

Author response to comments from anonymous referee #1 - manuscript bg-2016-63

We thank the reviewer for their constructive comments. Below please find our point by point responses to the referees' specific comments.

- 1) *The reviewer asks whether using a glycine-ammonium buffer (pH 10.5) would be beneficial prior to measuring fluorescence.*

As the reviewer notes, the pH effect for the MUF substrate was previously known. The objective of our research was to investigate if there was a pH dependency in the MCA assay, and whether our assay required buffering. We have shown that for this substrate it is important to control pH. We wished to maintain a constant and defined pH throughout the incubation as well as the fluorescence measurement, as the specific activity of enzymes' can vary with pH. Thus we favoured buffering at a pH relevant to enzyme activity, rather than allow pH to "roam" during the incubation, with potential variations in product formation between experiments.

- 2) *The reviewer notes that fluorescence was determined at the same wavelengths for MUF and MCA, but the fluorescence spectra of the two molecules are different. The review asked whether this slight difference could have any effect on the results and conclusions?*

For robustness we used filter blocks in our sea-going plate reader. 365 nm excitation and 460 nm emission wavelengths was the only block available that covered the correct wavelengths. Although we may be off the ideal excitation and emission wavelengths, which raises our limit of detection, this should not alter the results or conclusions, unless there was something else in the system giving an emission at that wavelength and whose emission was pH responsive. We ran trials with natural seawater to ensure there was no inherent interference with fluorescence. Variation in the wavelengths used for MUF & MCA also exists within current literature. For instance, Chrost (1992) use the same MUF excitation and emission as we report, but used 380 nm excitation and 440 nm emission for MCA (as stated by the reviewer), while Hoppe et al (1988) used 365 excitation and 445nm. Christian & Karl (1995) used 360 nm excitation and 447nm emission for MUF, while both Mass et al. (2013) and Piontek et al. (2009, 2010, 2013) uses 355nm excitation and 460 nm emission for both MUF & MCA. In our study, it was important to use wavelengths used by others for consistency and comparison of responses.

Author response to comments from anonymous referee #2 - manuscript bg-2016-63

We thank the reviewer for their constructive comments. Below please find our point by point responses to the referees' specific comments.

In my opinion, the authors' focus is to narrow on extracellular enzyme activities and substrate fluorescence. I suggest broadening the discussion of the results including OA effects on bacterial growth and production and changing the conclusion section as well as the abstract accordingly.

We have broadened the discussion to include additional bacterial responses to OA. The conclusion and abstract have been amended accordingly.

The authors should stress their point that buffering is necessary when determining enzyme rates in general, or at least when using MCA as a marker.

We have further emphasized this observation.

In contrast, the effect of pH on MUF fluorescence is well known (e.g. Mead et al. 1955), explicitly written in the Sigma product information and usually considered in enzyme rate measurements. Furthermore, I am not convinced by the authors' proposed effect of pH on MCA fluorescence.

We acknowledge previous studies on the effect of pH on MUF fluorescence, and have revised our interpretation on the effect of pH on MCA fluorescence (see below).

Specific comments:

p. 1 l. 17f: Change to "This study investigated the potential artefacts in determining the response of bacterial growth and activity to ocean acidification, and the relative effects of three different acidification techniques."

This has been changed in the revised manuscript.

p. 1 l.26ff: From the presented results I would conclude that "bubbling may stimulate carbohydrate degradation and bacterial growth".

This has been changed in the revised manuscript.

p. 2 l.32ff Add some more information on extracellular enzyme characteristics: Enzymes are considered as the rate limiting step in hydrolysis of HMW-substrate by bacteria. Both enzyme groups consist of several isoenzymes that catalyze the same reaction but may vary significantly in e.g. pH or temperature optimum and sensitivity (e.g. broad range or narrow optimum range). Define "extracellular enzyme". Do you include cell-attached and particle-attached enzymes or only free enzymes?

Additional information has been incorporated as requested. Yes, the presented enzyme activities also include any cell- or particle attached enzyme activity. The definition of 'extracellular' in the context of this work has been clarified in the revised manuscript.

p. 2 l.56: What are "indirect influences on longer timescales"? Please specify.

We are referring to potential indirect effects on enzymatic rate, due to factors including altering enzyme abundance, changes in substrate arising from plankton community composition change, grazing or viral lysis, abiotic influences on rate (e.g. through pH effects or carbamylation), and type of enzyme synthesized, as opposed to direct effects of change in pH on the enzyme activity. This has been clarified in the revised manuscript.

p. 3 l. 66ff: The pH sensitivity of MUF is well known (e.g. Mead et al. 1955 and SIGMA product information). Please clarify this in the text. p. 4 l. 107: see above

This has been clarified in the revised manuscript.

p. 5 l. 130ff: It would be very interesting to see the kinetic curves of the independent tests that the authors mention. Please provide a short table or graph. Enzyme kinetics and maximum velocities may vary from one seawater sample to the other (depending on isoenzymes present in the sample). Did you test enzyme kinetics both, in summer and spring?

The tests to calculate the optimum substrate concentrations were conducted by another researcher using seawater collected from a similar location. We agree that enzyme kinetics and maximum velocities may vary spatially and temporally; however, as the purpose of this study was to assess enzyme response to acidification, and to different methods of acidification, full enzyme kinetics were not required. We believe that the variance between samples collected from different sites acts as a reasonable proxy for the variance that would be found between different seasons at the same site.

p. 5 l. 132f: At which pH did you calibrate MUF?

At pH 7.8 and 8.1. This has been clarified in the revised manuscript.

p. 7 l. 182: Please include data (e.g. as supplementary graph)

As requested, the following table compares average LAP activity buffered in 0.1 M Tris and MOPS at pH 8.1 in coastal seawater. This has been added as supplementary material (S1) in the revised manuscript.

Trial	Tris activity (nmol l ⁻¹ h ⁻¹)	MOPS activity (nmol l ⁻¹ h ⁻¹)
1	51.54 (±2.32)	43.42 (±1.43)
2	35.92 (±0.81)	29.34 (±1.08)

p. 8 l. 216: Did you determine pH at the end of the incubations?

pH was determined at 0, 24, 48, 72 and 96 h in both trials. This data has been added to supplementary material (S2 and S3).

p. 8 l. 220: Was there a reason to incubate under artificial light instead of dark incubations?

The aim was to simulate natural conditions in coastal waters.

p. 9 l. 270ff: Can you please give the standard deviation of your calibration? An increase by 4% only seems to me very low and within the experimental detection limit. Previous studies did not detect a significant effect of pH on MCA fluorescence and I would not consider 4% to be significant. It would be useful, if you could provide a graph with the calibration curves of both fluorescent markers at pH 8.1 and pH 7.8!

We were prompted by the reviewer's concern to refer back to this data and consider other data not previously interpreted for this paper. We find a similar trend in this revised data set, (shown in revised Table 1), but not an acceptable level of significance ($p > 0.05$). We thank the reviewer for raising the issue, and in the light of this more recent analysis have revised this statement to clearly state that there

is no robustly significant difference in the fluorescence of Tris buffered MCA or MUF between 7.8 and 8.1. This result is included in the abstract.

p. 10 l. 299 I agree that different acidification methods had significant effects on BG activities, but I cannot see a significant effect on LAP activity from the presented data (Figure 1). In p. 11 l. 318 the authors state that, although cell-specific LAP activity showed evidence of a response to acidification, this was not significant in either trial. Please explain/clarify and give statistical evidence.

We agree with the reviewers comment that this sentence is confusing and requires rewording. The reference to "significant effects" in Line 299 refers to the response of different treatments at select sampling points only. For instance, activity in treatment A was significantly higher than the Control at 48 and 72 h during trial 1 (t-test, df=4, $p < 0.001$) (Fig 1), while activity in treatment B was frequently higher than the Control in both trials however was not significantly greater. So although there was 'evidence' of a response to acidification, these were not statistically significant. This has now been clarified in the revised manuscript.

It would be also interesting to see the data for AG and AAP activity.

Trial 1 AG and AAP activity, and trial 2 AAP activity have now been included in the supplementary material (S4, S5, and S6 respectively). AG activity was very low during trial 2, being below detection at several sampling points across all treatments; consequently we mention this, but do not include the data in the supplementary material.

p. 12 l. 337ff: What about total secondary production rates? How do you explain the difference towards the end of trial 2? Can you relate it to changes in BG activity?

The response in secondary production in the latter part of trial 2 (Fig. 3) does not reflect the response in BG activity in the different treatments (Fig. 1). The increase in BG activity is highest in treatment B relative to the other treatments, while the increase in secondary production occurs in treatment A. The response in secondary production observed from 72 h during trial 2 may be explained by other indirect factors, as discussed on l. 355.

p. 13 l. 367: The authors state that the introduction of CO₂-air gas mixtures using gas-permeable tubing would be the "most robust technique to investigate the response of bacterial processes to future OA conditions". This is ignoring the fact that there are more techniques commonly used which were not tested in this study and may be even "more robust". Furthermore, I would conclude from this study that different techniques may result in different results. They may under- or overestimate certain parameters at the same time but not all parameters equally.

We agree and have now highlighted the fact that this study only investigates some of the commonly used methods to artificially acidify seawater. We have altered the abstract and conclusion accordingly.

1 Assessing approaches to determine the effect of ocean 2 acidification on bacterial processes

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12

13 **Abstract**

14 Bacterial extracellular enzymes play a significant role in the degradation of labile organic
15 matter and nutrient availability in the open ocean. Although bacterial production and
16 extracellular enzymes may be affected by ocean acidification, few studies to date have
17 considered the methodology used to measure enzyme activity and bacterial processes. This
18 study investigated the potential artefacts in determining the response of ~~bacterial growth and~~
19 ~~activity~~ ~~bacterial~~ ~~and~~ extracellular glucosidase and aminopeptidase ~~activity~~ to ocean
20 acidification, and the relative effects of three different acidification techniques. Tests
21 confirmed that the ~~observed effect of pH on~~ fluorescence of ~~the~~ artificial fluorophores ~~was~~
22 ~~affected by pH,~~ ~~and that the influence of addition of the~~ MCA fluorescent substrate ~~on alters~~
23 seawater ~~sample pH,~~ ~~were both negated~~ ~~overcome~~ by the use of Tris buffer. In experiments
24 testing different acidification methods, bubbling with CO₂ gas mixtures resulted in higher β-
25 glucosidase activity ~~and 15–40 % higher bacterial abundance,~~ relative to acidification ~~by their~~
26 ~~introduction~~ via gas-permeable silicon tubing, ~~or by and~~ acid addition (HCl). ~~In addition,~~
27 ~~bacterial numbers were 15–40 % higher with bubbling relative to seawater acidified with gas-~~
28 ~~permeable silicon tubing and HCl.~~ ~~BB~~ubbling may ~~therefore stimulate~~ ~~lead to overestimation~~

29 | ~~of~~ carbohydrate degradation and bacterial ~~growth abundance~~, ~~leading to the~~ and consequently
30 | incorrect interpretation of the impacts of ocean acidification on organic matter cycling.

31 | **1 Introduction**

32 | Proteins and carbohydrates constitute two of the most common labile organic substrates in
33 | the ocean (Benner, 2002; Benner et al., 1992; McCarthy et al., 1996), both of which are
34 | essential for cellular growth and repair (Azam et al., 1983; Simon and Azam, 1989). ~~Labile~~
35 | ~~substrate availability is limited by bacterial enzyme--driven hydrolysis of high molecular~~
36 | ~~weight organic material (Azam and Cho, 1987, Munster, 1991).~~ Two groups of ~~bacterial~~
37 | extracellular enzymes (~~attached or free of the cell~~ released into surrounding water) commonly
38 | studied for their role in protein and carbohydrate degradation are aminopeptidases and
39 | glucosidases, respectively. ~~The activity of individual enzymes are responsive to changes in~~
40 | ~~environmental factors, and so overall glucosidase and peptidase activities will have different~~
41 | ~~pH optima~~ Each enzyme group Enzyme activity is sensitive to ~~changes in different~~
42 | ~~environmental factors, for example, and consequently degradation of proteins and~~
43 | ~~carbohydrates will vary accordingly. M~~ many of enzymes are pH sensitive and have different
44 | ~~pH optima~~ (Tipton and Dixon, 1979, Piontek et al., 2013), ~~and consequently~~ Consequently a
45 | change in ocean pH may result in a decline or increase in activity of extracellular enzymes as
46 | these are directly exposed to the external seawater pH (Orsi and Tipton, 1979; Tipton and
47 | Dixon, 1979).

48 | Atmospheric CO₂ has increased by 40 % since the 18th century (IGBP-IOC-SCOR, 2013;
49 | IPCC, 2013), which is of concern as CO₂ freely exchanges with the ocean and directly alters
50 | ocean carbonate chemistry and pH. As a result ocean pH has declined from 8.2 to 8.1, with a
51 | continued decline to 7.8 predicted by the year 2100. This decline in ocean pH and the
52 | associated change in carbonate chemistry, referred to as ocean acidification (OA), will
53 | significantly impact metabolic reactions and influence carbon cycling in the ocean (Endo et
54 | al., 2013; Engel et al., 2014; Piontek et al., 2010; Riebesell et al., 2007). For this reason,
55 | researchers have investigated the sensitivity of a wide range of biotic and abiotic factors to
56 | future changes in ocean pH and the carbonate system.

57 | Bacterial extracellular enzyme activity has been investigated in OA studies (reviewed in
58 | Cunha et al., 2010) due to the important role they play in the degradation of ~~labile high~~
59 | ~~molecular weight~~ organic matter (Azam and Ammerman, 1984; Azam and Cho, 1987; Law,

60 1980; Münster, 1991) and the vertical flux of carbon to the deep ocean (Piontek et al., 2010;
61 Riebesell and Tortell, 2011; Segschneider and Bendtsen, 2013). Current research suggests
62 that bacterial extracellular enzyme activities may increase under future OA conditions
63 (Grossart et al., 2006; Maas et al., 2013; Piontek et al., 2010, 2013; Yague and Estevez,
64 1988). This may result from the direct effect of pH on the ionisation state of the enzyme's
65 component amino acids (Dixon, 1953), or from indirect influences potentially altering
66 enzyme production on longer timescales (Boominadhan et al., 2009). Examples of the latter
67 include ~~The latter may be arise in response to~~ changes in the concentration and composition
68 of high molecular weight organic substrate due to the effect of pH on phytoplankton and
69 bacterioplankton community composition (Endo et al., 2013; Engel et al., 2008; Riebesell,
70 2004; Witt et al., 2011), bacterial secondary production and cell numbers (Endres et al., 2014;
71 Maas et al., 2013), and phytoplankton-derived organic exudation (Engel, 2002; Engel et al.,
72 2014).

73 Bacterial extracellular enzyme activity is regularly determined using artificial fluorogenic
74 substrates. These substrates consist of a fluorescent moiety covalently linked to one or more
75 natural monomer molecules (Arnosti, 2011; Kim and Hoppe, 1984). The molecule is non-
76 fluorescent until it is hydrolysed by an extracellular enzyme, which triggers a fluorescent
77 response, allowing it to be detected and quantified (Hoppe, 1993). The sensitivity of the
78 analytical method to pH has been assessed in terrestrial soils (Malcolm, 1983; Niemi and
79 Vepsäläinen, 2005), however limited information is available on how these components
80 respond to a reduction in seawater pH (Piontek et al., 2013). If pH does have a significant
81 effect on the individual assay components, and this is not corrected, then calculated enzyme
82 kinetics will under or overestimate the true activity rates.

83 Several different methods ~~are have been commonly~~ used to artificially adjust seawater pH in
84 experimental systems (Cornwall and Hurd, 2015; reviewed in Riebesell et al., 2010). The
85 simplest acidification method involves the addition of a strong acid (typically HCl). The acid
86 decreases the sample pH through the formation of hydronium ions and modifies total
87 alkalinity (TA), but does not alter dissolved inorganic carbon (DIC) in a closed system
88 (Emerson and Hedges, 2008); consequently although it is relatively simple to adjust pH using
89 acid, the balance of carbonate species does not reflect the changes that will occur in response
90 to increased CO₂ uptake unless corrected for by the addition of a base (Iglesias-Rodriguez et
91 al., 2008; Riebesell et al., 2010). Another method for acidifying seawater is the use of CO₂-

92 Air gas mixtures, which alter the seawater carbonate species in ratios predicted to occur from
93 the uptake of atmospheric CO₂ under future scenarios (Gattuso and Lavigne, 2009; Riebesell
94 et al., 2010; Rost et al., 2008; Schulz et al., 2009). Schulz et al. (2009) suggest that microbial
95 organisms are likely to respond to changes in carbonate species (e.g. CO₂, HCO₃⁻ or CO₃²⁻),
96 rather than changes in overall DIC or TA. A review by Hurd et al. (2009) concluded that
97 differences in carbonate chemistry arising from the use of different acidification
98 methodologies can influence phytoplankton photosynthesis and growth rates, as well as
99 particulate organic carbon production per cell, and so it is important to ensure changes in all
100 carbonate system species reflect that projected from an increase in CO₂ (Cornwall and Hurd,
101 2015).

102 In addition to the method of acidification, the mode of application also needs to be
103 considered. A commonly used method of introducing CO₂-Air gas mixtures into seawater is
104 by bubbling. This method is simple to implement and maintain for extended periods,
105 however, the physical disturbance associated with bubbling CO₂ gas may influence
106 coagulation of organic matter (Engel et al., 2004; Kepkay and Johnson, 1989; Mopper et al.,
107 1995; Passow, 2012; Schuster and Herndl, 1995; Zhou et al., 1998), as well as microbial
108 interactions (Kepkay and Johnson, 1989). This mechanical disturbance may be particularly
109 exacerbated when bubbling is used in small-volume incubations at the laboratory/microcosm
110 experimental scale (<20 litres). An alternative method of introducing CO₂ gas is by using
111 gas-permeable tubing (Law et al., 2012; Hoffmann et al., 2013), which eliminates physical
112 artefacts associated with bubbling whilst achieving realistic future carbonate chemistry.
113 Previous research has been conducted comparing the effect of acid addition and CO₂ gas
114 bubbling on phytoplankton growth, with no significant ~~effect-difference~~ detected (Chen and
115 Durbin, 1994; Hoppe et al., 2011; Shi et al., 2009). However, to date no comparison of the
116 bacterial response to seawater acidified with acid and CO₂ gas aeration has been carried out.
117 In addition, there are no published comparisons of CO₂ gas ~~mixtures~~ introduced through gas-
118 permeable silicon tubing with bubbling to assess their suitability for OA research.
119 Consequently the aims of the following study were two-fold; to ~~determine-identify any~~
120 ~~artefacts associated with~~ the ~~effect-of pH-on-the-sensitivity-of~~ use of fluorogenic substrates
121 ~~used-in bacterial-extracellular~~ enzyme analysis, and also to compare the response of bacterial
122 processes to different ~~approaches-methods~~ of acidification in small-volume incubations.

123

124 **2 Material and methods**

125 **2.1 pH determination**

126 Sample pH was determined using a CX-505 laboratory multifunction meter (Elmetron)
127 equipped with a platinum temperature integrated pH electrode (IJ44C-HT enhanced series;
128 accuracy 0.002 pH units), calibrated using Tris buffers (Cornwall and Hurd, 2015) and
129 regularly cleaned using potassium chloride reference electrolyte gel (RE45-Ionode).
130 Electrode pH measurements were validated using a pH spectrophotometer with colorimetric
131 determination using a thymol blue dye solution (Law et al., 2012; McGraw et al., 2010).
132 Following recommendations in the European Project on Ocean Acidification (Riebesell et al.,
133 2010), pH values in this research reflect the total hydrogen ion scale (pH_T).

134

135 **2.2 Extracellular enzyme activity**

136 The activity of two proteases was examined, with arginine aminopeptidase activity (AAP)
137 quantified using L-arginine-7-amido-4-methylcoumarin hydrochloride (Arg-MCA), and
138 leucine aminopeptidase activity (LAP) quantified using L-leucine-7-amido-4-
139 methylcoumarin hydrochloride (Leu-MCA). Two glucosidases were also examined; α -
140 glucosidase activity (AG) was quantified using 4-Methylumbelliferyl α -D-glucopyranoside
141 (α -MUF), and β -glucosidase activity (BG) was quantified using 4-Methylumbelliferyl β -D-
142 glucopyranoside (β -MUF, all from P212121 LLC, USA). Artificial fluorogenic substrate was
143 added to each seawater sample to give a final substrate assay concentration of 39 μ M, which
144 was determined from independent tests to be the optimum concentration for calculating the
145 maximum velocity of enzyme hydrolysis in seawater samples (data not shown). A four point
146 calibration curve (0, 4, 40, 200 nM final concentration) was created [at both pH 7.8 and 8.1](#)
147 using 4-methylumbelliferone (MUF) for glucosidase activity, with a separate calibration curve
148 (0, 40, 400, 4000 nM final concentration) created using 7-amino-4-methylcoumarin (MCA)
149 for protease activity (Sigma-Aldrich). UltraPure distilled water (InvitrogenTM, Life
150 Technologies) was used as a sample blank. Each sample was assayed in triplicate using a
151 single 96-microwell flat bottom black assay plate (Nunc A/S), with a separate enzyme assay
152 performed for glucosidase and protease activity. Each assay plate was read at 5 min intervals
153 for a minimum of 3 h using a Modulus microplate reader (Turner Biosystems) at 365 nm
154 excitation and 460 nm emission wavelength as in Burrell et al., (2015). Incubation assay

155 temperature matched the seawater temperature at the sampling site. The potential for
156 outgassing and associated increase in sample pH during the 3 h enzyme assay was not tested.
157 The maximum potential enzyme rate (V_{\max} , $\text{nmol l}^{-1} \text{h}^{-1}$) was approximated from the
158 saturating substrate concentration of 39 μM . Triplicate V_{\max} approximations were averaged
159 per sample. Cell-specific rates were calculated by dividing the activity per litre by bacterial
160 cell numbers per litre. The assay tests were carried out using surface seawater collected from
161 the south coast of Wellington, New Zealand ($41^{\circ}20'53.0''\text{S}$, $174^{\circ}45'54.0''\text{E}$).

162 **2.3 Enzyme assays**

163 **2.3.1 The effect of pH on fluorophore fluorescence**

164 The effect of pH on fluorophore fluorescence was investigated at both typical (Hoppe, 1983)
165 and elevated fluorophore concentrations using two different buffer solutions, the organic
166 solvent 2-methoxyethanol (Sigma-Aldrich) and 0.1 M Tris/HCl. The pH of MUF and MCA
167 fluorophore working standard (200 μM) diluted in 1 % 2-methoxyethanol (Sigma-Aldrich)
168 was first recorded (pH 6.22 and 6.58 at 18.6 $^{\circ}\text{C}$ respectively). Each fluorophore was then
169 diluted to 4000, 20000 and 40000 nM (referred to as high concentrations) at four pH values
170 (8.2, 8.1, 7.9 and 7.8) in triplicate by addition of 0.1 N aqueous NaOH. The MUF and MCA
171 fluorophore working standards made up in ~~in~~ 0.1 M Tris/HCl were prepared at pH 8.1 and 7.8
172 only, and also carried out at lower concentrations (MUF: 4, 40, 200 nM; MCA: 40, 400, 4000
173 nM).

174 **2.3.2 The effect of artificial fluorogenic substrate on seawater pH**

175 Individual seawater samples were adjusted to pH 7.95 and 7.70 using 0.1 M HCl. All four
176 artificial fluorogenic substrates previously described were made up to working standards
177 using 1 % 2-methoxyethanol (Sigma-Aldrich). A time-zero reference pH was recorded for
178 each seawater sample and, following the addition of each substrate at 39 μM final
179 concentration, sample pH was recorded immediately and after 30 min. Each artificial
180 fluorogenic substrate was run in triplicate at both pH values, and compared to controls
181 without substrate addition at both pH levels.

182 **2.3.3 Buffering artificial substrates**

183 Duplicate trials were undertaken to determine if 0.1 M Tris/HCl could successfully buffer
184 MCA substrate at the working concentration (39 μM) when added to seawater of similar pH.

185 Tris buffer contains an amine group which can affect peptidase activity (Baker and Prescott,
186 1983; Desmarais et al., 2002; Saishin et al., 2010), and so tests were carried out to compare
187 the impact of different buffers. LAP activity was compared in seawater using LAP substrate
188 (39 μ M final concentration) buffered with 0.1 M Tris/HCl or 3-(N-
189 morpholino)propanesulfonic acid (MOPS) with pH adjusted to 8.1. Enzyme activity was also
190 determined in seawater (pH 8.18). A non-buffered LAP substrate addition was not included
191 due to the acidic nature of the aminopeptidase substrate (non-buffered LAP substrate was pH
192 5.87). MOPS has been used as a buffer in studies of the effects of pH on enzymes (Piontek et
193 al. 2010), and so was an appropriate comparison. Borate buffers were not trialled because
194 they have a bactericidal effect on microbial activity (Houlsby et al., 1986). In two separate
195 test experiments using coastal seawater Tris/HCl buffer did not inhibit LAP activity relative
196 to MOPS but instead showed a minor stimulatory effect with 16-18% higher LAP activity
197 (~~data not shown~~ [supplementary material S1](#)). Tris/HCl was selected for subsequent use as its
198 optimal buffer range is pH 7.8-9.0, making it ideal for OA incubations, and it has a pKa of
199 8.06, so is appropriate for artificial fluorescent substrates (Hoppe, 1993).

200 ~~Following~~ Based on the buffer above ~~trials~~ ests, the following methodology was used for the
201 seawater acidification tests. Tris buffered Leu-MCA and Arg-MCA substrate working
202 standards were made by diluting 500 μ l of MCA substrate stock (16 mM) with 4.5 ml of 0.1
203 M Tris/HCl buffer. Duplicate Tris/MCA substrate solutions were adjusted to pH 8.1 and 7.8
204 by adding 10 % HCl and the pH of duplicate 10 ml aliquots of coastal seawater was also
205 adjusted to pH 8.1 and 7.8. For each pH treatment, 250 μ l of Tris/MCA substrate solution
206 was added to 10 ml of seawater fixed at the corresponding pH. pH was recorded at room
207 temperature using a pH electrode as described above.

208

209 **2.4 Seawater acidification approach**

210 The influence of acidification technique on biotic parameters was investigated in two separate
211 experiments conducted under controlled temperature conditions in late summer (May 2013 -
212 trial 1) and in early spring (October 2013 - trial 2). Coastal seawater was first filtered through
213 a 15 μ m filter and then a 1 μ m inline cartridge filter. Three different methods were used to
214 acidify seawater to that predicted by the end of the century (pH 7.80) (IPCC, 2013): (A) acid
215 addition using 0.1 M HCl; (B) bubbling CO₂-Air gas mixture through an acid-washed

216 aquarium airstone, and (P) CO₂-Air gas mixture introduced through gas-permeable silicon
217 tubing (Tygon Tubing R-3603; ID 1.6 mm; OD 3.2 mm; Law et al, 2012). Treatment P was
218 acidified to a pH of 7.8 by the sequential application of 100 % synthetically produced CO₂
219 gas for 25 min, followed by 10 % CO₂ gas (in 20.8 % O₂ in N₂, BOC Gas Ltd) for 60 min at a
220 flow rate of < 26 ml min⁻¹. The initial use of pure and 10 % CO₂ gas made it possible to reach
221 the target pH within 3 h. Treatment B was acidified by bubbling seawater with 742 µatm CO₂
222 gas (in 20.95 % O₂ in N₂, BOC Gas Ltd) for 143 min at < 25 ml min⁻¹ to achieve the target
223 pH 7.80. The volume of 0.1 M HCl required to acidify treatment A to pH 7.8 (2.0 ml - trial 1,
224 3.1 ml - trial 2) was calculated based on the sample volume, DIC and alkalinity (pers. comm.
225 Dr K. Currie, NIWA/University of Otago) using an algorithm from Dickson et al. (2007). To
226 ensure a consistent rate of pH change across treatments, treatment B and A were adjusted to
227 match that of the slower treatment P (150 min), with the pH of each sample verified using a
228 pH electrode. Each treatment and an ambient seawater Control were then incubated in
229 triplicate in acid-washed milli-Q water-rinsed 4.3 Litre low-density polyethylene (LDPE)
230 cubitainers (ThermoFisher Scientific), without a headspace. pH was monitored throughout
231 each 96 h incubation (supplementary material S2 and S3), however No further pH
232 adjustment took place, ~~during the 96 h incubation.~~

233 Each cubitainer was housed in one of two identical perspex incubation chambers (1730 mm
234 long, 450 mm high by 325 mm deep), set at *in situ* ambient seawater temperature (15.1 °C -
235 trial 1, 15.5 °C - trial 2). Artificial light (700 - 900 µE m⁻² s⁻¹) was maintained in each
236 cubitainer through external fluorescent light banks (Philips TLD 36 W/840); neutral density
237 polycarbonate screening ensured light intensities were uniform between incubation chambers,
238 while adjustable timers ensured an automated diurnal 12 h light/dark cycle. Mixing of water
239 in each cubitainer was achieved using an inflating diaphragm positioned underneath each
240 cubitainer, with the inflation and collapse of the diaphragm under the weight of the sample
241 resulting in continual water mixing. Cubitainers were also manually removed and inverted
242 three times prior to each sampling. Time-zero sampling occurred after initial pH adjustment.
243 Assay fluorophore and substrate standard solutions were adjusted to treatment pH.

244 **2.4.1 Bacteria and picoplankton cell numbers**

245 Triplicate samples were collected in 2 ml Cryovials (Raylab Ltd) and frozen in liquid
246 nitrogen (Hall et al., 2004) for up to 12 weeks prior to analysis. Bacterial cell numbers were

247 determined by flow cytometry (FACSCalibur, Becton-Dickinson) following staining with
248 SybrGreenII (Invitrogen) (Lebaron et al., 1998), and count events were normalised to volume
249 using TruCount bead solution (BD Biosciences) (Button and Robertson, 1993). Total
250 eukaryotic picoplankton numbers ($< 2 \mu\text{m}$) were determined by fluorescence of chlorophyll
251 (wavelength 670 nm), phycoerythrin (585 nm), and phycourobilin (530 nm) as well as
252 forward light-scatter providing an estimate of cell size. Final count values were recorded as
253 cells ml^{-1} .

254 **2.4.2 Bacterial secondary production**

255 Potential bacterial secondary production (BSP) was measured using ^3H -leucine (^3H -Leu) of
256 high specific activity ($> 80 \text{ Ci mmol}^{-1}$, SciMed Ltd) in triplicate 1.7 ml samples. Following
257 the TCA precipitation and centrifugation methodology (Kirchman, 2001; Smith and Azam,
258 1992), ^3H -Leu incorporation was determined using a liquid scintillation counter (Tri-Carb
259 2910 TR) and converted to secondary production using a protein conversion factor (1.5 kg C
260 mol^{-1} leucine) (Simon and Azam, 1989). Cell-specific rates were calculated by dividing the
261 BSP rate by total bacterial cell numbers.

262 **2.4.3 Dissolved Inorganic Carbon and Total Alkalinity**

263 Pre-combusted 12 ml sample DIC vials (Labco Ltd) were triple rinsed with sample seawater
264 and filled, ensuring no air bubbles. One drop of saturated HgCl_2 was added to each DIC
265 sample, with storage at room temperature. DIC was determined using evolved CO_2 gas after
266 sample acidification on a Marianda AIRICA system, the accuracy of this method was
267 estimated to be $\pm 5 \mu\text{mol kg}^{-1}$, as determined by analysis of Certified Reference Material.
268 Alkalinity samples were collected by filling a 1 liter screw top bottle, and following the same
269 sample preparation and storage procedures as DIC above. Samples were later analysed by
270 potentiometric titration in a closed cell (Dickson et al., 2007) with an accuracy of $\pm 2 \mu\text{mol}$
271 kg^{-1} , also determined by analysis of Certified Reference Material.

272 **2.5 Statistical analysis**

273 Statistica v.10 (StatSoft Inc., USA) was used for basic graphics and descriptive statistics.
274 Data was tested for normality and equality of variance prior to statistical analysis. Data was
275 $\log(x+1)$ transformed due to the small sample size at each sampling point. Standard
276 hypothesis formulations were used for each Analysis of Variance (ANOVA), the null
277 hypothesis (H_0) was $\mu = 0$. The significance level of each test was $p \leq 0.05$. If H_0 was

278 rejected, a Tukey's HSD post-hoc analysis test was run to identify individual variable
279 responses.

280

281 **3 Results and discussion**

282 **3.1 Enzyme assay methodology**

283 MUF and MCA fluorescence was lower at pH 7.8 relative to pH 8.1, as previously reported
284 in soils (Niemi and Vepsäläinen, 2005). The fluorescence of the unbuffered MUF 2-
285 methoxyethanol at 40000 nM was 20 % higher at pH 8.1 than at pH 7.8 (t-test, $p < 0.05$),
286 while MUF Tris buffered fluorescence at 200 nM was ~~15-3.2~~ % higher at pH 8.1 (t-test, $p <$
287 ≥ 0.05 ; Table 1). MCA 2-methoxyethanol fluorescence at 40000 nM was 25% higher at pH
288 8.1 than fluorescence at pH 7.8 (t-test, $p < 0.05$), while MCA Tris buffered fluorescence at
289 200 nM was ~~9-1.7~~ % higher at pH 8.1 than at pH 7.8 (t-test, $p < \geq 0.05$; Table 1). Due to the
290 basicity of the MCA amino group, fluorescence intensity is less affected by pH and it has
291 been suggested that buffering is not required in seawater (Piontek et al., 2013; Endres et al.,
292 2014), whereas buffering of MUF has been reported (Mead et al., 1955; Piontek et al., 2010;
293 2013, Endres et al., 2013). ~~Our~~These results confirm that pH has a significant effect on
294 unbuffered MUF and MCA fluorescence ~~at both high and typical working concentrations,~~
295 ~~while and that 0.1 M Tris buffer minimises any pH effect at typical working concentration, and~~
296 ~~so fluorophore calibrations should be carried out at the same pH as the sample.~~

297 Although there is awareness of the effect of pH on fluorophore fluorescence (Mead et al.,
298 1955; Piontek et al., 2013; Endres et al., 2014), few studies consider the effect of fluorescent
299 substrate addition on seawater pH. ~~Due to the basicity of the MCA amino group, fluorescence~~
300 ~~intensity is less affected by pH and it has been suggested that buffering is not required~~
301 ~~(Piontek et al., 2013; Endres et al., 2014), whereas buffering of MUF has been reported~~
302 ~~(Mead et al., 1955; Piontek et al., 2010; 2013, Endres et al., 2013).~~ Immediately following the
303 addition of non-buffered Leu-MCA or Arg-MCA substrate to seawater at pH 7.95 or 7.70, pH
304 decreased by at least 0.05 units for each substrate, and remained significantly lower 30 mins
305 after addition when compared to time-zero pH (one-way ANOVA, $p < 0.05$). As both MCA
306 substrates are hydrochloride salts, addition resulted in a significant pH change, as previously
307 reported by Hoppe (1993). In tests ~~of~~adding -Tris--buffered MCA substrate solutions adjusted

308 | to seawater pH 7.8 and 8.1 to seawater at the same pH, the resulting pH change ranged from
309 | 0.003 to 0.03 units (± 0.001 SE). As the addition of buffer solution ~~reduced~~ minimised the pH
310 | change, both MCA substrates and fluorophores were subsequently produced using 0.1 M
311 | Tris/HCl, with pH adjusted to that of the respective experimental treatments and Control. In
312 | contrast to MCA, no statistically significant change in pH was recorded immediately
313 | following, or 30 mins after, addition of either α -MUF or β -MUF substrate to seawater at pH
314 | 7.95 or 7.70, indicating that these are neutral compounds. However, to eliminate possible
315 | bias, MUF substrates were also buffered using Tris/HCl.

316

317 **3.2 Seawater acidification**

318 | Having established that the analytical procedures for determining extracellular enzyme
319 | activity are affected by, and alter pH, the influence of acidification technique was then
320 | considered in two separate trials in different seasons. Overall, the experiments showed that
321 | different acidification techniques had significant effects on BG and LAP activity at select
322 | time points in both trials (Fig. 1), while the response of AG and AAP activity was variable
323 | with no consistent treatment response relative to the Control (~~data not shown~~ supplementary
324 | material S4, S5, and S6). Overall, BG and AG activity declined from time-zero to 96 hrs in
325 | the Control and treatments in trial 1, but were both significantly higher in the treatments
326 | relative to the Control from time-zero to 72 h, with BG activity approximately three-fold
327 | higher than AG activity (data not shown). Cell-specific BG activity was at least an order of
328 | magnitude higher in treatment B, P and A relative to the Control at time-zero (one-way
329 | ANOVA, $p < 0.05$) (Fig. 2), which is consistent with a direct effect of acidification (Piontek
330 | et al., 2013). Cell-specific BG activity was highest in treatment B from 24 h to 72 h by at
331 | least 14 % relative to treatment A and P (Fig. 1). In contrast to trial 1, cell-specific BG
332 | activity increased significantly throughout trial 2 (repeated measures ANOVA, $p < 0.05$). The
333 | opposing temporal trends between trials may signify seasonal differences in the response of
334 | glucosidase to OA, potentially reflecting differences in microbial community composition
335 | (Endo et al., 2013) or substrate availability (Morris and Foster, 1971). There was no
336 | significant difference in BG activity between treatments at time-zero in trial 2 (one-way
337 | ANOVA, $p > 0.05$) (Fig. 2), and BG activity was again highest in treatment B from 48 h,
338 | with activity at least 18 % higher relative to treatment P and A (Fig. 1). Bulk water LAP and
339 | AAP activity varied between treatments for trials 1 and 2. For example, both LAP and AAP

340 activity were highest in treatment P throughout trial 1, whereas LAP activity was highest in
341 treatment B from 72 h to 96 h in trial 2 (data not shown). Although cell-specific LAP activity
342 showed evidence of a response to acidification at select time points, there was no consistent
343 significant response ~~is was not significant in~~ throughout either trial (Fig. 1).

344 Although treatment B was only bubbled with gas mixtures for the pre-incubation period (143
345 mins), this had a greater effect on BG activity than in the other treatments, indicating
346 potential artefacts associated with bubbling. Bubbling may have ruptured picoplankton cells
347 or increased their susceptibility to viral lysis, leading to an increase in the release of labile
348 organic carbohydrates. This is potentially supported by the decline in total eukaryotic
349 picoplankton cell numbers in treatment B (trial 1 – 2.8×10^3 to 2.6×10^3 cells ml⁻¹, trial 2 –
350 1.7×10^3 to 1.3×10^3 cells ml⁻¹) in both trials (repeated measures ANOVA, $p < 0.01$). An
351 increase in enzyme activity would theoretically increase the availability of low molecular
352 weight organic substrate for bacterial assimilation, and may explain the significant increase in
353 bacterial cell numbers in treatment B relative to the Control at 96 h in both trials (one-way
354 ANOVA, $p < 0.05$) (Fig. 2). An increase in bacterial abundance in response to bubbling has
355 been previously reported by (Kepkay and Johnson, 1989) who suggested that surface DOC
356 coagulation facilitated by bubbling resulted in increased respiration and bacterial numbers. It
357 is possible that bubbling increased the abiotic coagulation of organic matter (Riley, 1963) and
358 formation of high molecular weight substrate such as transparent exopolymer particles
359 (Mopper et al., 1995; Passow, 2012; Schuster and Herndl, 1995; Zhou et al., 1998), which
360 could explain the elevated cell-specific BG activity (Fig. 1).

361 All acidification treatments had a significant negative effect on cell-specific BSP from 24 h to
362 48 h in trial 1 (one-way ANOVA, $p < 0.05$) (Fig. 3). During trial 2, cell-specific BSP was
363 significantly lower in treatments B and P when compared to the Control from 72 h to 96 h
364 (one-way ANOVA, $p < 0.05$), while BSP was twice as high in treatment A during this period
365 (Fig. 3). Although a clear treatment response was not observed in either trial, the low cell-
366 specific BSP in treatment B relative to the Control and treatment A at 96 h in trial 2 was
367 surprising as enzyme activity and bacterial cell numbers were elevated. Existing literature
368 also reports variable BSP responses to acidified conditions. Arnosti et al., (2011) and Teira et
369 al., (2012) detected no significant BSP response, while Grossart et al., (2006) detected an
370 increase, and Maas et al., (2013) and Siu et al., (2014) recorded a decrease in BSP rates with
371 increasing CO₂. As the same response was not observed in trial 1, it is possible that additional

372 indirect factors such as bacterial community composition or substrate type may have
373 influenced BSP under OA conditions (Piontek et al., 2013).

374

375 **4 Conclusions**

376 Artificial fluorogenic substrates have been used to investigate bacterial extracellular enzyme
377 activities in aquatic environments for decades (Hoppe, 1983; Somville and Billen, 1983).
378 Although the technique has several limitations, including that the artificial fluorogenic
379 substrate may not represent the naturally occurring substrate (Chróst, 1989), ~~and so that~~ the
380 observed activity only represents potential hydrolysis (Arnosti, 1996; Unanue et al., 1999),
381 the technique is rapid and easily applied in the field and most importantly, allows for a
382 standardised method for comparison of results in different OA studies. This study confirmed
383 that specific artificial fluorogenic substrates used to determine extracellular enzyme activity
384 ~~are affected by, or can alter, sample pH, and, and so consequently that~~ buffering is required,
385 particularly when used in OA research. Seawater acidification stimulated β -glucosidase
386 activity as previously reported (Piontek et al., 2010; Burrell et al., 2015), but the use of
387 different methodological approaches will result in may generate different variable results, and
388 may under- or overestimate certain parameters at the same time but not all parameters
389 equally. can influence the magnitude of this response. Simple aAcid addition does not
390 produce realistic seawater carbonate chemistry predicted in a future ocean (Riebesell et al.,
391 2010), and bubbling with CO₂ gas has a significant effect on β -glucosidase activity and
392 bacterial cell numbers, indicating ~~that there are~~ artefacts associated with bubbling. It should
393 be noted that these effects were observed in small-volume laboratory-scale experiments (<10
394 Litres), and may have less impact in larger-scale experiments. Although not all techniques
395 previously used to artificially adjust seawater pH were trialled- (Riebesell et al., 2010), the
396 results presented here ~~Nevertheless, the results~~ indicate that introducing CO₂-air gas mixtures
397 using gas permeable-silicon tubing the most is a robust highly an effective technique to for
398 investigate- investigating the response of bacterial processes to future OA conditions, that and
399 appears superior to tested- alternatives methods is CO₂-Air gas mixtures introduced using gas
400 permeable- silicone tubing. This approach should be considered for broader use in standardised
401 protocols for ocean acidification (Riebesell et al., 2010; Cornwall and Hurd, 2015) to achieve
402 robust meta-analyses and international inter-comparisons.

403

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410

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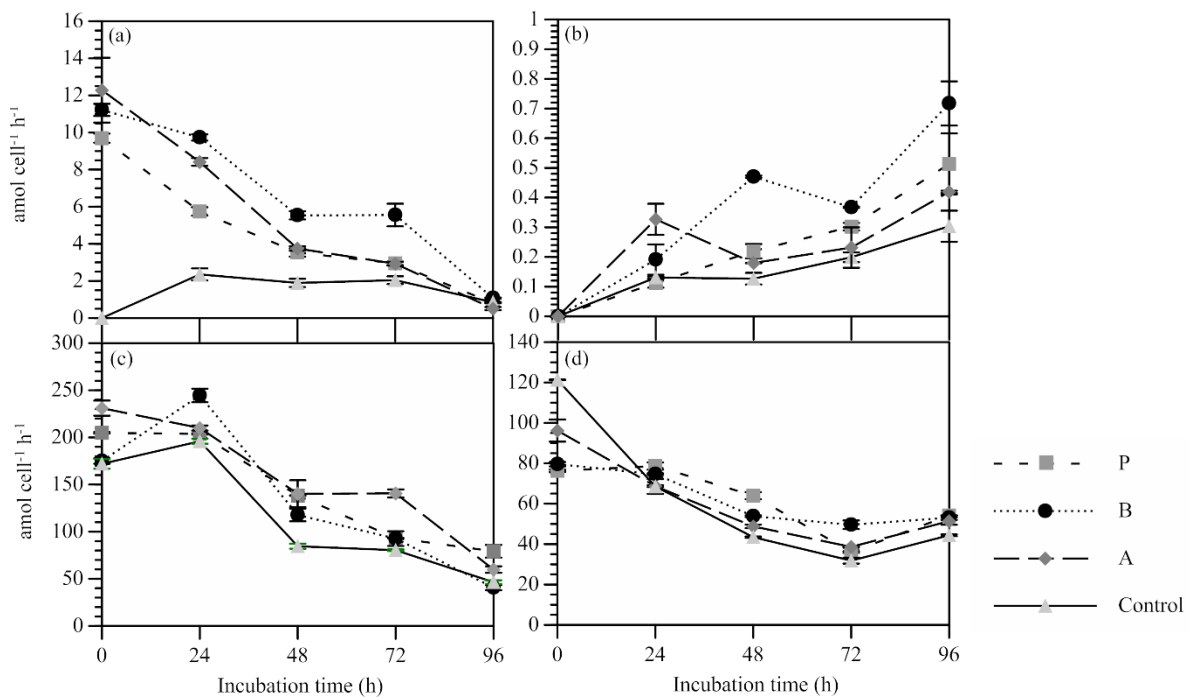
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647 **Figure Legends**

648 Table 1. Mean fluorophore fluorescence at pH 8.1 ad 7.8 (RFU, n=3, \pm SE).

Concentration (nM)	Fluorophore	pH 8.1	pH 7.8
0.1M Tris	MUF	1604.2421.44 (\pm 3.4317.86)	1553.18373.33 (\pm 38.412.49)
	MCA	13653.694948.90 (\pm 1518.052.52)	13420.72626.54 (\pm 2005.052.52)

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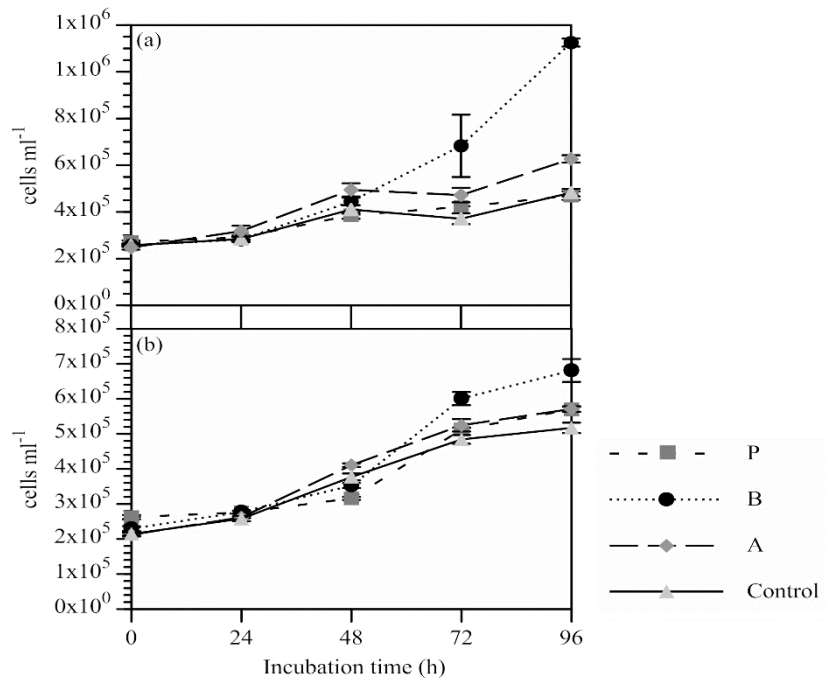
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651 Figure 1. Cell-specific extracellular enzyme activity (mean \pm SE, n=3) in response to
652 seawater acidified with 0.1 M HCl (A), bubbled with CO₂-Air gas mixture (B) and CO₂-Air
653 gas mixture introduced through gas-permeable silicon tubing (P). (a) BG activity in trial 1,
654 (b) BG activity in trial 2, (c) LAP activity in trial 1, (d) LAP activity in trial 2.

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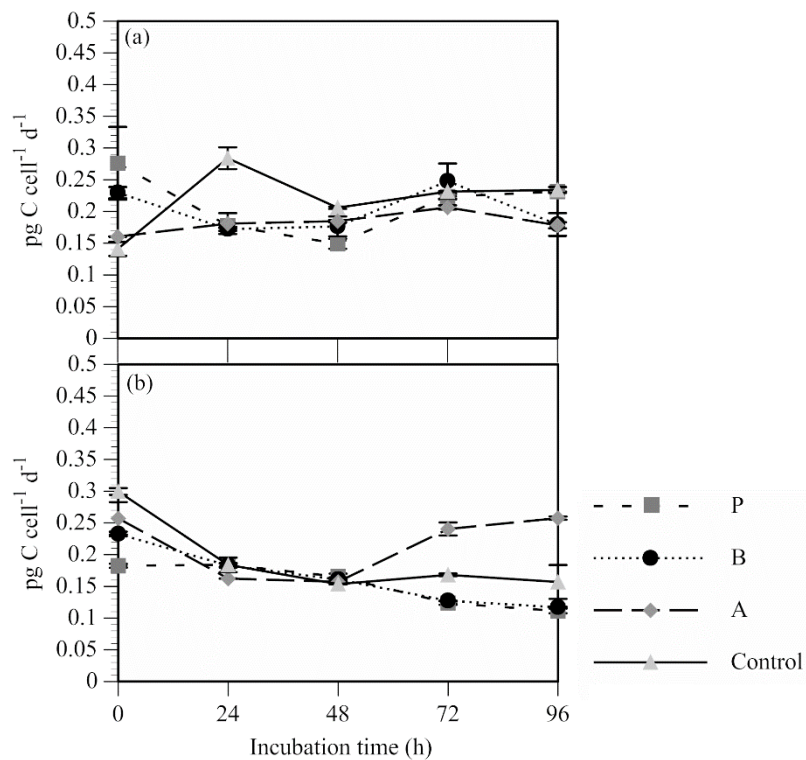
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659 Figure 2. Bacterial cell numbers (mean \pm SE, n=3) in response to seawater acidified with 0.1
660 M HCl (A), bubbled with CO₂-Air gas mixture (B) and CO₂-Air gas mixture introduced
661 through gas-permeable silicon tubing (P). (a) trial 1, (b) trial 2.



662

663 Figure 3. Cell-specific bacterial secondary production (mean \pm SE, n=3) in response to
664 seawater acidified with 0.1 M HCl (A), bubbled with CO₂-Air gas mixture (B) and CO₂-Air
665 gas mixture introduced through gas-permeable silicon tubing (P). (a) trial 1, (b) trial 2.