores- Molina et al. 2016;
Pospíšil 2016). It has been well described the role of phenolic compounds as antioxidants
to counteract ROS-excess (Moenne et al. 2016); thus, the records may indicate that *S. capensis* induces the production of phenolic compounds to inactivate ROS and avoid
oxidative damage.

346 Short-term assessment of temperature and acidification conditions on 347 physiological and metabolic features of L. crispatum and S. capensis reinforce the importance of local stressors to promote local baseline shifts. In addition, similar records 348 349 on other macroalgae species suggest that short-term responses may be similar to those 350 expressed in a long-term and, thus, may provide a good representation of potential 351 tolerance and survival thresholds under upcoming environmental conditions. For 352 instance, Chen et al. (2017) observed that within 60 min, light-saturated net 353 photosynthetic O<sub>2</sub> evolution rate (NPRm) in the green macroalga Ulva lactuca decreased 354 steadily when the pH dropped from 8.2 to 7.5 and, even more, to 6.5. After this time, 355 NPRm decrease stopped and levels were maintained without a trend of change. Indeed, 356 similar NPRm levels were observed in the alga exposed to pHs 7.5 and 6.5 at 60 min, 90 357 min and 8 days of experiments. Similar records on short-term exposure (within 48 h) to 358 future climate change scenarios have recorded in other macroalgae species belonging to different phylogenetic groups (e.g. Celis-Pla et al. 2014; Cruces et al. 2017; Flores-359 360 Molina et al. 2016).

In conclusion, we observed that both macroalgae species, the coralline *L. crispatum* and the aragonite *S. capensis*, displayed similar physiological responses to acidification. Moreover, and considering the interaction of higher sea surface temperatures and acidification, the information suggests that both assessed macroalgae species may be able to cope and develop under expected future marine environmental scenarios mediated by Climate Change.

367

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#### 580 Figures

**Figure 1.** Changes in maximum fluorescence ( $\Delta$  *F/Fm'*) emitted by (**A**) *Lithothamnium crispatum* and (**B**) *Sonderophycus capensis* exposed to increased temperatures and decreased pH for up to 48 h. Data correspond to mean  $\pm$  SD (*n*=9). Letters represent statistical differences at 95 % confidence interval (SNK tests, *p*< 0.05).

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**Figure 2.** Changes in relative carbon content (RCC), as a proxy of calcification rates, observed in (A) *Lithothamnium crispatum* and (B) *Sonderophycus capensis* exposed to increased temperature and decreased pH for up to 48 h. Data correspond to mean  $\pm$  SD (*n*=3). Letters represent statistical differences at 95% confidence interval (SNK tests, *p*< 0.05).

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**Figure 3.** Change in growth rate measured in (A) *Lithothamnium crispatum* and (B) *Sonderophycus capensis* exposed to increased temperature and decreased for up to 48 h. Data correspond to mean  $\pm$  SD (*n*=3). Letters represent statistical differences at 95% confidence interval (SNK tests, *p*< 0.05).

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**Figure 4.** Change in total phenolic compound in *Sonderophycus capensis* exposed to increased temperature and decreased pH for up 48 h. Data correspond to means  $\pm$  SD (*n*=6). Letters represent statistical differences at 95% confidence interval (SNK tests, *p*< 0.05).

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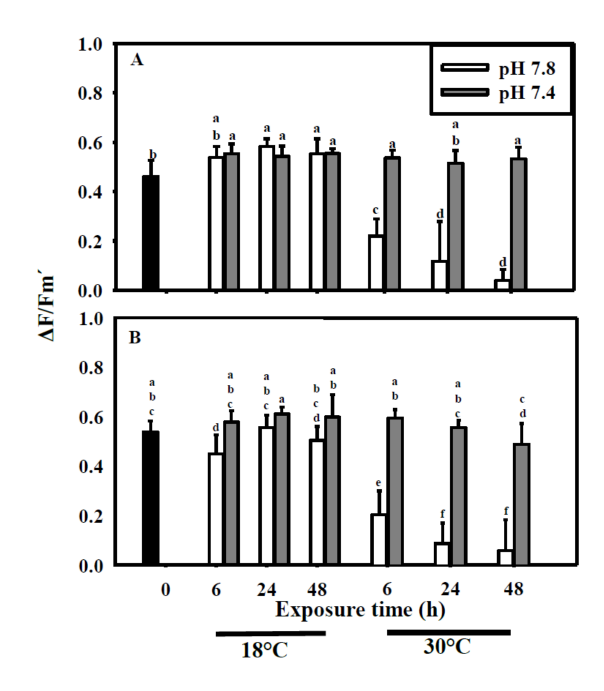
**Table 1** Summary of the chemical characteristics of experimental seawater, including T°, pH,  $pCO_2$ , HCO<sub>3</sub><sup>-</sup> and CO<sub>3</sub><sup>2-</sup> concentration,  $\Omega_{\text{calcite}}(\Omega \text{ Ca})$  and  $\Omega_{\text{aragonite}}(\Omega \text{ Ar})$ .

| 610 | T(°C) | рН    | pCO <sub>2</sub> | HCO <sub>3</sub> -         | CO <sub>3</sub> <sup>2-</sup> | ΩCa              | ΩAr              |
|-----|-------|-------|------------------|----------------------------|-------------------------------|------------------|------------------|
| 611 |       |       | (ppm)            | $(\mu mol \ kg \ SW^{-1})$ | $(\mumolkg\;SW^{\text{-}1})$  |                  |                  |
|     | 18±2  | 7.8±0 | 432±14           | 1164±51                    | 62.563±3.2                    | $1.491 \pm 0.07$ | 0.96±0.04        |
| 612 | 18±2  | 7.4±2 | 2035±19          | 1387±13                    | 18.816±1.8                    | $0.449 \pm 0.04$ | 0.290±0.02       |
|     | 30±2  | 7.4±2 | 2504±23          | 1590±15                    | 33.393±3                      | $0.811 \pm 0.07$ | $0.543 \pm 0.05$ |
| 613 | 30±2  | 7.8±0 | 455±1.10         | 1148±0.24                  | 95.524±0.6                    | 2.319±0.01       | 1.553±0.01       |

615 Note: n = 3 (means  $\pm$  SDs). Abbreviations: T (°C) = Temperature,  $pCO_2$  = partial 616 pressure of carbon dioxide,  $HCO_3^-$  = bicarbonate,  $CO_3^{2-}$  = carbonate,  $\Omega$  Ca = Calcite 617 saturation,  $\Omega$  Ar = Aragonite Saturation.

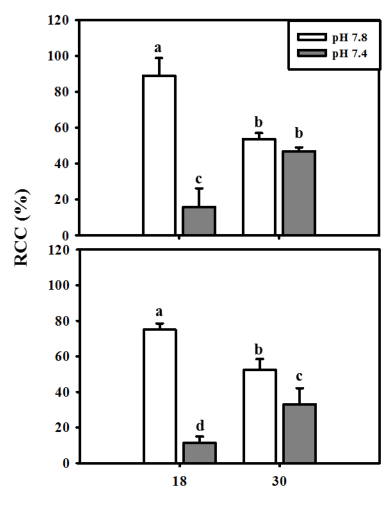
630 **Table 2** Summary table of Factorial ANOVA of the effects of temperature and pH 631 treatment on  $\Delta F/Fm'$  and calcification percentage in *Lithothamnion crispatum* and 632 *Sonderophycus capensis*.

|                                 |           |                       | Lith | othamnior | n crispati | ит       | Sona      | erophycus co | ipensis |         |
|---------------------------------|-----------|-----------------------|------|-----------|------------|----------|-----------|--------------|---------|---------|
|                                 |           |                       | df   | MS        | F          | p        | df        | MS           | F       | р       |
| 634                             |           | ∆F/Fm´                |      |           |            |          |           |              |         |         |
|                                 |           | Temperature (T)       | 1    | 0.25      | 110.53     | p<0.01   | 1         | 0.11         | 25.940  | p<0.01  |
| 635                             |           | pН                    | 1    | 0.25      | 108.05     | p<0.01   | 1         | 0.60         | 131.84  | p<0.01  |
| 055                             |           | T*pH                  | 1    | 0.20      | 88.84      | p<0.01   | 1         | 0.15         | 33.76   | p<0.01  |
|                                 |           | Error                 | 32   | 0.002     |            |          | 32        | 0.004        |         |         |
| 636                             |           | Calcification         |      |           |            |          |           |              |         |         |
|                                 |           | Т                     | 1    | 13.84     | 0.24       | 0,63     | 1         | 0.49         | 0.01    | 0,911   |
| 637                             |           | pH                    |      | 4793.10   |            | p<0.01   | 1         | 5159.99      | 141.58  | p<0.01  |
| 037                             |           | T*pH                  | 1    | 3301.41   | 59.40      | p<0.01   | 1         | 1474.34      | 40.45   | p<0.01  |
|                                 |           | Error                 | 8    | 55.58     |            |          | . 8       | 36.45        |         |         |
| 638                             | Note:     |                       |      |           |            |          |           |              |         |         |
| 639                             | n = 8 for | $\Delta F/Fm'; n = 3$ | for  | calcifi   | cation     | percenta | ige. Abbr | eviations    | T = Te  | mperatu |
| 640                             | degrees   | of freedom.           |      |           |            |          |           |              |         |         |
|                                 |           |                       |      |           |            |          |           |              |         |         |
| 641                             |           |                       |      |           |            |          |           |              |         |         |
| 041                             |           |                       |      |           |            |          |           |              |         |         |
|                                 |           |                       |      |           |            |          |           |              |         |         |
| 642                             |           |                       |      |           |            |          |           |              |         |         |
|                                 |           |                       |      |           |            |          |           |              |         |         |
|                                 |           |                       |      |           |            |          |           |              |         |         |
| 643                             |           |                       |      |           |            |          |           |              |         |         |
| 643                             |           |                       |      |           |            |          |           |              |         |         |
|                                 |           |                       |      |           |            |          |           |              |         |         |
| 643<br>644                      |           |                       |      |           |            |          |           |              |         |         |
|                                 |           |                       |      |           |            |          |           |              |         |         |
| 644                             |           |                       |      |           |            |          |           |              |         |         |
|                                 |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645                      |           |                       |      |           |            |          |           |              |         |         |
| 644                             |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645                      |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645<br>646               |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645                      |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645<br>646               |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645<br>646               |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645<br>646<br>647        |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645<br>646<br>647<br>648 |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645<br>646<br>647        |           |                       |      |           |            |          |           |              |         |         |
| 644<br>645<br>646<br>647<br>648 |           |                       |      |           |            |          |           |              |         |         |



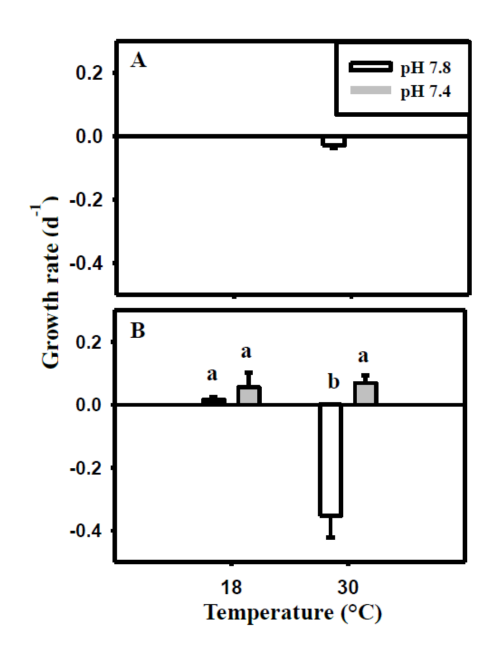


654 Figure 1



Temperature (°C)

**Figure 2** 





667 Figure 3

