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1	Community-level sensitivity of a calcifying ecosystem to acute in situ CO2 enrichment
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14	Running head: CO <sub>2</sub> enrichment of calcifying ecosystem
15	

### 16 Abstract

17 The rate of change in ocean carbonate chemistry is a vital determinant in the magnitude of impacts observed. Benthic marine ecosystems are facing an increasing risk of acute CO<sub>2</sub> 18 19 exposure, that may be natural or anthropogenically-derived (e.g. engineering and industrial activities). However, our understanding of how acute CO<sub>2</sub> events impact marine life is 20 restricted to individual organisms, with little understanding for how this manifests at the 21 22 community level. Here, we investigated, in situ, the effect of acute CO<sub>2</sub> enrichment on the coralline algal ecosystem - a globally ubiquitous, ecologically and economically important 23 24 habitat, but one which is likely to be sensitive to CO<sub>2</sub> enrichment due to its highly calcified 25 reef-like structures engineered by coralline algae. Most notably, we observed a rapid 26 community-level shift to favour net dissolution rather than net calcification. Smaller changes 27 from net respiration to net photosynthesis were also observed. There was no effect on the net 28 flux of dimethylsulphide / dimethylsulphoniopropionate (algal secondary metabolites), nor 29 the nutrients nitrate and phosphate. Following return to ambient CO<sub>2</sub> levels, only a partial 30 recovery was seen within the monitoring timeframe. This study highlights the sensitivity of biogenic carbonate marine communities to acute CO<sub>2</sub> enrichment, and raises concerns over 31 32 the capacity for the system to 'bounce back' if subjected to repeated acute high-CO<sub>2</sub> events.

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Keywords: calcification, photosynthesis, community, ecosystem, maerl bed, carbon dioxide,
acidification

## 36 Introduction

37 Long-term environmental change as a result of rising atmospheric CO<sub>2</sub> levels are projected to have significant impacts on marine organisms, especially those with calcified 38 body parts (Kroeker et al. 2010). Simultaneously, the risk of exposure to acute periods of 39 40 high-CO<sub>2</sub> conditions is also increasing, due to coastal / marine processes (e.g. tides (Abril et al.), upwelling (Lachkar 2014)), land runoff (Strong et al. 2014) and the development of 41 42 engineering activities such as carbon capture and storage (Blackford et al. 2015). Research has shown that the rate of environmental change is critical in determining the extent of 43 organismal damage, and that acute high-CO<sub>2</sub> exposures can have long-lasting effects (Burdett 44 45 et al. 2012, Kamenos et al. 2013). However, our understanding of how marine ecosystems 46 (rather than individuals) impact, and are impacted by, acute changes in ocean carbon chemistry is poorly understood (Pfister et al. 2014). This is despite the known importance of 47 48 key biological processes such as calcification, photosynthesis, respiration and nutrient uptake in driving marine ecosystem variability. 49

50 In the natural environment, an organism's response to environmental change is mediated 51 by community dynamics within the ecosystem. Failure to take these community-level interactions into account prevents macro-scale predictions of future ecosystem change 52 (Queirós et al. 2014). To date, the majority of acute or chronic environmental change 53 experiments have focused on one, or maybe two, environmental factors (e.g. increased  $CO_2$  / 54 55 temperature), and consider organisms in isolation (Riebesell & Gattuso 2015). However, whilst informing our mechanistic understanding of physiological responses, these types of 56 57 experiments are not representative of real-world impacts due to laboratory artefacts and the lack of appreciation for community-wide interactions (Cornwall & Hurd 2015, Riebesell & 58 Gattuso 2015). Consequently, efforts in developing methods for in situ experimentation have 59 60 recently increased.

61 Natural CO<sub>2</sub> vents, where the water column is enriched with CO<sub>2</sub> due to benthic bubbling of volcanic gases, have proven useful for understanding the impacts of long-term exposure to 62 63 a high CO<sub>2</sub> environment on marine ecosystem structure (Hall-Spencer et al. 2008, Fabricius et al. 2011, Kamenos et al. 2016). However, these study areas are typically characterised by 64 conditions more extreme or more variable than those predicted for the future, due to variation 65 in physical factors such as water currents and venting rates (Hall-Spencer et al. 2008). 'Free 66 67 Ocean CO<sub>2</sub> Enrichment' (FOCE) experimental setups attempt to bridge the gap between the precise control of laboratory experiments and the natural setting of CO<sub>2</sub> vents (Gattuso et al. 68 69 2014), by artificially exposing organisms or communities to a high  $CO_2$  environment. This also allows the effects of both chronic and acute CO<sub>2</sub> enrichment to be tested. Partially-70 artificial designs (where organisms are manually placed in the chambers, rather than 71 72 examining the natural system) have been conducted on tropical reefs (Kline et al. 2012) and in the deep sea (Barry et al. 2014), whilst smaller chambers deployed on tropical seagrass 73 beds have investigated the community-level response of this vegetated habitat to short-term 74 CO<sub>2</sub> enrichment (e.g. Campbell & Fourqurean 2014). 75

One of the potentially most susceptible groups of organisms to both long and short-term 76 77 CO<sub>2</sub> enrichment are the red coralline algae (Kroeker et al. 2010) – key ecosystem engineers 78 in the coastal zone (Riosmena-Rodríguez 2017). Coralline algal beds - supported by a freeliving coralline algal framework – are globally distributed (van der Heijden & Kamenos 79 2015), highly diverse (BIOMAERL 1999, Barbera et al. 2003) and biogeochemically active 80 81 (Burdett et al. 2015b, van der Heijden & Kamenos). However, the community susceptibility of coralline algal habitats is currently unknown, despite the real-world relevance of this 82 question compared to laboratory-based single organism studies (Gattuso et al. 2014). 83 Coralline algal beds are listed as 'Vulnerable' or 'Endangered' by the IUCN (Gubbay et al. 84 2016), a status driven by the sensitivity of coralline algae to environmental change, but also 85

due to the paucity of data available on the functioning of these habitats at the communitylevel.

Our understanding of coralline algal community functioning remains limited, even under 88 ambient conditions. Despite substantial gross primary production, coralline algal 89 90 communities exhibit net heterotrophy (i.e. O<sub>2</sub> uptake; Attard et al. 2015), acting as both a CO<sub>2</sub> source (Martin et al. 2007a) and organic carbon sink (Attard et al. 2015). While nutrient 91 92 availability is not thought to limit the growth of coralline algal ecosystems (Steller et al 93 2009), there is evidence that coralline algal communities act as a nutrient source, at least in the Mediterranean (Martin et al. 2007b). Coralline algae also represent a globally significant 94 95 stock of dimethylsulphoniopropionate (DMSP; Burdett et al. 2015a) – an algal secondary 96 metabolite that is the major precursor to the climate-gas dimethylsulphide (DMS). DMSP and DMS (DMS/P) drive a range of community interactions (e.g. grazing behaviour; Lyons et al. 97 98 2007), but it is not yet known if coralline algal communities are a net source or sink of these compounds. At an individual level, we know that CO<sub>2</sub> enrichment can affect the 99 100 photosynthesis, calcification and DMSP production of coralline algae (Burdett et al. 2012; Kamenos et al. 2013), but it is not currently understood how this is manifest at a community 101 102 level, despite the significant implications for ecosystem functioning.

Here, we investigated the effect of acute in situ CO<sub>2</sub> enrichment on key community-level, 103 biologically-driven processes in a temperate coralline algal bed. Periodic CO<sub>2</sub> enrichment is a 104 105 risk to marine habitats in this region due to the prevalence of human activities such as aquaculture – a rapidly expanding industry in Scotland and globally (OECD-FAO 2014). 106 107 Diel-scale pulsed release of  $CO_2$  can occur from aquaculture infrastructures due to periodicity in fish metabolism, e.g. after feeding (Forsberg 1997, Zakeś et al. 2003). In addition, the 108 development of carbon capture and storage facilities may further accentuate the risk of 109 110 periodic acute CO<sub>2</sub> release in the future (Blackford et al. 2015). Lithothamnion glaciale, the

- 111 coralline algal ecosystem engineer of this system, is known to be highly sensitive to acute
- 112 CO<sub>2</sub> exposure (Burdett et al. 2012, Kamenos et al. 2013), but sensitivity at a community level
- 113 remains unclear. Here, we investigated the integrated community-level response of a
- 114 coralline algal habitat to short-term CO<sub>2</sub> enrichment via in situ experimentation.

#### 116 Materials and Methods

# 117 Study site and experimental set-up

The experiment was performed on a coralline algal bed in Loch Sween, on the west 118 coast of Scotland, UK, at a depth of 6 m. The ecosystem framework is dominated by the free-119 120 living non-geniculate red coralline alga *Lithothamnion glaciale*, supporting a highly diverse community across multiple trophic levels. This includes both calcified and non-calcified 121 macroalgae (including Laminariale) and invertebrates, being particularly rich in Mullusca 122 (e.g. Aequipecten opercularis - queen scallops [~4 per 20 m<sup>2</sup>]) and particularly abundant in 123 Ophiuroidea (sea stars & brittle stars, e.g. *Ophiocomina nigra* [up to 10,000 per m<sup>2</sup>] and 124 Asterias rubens [~11 per 20 m<sup>2</sup>]) (BIOMAERL 1999, Barbera et al. 2003, Kamenos 2004). 125 Community biodiversity was not further quantified in this study. Four benthic chambers (28 126 litre volume, diameter = 38 cm) were deployed within the coralline algal bed by SCUBA 127 divers pushing them into the seabed. Chambers were left open for 24 hours to allow the water 128 129 within the chambers to equilibrate with the surrounding environment. Following 130 equilibration, lids were fitted and the experiment begun, which consisted of three phases: (1) 131 before CO<sub>2</sub> enrichment at ambient (control) conditions (15 hours), (2) during CO<sub>2</sub> enrichment (28 hours) and (3) post-enrichment recovery (37 hours). 132

133 Chambers were individually connected to the surface via a flow-through system, 134 which continually pumped water through the chamber via the surface at a rate of 120 L hr<sup>-1</sup> 135 (Swell UK Filter pump 5000). Pumps were located perpendicular from the chambers in 136 relation to the tidal current, to prevent the re-pumping of water through the system.  $CO_2$ 137 enrichment was achieved by bubbling pure  $CO_2$  directly into a mixing chamber on the 138 surface, prior to the water being directed to the main in situ chambers. pH (total scale) of 139 water in the mixing chamber was monitored using a pH probe (VitalSINE, daily 3 point

calibration following the manufacturer's instructions) and the rate of CO<sub>2</sub> bubbling was 140 adjusted as required to maintain a stable ~0.2 pH unit offset from the incoming water supply. 141 142 Actual pH change in the chambers (reflecting both the CO<sub>2</sub> addition and biogeochemical community processes) was determined by sampling the in-chamber water during the 143 experimental periods and analysing for total alkalinity (A<sub>T</sub>) and dissolved inorganic carbon 144 145 (C<sub>T</sub>), from which pH is calculated (details below). Flow-through circulation was maintained 146 for the duration of the experiment, except during 2-hour incubation periods when the water flow was stopped, but within-chamber circulation was maintained by stirring paddles (Attard 147 148 et al. 2015). Water samples were taken for determination of dissolved oxygen, carbonate chemistry, nutrients and dimethylated sulphur at the beginning and end of a 2-hour incubation 149 periods, which was carried out every ~12 hours during the experiment (i.e. around midday 150 and midnight during the three experimental phases). Measurements from the beginning and 151 end of the incubation were used for the determination of seabed flux measurements of each 152 parameter to gain understanding of the community response to CO<sub>2</sub> enrichment. All water 153 samples were collected in borosilicate glass syringes using SCUBA diving. Immediately after 154 collection, water samples were returned to the shore and prepared for various water chemistry 155 parameters, as detailed below. 156

## 157 Net photosynthesis / respiration (dissolved oxygen)

Winkler reagents (200  $\mu$ l each of 3M MnSO<sub>4</sub>.H<sub>2</sub>O solution and 200  $\mu$ l of 8M NaOH+4M NaI) were added to 12 ml unfiltered water samples for subsequent dissolved oxygen (DO) determination, and stored in the dark at 4°C until analysis. DO concentrations were determined using the Winkler titration method (Grasshoff et al. 2007): The sample was acidified with 200  $\mu$ l 5M sulphuric acid and titrated against 0.05M sodium thiosulphate solution with potassium iodate as a standard.

#### 164 Net calcification / dissolution (carbonate chemistry)

Samples for A<sub>T</sub> and C<sub>T</sub> were stored in borosilicate glass vials (Labco Ltd, UK) and poisoned 165 with mercuric chloride, following Dickson et al. (2007). AT was measured on a Metrohm 848 166 Titrino Plus using the 2-stage open-cell potentiometric titration method on 10 ml sample 167 volumes with 0.01 M HCl (Dickson et al. 2007). All A<sub>T</sub> samples were analysed at 25  $\pm$  0.1°C 168 with temperature regulation using a water-bath (Julabo 19). C<sub>T</sub> was determined by infra-red 169 170 detection of CO<sub>2</sub> from acidified samples on a dissolved inorganic carbon analyser (Marianda Airica). Additional carbonate chemistry parameters ( $pH_{NBS}$ ,  $pCO_2$ ,  $[HOC_3^-]$ ,  $[CO_3^{2-}]$ , 171 aragonite saturation state  $[\Omega_{Arg}]$ ) were calculated from A<sub>T</sub> and C<sub>T</sub> using CO2SYS (Pierrot et 172 173 al. 2006) with dissociation constants from Mehrbach et al. (1973), refit by Dickson and 174 Millero (1987) and KSO4 using Dickson (1990). In situ water temperature (°C), salinity and pH was measured hourly throughout the experimental period using an Exo2 multiparameter 175 176 sonde (YSI Inc). Nitrate and phosphate concentrations were calculated throughout the experimental period (below) and included in carbonate chemistry calculations. Net 177 community calcification rates were calculated using the alkalinity anomaly technique 178 (Chisholm & Gattuso 1991), based on the change in seawater A<sub>T</sub> during the incubation 179 period. For each mole of CaCO<sub>3</sub> precipitated (i.e. calcification), A<sub>T</sub> is lowered by two molar 180 181 equivalents. Therefore, the change in alkalinity can be converted to the mass of CaCO<sub>3</sub> precipitated. Certified seawater references materials for oceanic CO<sub>2</sub> (Scripps Institution of 182 Oceanography, University of California, San Diego) were used as A<sub>T</sub> and C<sub>T</sub> standards, 183 184 following Dickson et al. (2007).

## 185 Net DMS+DMSP (DMS/P<sub>T</sub>) flux

186 Samples for total (dissolved+particulate) DMS+DMSP (DMS/P<sub>T</sub>) were stored in 50 ml

187 crimp-top serum vials (Wheaton) fitted with Pharma-Fix lids. NaOH was added to a final

concentration of 0.03 M to hydrolyse DMSP into DMS. Samples were analysed by purgeand-trap gas chromatography (Turner et al. 1990), using an SRI 8610C GC fitted with a flame photometric detector (nitrogen carrier gas @ 8 psi). Sample concentrations were quantified via comparison to a DMSP standard (Research Plus Inc); sample detection limit was <1 nmol L<sup>-1</sup>, precision and accuracy for standards and samples was within 1%.

193 Net nitrate and phosphate flux

Unfiltered samples for nitrate and phosphate were stored in HDPE bottles (Fisher Scientific)
and frozen within 1 hour of collection. 10 ml samples were analysed for nitrate following the
cadmium reduction spectrophotometric method (Grasshoff et al. 2007); absorbance was
measured at 400 nm, with sodium nitrate used as a standard. 10 ml samples were analysed for
phosphate using the ammonium molybdate/ascorbic acid method (Grasshoff et al. 2007);
absorbance was measured at 885 nm, with potassium phosphate used as a standard.

200

# 201 Statistical analyses

Where parametric assumptions for normality and homogeneity of variance were met, 202 parametric tests were used to interrogate the data. One-way ANOVAs were used to test for 203 differences between ambient, CO<sub>2</sub> enrichment and recovery experimental phases in terms of 204 carbonate chemistry and net fluxes of DO, calcification rate, DMS/P<sub>T</sub>, nitrate and phosphate 205 206 (i.e. experimental phase included as a factor; no data transformation was required). Correlation tests were used to test correlation significance between fluxes of dissolved 207 oxygen, calcification, DMS/P<sub>T</sub>, nitrate and phosphate. Kruskall-Wallis tests were used to test 208 209 for differences in DO fluxes (parametric assumptions could not be met). Analyses were conducted using Minitab V14.1. 210

#### 211 **Results**

#### 212 Environmental conditions

213 Water temperature was  $15.3\pm0.32^{\circ}$ C and salinity was  $33.0\pm0.38$  throughout the experimental period (mean±SD, n=80). No significant difference in T<sub>A</sub> was observed between the three 214 215 experimental phases ( $F_{2,20} = 0.11$ , p = 0.89; Table 1). In contrast, C<sub>T</sub> was significantly higher during the CO<sub>2</sub> enrichment compared to the ambient / recovery phases ( $F_{2,20} = 31.6$ , p < 216 0.001; Table 1), resulting in a significant increase in  $HCO_3^-$  (F<sub>2,20</sub> = 10.45, p = 0.001) and 217 218  $pCO_2$  (F<sub>2,20</sub> = 4.24, p = 0.03). Mean aragonite saturation state and pH were reduced during CO<sub>2</sub> enrichment compared to the ambient / recovery phases, but not to the extent that 219 significant differences were observed ( $\Omega$ Ar: F<sub>2,20</sub> = 1.47, p = 0.26; pH: F<sub>2,20</sub> = 2.76, p = 0.09; 220 Table 1). Average in situ pH at the site in the 38 days before and during the experiment was 221 8.04±0.04 (mean±SD) (Figure S1). 222

## 223 Net photosynthesis / respiration (dissolved oxygen)

At ambient conditions, an average net uptake of O<sub>2</sub> (i.e. net respiration) was observed, 224 225 characterised by a small net release of O<sub>2</sub> during the day (i.e. net photosynthesis) to net respiration during the night (Figure 1). During the CO<sub>2</sub> enrichment average net O<sub>2</sub> release 226 increased compared to the ambient / recovery phases, reducing the difference between day 227 (higher net  $O_2$  release) and night (lower net  $O_2$  release / net uptake) measurements ( $F_{2,27}$  = 228 2.98, p = 0.07). During the recovery phase, net O<sub>2</sub> uptake decreased towards initial levels, but 229 did not quite reach the magnitude of net photosynthesis originally observed. When compared 230 231 separately, net oxygen flux was significantly higher in CO<sub>2</sub>-enriched conditions than ambient or recovery periods during the night ( $H_1 = 4.20$ , p = 0.040), but not during the day ( $H_1 = 1.70$ , 232 p = 0.192), reflecting the observed overall trend towards increased O<sub>2</sub> flux under CO<sub>2</sub> 233 enrichment (Figure 1). 234

## 235 Net calcification / dissolution (carbonate chemistry)

A significant reduction in net calcification was observed during the CO<sub>2</sub> enrichment compared to the ambient / recovery phases ( $F_{2,25} = 5.49$ , p = 0.01; Figure 1). Under ambient CO<sub>2</sub> conditions, the coralline algal community consistently exhibited a net calcification. During CO<sub>2</sub> enrichment, a significant shift towards net dissolution was observed. The recovery phase was characterised by an intermediate rate of net calcification. A significant negative correlation between DO flux and net calcification rate was observed (r = -0.40, p =0.05; Figure 1).

### 243 Net DMS/P<sub>T</sub> flux

Under ambient CO<sub>2</sub> conditions, there was a net uptake of DMS/P<sub>T</sub> by the coralline algal community of between  $11 - 24 \mu \text{mol m}^{-2} \text{ h}^{-1}$  (Table 2). During CO<sub>2</sub> enrichment there was a small reduction in net uptake rates, manifest as a shift towards the occasional net release of DMS/P<sub>T</sub>, but this change was not significant between experimental phases (F<sub>2,27</sub> = 0.62, p = 0.54; Table 2). DMS/P<sub>T</sub> flux was not significantly correlated with any of the other biogeochemical parameters, at p < 0.05.

# 250 Net nitrate and phosphate flux

Average net nutrient release and uptake rates were balanced (i.e. flux of ~zero), and no significant change was observed during CO<sub>2</sub> enrichment compared to the ambient / recovery phases (nitrate:  $F_{2,25} = 0.80$ , p = 0.46; phosphate:  $F_{2,25} = 0.01$ , p = 0.99; Table 2). Net benthic flux of phosphate, but not nitrate, was significantly correlated with benthic oxygen flux (r = 0.46, p = 0.02). No other significant correlations between net O<sub>2</sub>, nitrate, phosphate and DMSPt flux and net calcification rate (at p < 0.05) were observed.

#### 258 Discussion

Despite the known issues with investigating the effect of elevated  $CO_2$  in a laboratory setting, only a handful of in situ  $CO_2$  enrichment experiments have been conducted, and even less on the whole natural community. This is the first community-level in situ acute  $CO_2$ enrichment study in mid/high latitudes, and the first to consider the rate of recovery following acute  $CO_2$  perturbation. In this study, there was a rapid community level response to acute  $CO_2$  enrichment. This was particularly evident for net calcification, demonstrating the sensitivity of the whole community to acute  $CO_2$  exposure, not just individual species.

Unlike single-organism laboratory experiments, this study integrated the response of 266 the whole community. Whilst this means we are unable to assign individual species to 267 specific biogeochemical changes, the results obtained are relevant to real-world challenges 268 such as the designation of marine management strategies, which by necessity incorporate 269 whole communities (even if a particular species is the target focus). At the level of CO<sub>2</sub> 270 271 enrichment used in this study, the skeleton and epithelial cell surface of Lithothamnion 272 glaciale is compromised (Burdett et al. 2012, Kamenos et al. 2013), allowing for skeletal 273 dissolution (Langdon et al. 2000) – supporting the observed shift towards net community dissolution. This may have also been facilitated by dissolution of carbonate sediment and 274 dead sections of coralline algae, which cannot exert biological control and buffering against 275 changes in carbonate chemistry (Kamenos et al. 2013). Like other reef-based marine 276 ecosystems, this coralline algal community is highly diverse across multiple trophic levels 277 (BIOMAERL 1999, Barbera et al. 2003, Kamenos 2004). Calcifying invertebrates are 278 especially abundant (e.g. Ophiocomina nigra, which can make up 47% of total faunal 279 biomass; BIOMAERL 1999), and CO<sub>2</sub> enrichment is known to lead to a reduction in 280 calcification rate / increase in dissolution rate of these organisms (Kroeker et al. 2010). Thus, 281 282 these organisms are likely to have also contributed to the observed shift towards net

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dissolution, impacting their contribution to coastal CO<sub>2</sub> flux (Davoult et al. 2009). Due to the 283 high heterotrophic diversity of coralline algal beds (Barbera et al. 2003), only a small net 284 285 photosynthesis during the day was observed, supporting previous measurements using the Eddy correlation technique (Attard et al. 2015) and providing confidence that results recorded 286 do not represent treatment artefacts. CO<sub>2</sub> enrichment led to a small increase in net O<sub>2</sub> release, 287 suggesting an increased capacity for net photosynthesis – supporting the likely benefits of 288 289 elevated CO<sub>2</sub> conditions for aquatic photosynthetic organisms (Kroeker et al. 2010). Photosynthetic use of CO<sub>2</sub> can also provide a potential refuge for calcifying species by 290 291 buffering against the damaging effects of CO<sub>2</sub> enrichment (e.g. crustose coralline algae; Cornwall et al. 2014, Short et al. 2014, Kamenos et al. 2016), although this was not observed 292 in this study. Increased photosynthetic capacity may also increase the carbon sequestration 293 294 potential of these ecosystems (a key process in blue carbon storage; van der Heijden & Kamenos 2015), but a shift towards net dissolution may impact the stability of coralline algal 295 carbonate deposits. The balance and interaction of photosynthesis and calcification / 296 dissolution, and subsequent impact on carbon sequestration / storage is exemplified by the 297 observed correlation between net O<sub>2</sub> flux and net calcification. 298

299 Change in the community-level flux of dimethylated sulphur compounds appears to 300 be robust to acute CO<sub>2</sub> enrichment, despite the known sensitivity of coralline algal DMSP dynamics to acute CO<sub>2</sub> exposure (Burdett et al. 2012). Thus, it may be hypothesised that 301 although DMS/P<sub>T</sub> concentrations did not change, the proportion of the molecular species (e.g. 302 303 dissolved vs particulate, DMSP vs DMS) may have been altered, but this was not calculable by the approach employed. Nutrient fluxes were also insensitive to acute CO<sub>2</sub> enrichment, at 304 least at the level used in this study. However, the correlation between phosphate and DO 305 suggests that a larger CO<sub>2</sub> perturbation (in duration and / or magnitude) may impact 306 phosphorus cycling processes. 307

308	Acute CO <sub>2</sub> enrichment is just one aspect of carbon-chemistry pressures on marine
309	habitats. In addition, the combined effects of acute CO <sub>2</sub> enrichment and chronic, long-term
310	changes in carbonate chemistry may exacerbate biological responses. This has yet to be tested
311	at the community scale, despite the known importance of both acute and chronic $CO_2$
312	enrichment in driving responses in marine organisms. Surprisingly, even after a recovery
313	phase almost 1.5 times the length of the $CO_2$ enrichment, a full recovery (i.e. complete return
314	of all parameters to the initial measured rates) was not seen, at least in terms of the
315	parameters measured here, suggesting that, at best, there is considerable lag in community
316	recovery response times. This calls into question the capacity for the system to 'bounce back'
317	following repeated exposure to acute CO <sub>2</sub> inputs, which would be likely given the sources of
318	short-term CO <sub>2</sub> enrichment (e.g. aquaculture, CCS). Previous studies have shown that
319	damage to the coralline algal skeletal structure under CO2-enriched conditions can rapidly
320	occur (Burdett et al. 2012, Kamenos et al. 2013). In situ, this effect may manifest through to
321	the community level. Results from this study and others (e.g. Hall-Spencer et al. 2008,
322	Fabricius et al. 2011) collectively suggest that CO <sub>2</sub> enrichment may cause change across
323	biological scales, from the individual to community levels. If these changes persist in the
324	long-term, we may observe permanent transitions in community composition, perhaps one
325	that favours net photosynthesis, thereby tipping the balance in terms of biodiversity, and / or
326	net dissolution. Such transitions would not favour the growth of carbonate-depositing
327	ecosystem engineers such as coralline algae.

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459	Table 1. System parameters under ambient, CO2 enrichment and recovery phase conditions,
460	in benthic chambers deployed on a coralline algal bed in Loch Sween, Scotland. Water
461	temperature, salinity, photosynthetically active radiation (PAR), $A_T$ (total alkalinity) and $C_T$
462	(dissolved inorganic carbon) were directly measured; all other carbonate parameters were
463	calculated as detailed in the methods (pH is on NBS scale; $\Omega_{Arg}$ = aragonite saturation state).
464	Data presented as mean±SD (n=18, except for temperature and salinity, where n=80). Bold
465	text denotes parameters that were significantly different during the CO <sub>2</sub> enrichment phase (at
466	p < 0.05).

	Ambient conditions	CO <sub>2</sub> enrichment	Recovery period
Temperature (°C)	15.3±0.32	15.3±0.32	15.3±0.32
Salinity	33.0±0.38	33.0±0.38	33.0±0.38
Max PAR (µmol photons m <sup>-2</sup> s <sup>-1</sup> )	158	158	158
A <sub>T</sub> (μmol kg <sup>-1</sup> )	2190.7±87.2	2202.0±123.28	$2210.8 \pm 68.2$
Ст (µmol kg <sup>-1</sup> )	2084.8±12.8	2168.9±31.20	2066.2±23.2
pH <sub>NBS</sub>	7.9±0.2	7.7±0.39	8.0±0.2
<i>p</i> СО <sub>2</sub> (µаtm)	821.6±343.4	1747.7±1403.33	646.7±320.6
HCO3 <sup>-</sup> (µmol kg <sup>-1</sup> )	1961.1±27.5	2033.5±20.35	1927.6±49.2
CO <sub>3</sub> <sup>2-</sup> (µmol kg <sup>-1</sup> )	92.0±45.9	67.8±50.77	113.5±45.5
$\Omega_{ m Arg}$	1.4±0.7	$1.0\pm0.78$	$1.7 \pm 0.7$

- 469 **Table 2.** Community response of acute in-situ CO<sub>2</sub> enrichment in terms of net DMSPt, nitrate
- and phosphate flux, under initial ambient CO<sub>2</sub> conditions, during CO<sub>2</sub> enrichment and during

	Ambient conditions	CO <sub>2</sub> enrichment	Recovery period
Net DMSPt flux ( $\mu$ mol m <sup>-2</sup> h <sup>-1</sup> )	-23.13±27.12	-13.46±28.12	-11.47±11.39
Net nitrate flux (mg m <sup>-2</sup> h <sup>-1</sup> )	-11.40±36.11	-0.55±19.90	7.71±27.37
Net phosphate flux (mg m <sup>-2</sup> $h^{-1}$ )	$0.04 \pm 0.44$	$0.02 \pm 0.24$	$0.05 \pm 0.29$

471 the recovery phase at ambient  $CO_2$ . Data presented as mean $\pm$ SD.

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Figure 1. Community response of acute in-situ CO<sub>2</sub> enrichment in terms of net dissolved
oxygen flux and net calcification rate, under initial ambient CO<sub>2</sub> conditions (black circle),
during CO<sub>2</sub> enrichment (white circle) or during the recovery phase at ambient CO<sub>2</sub> (grey
circle). Data presented as mean±SD.



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