

1 **Climate sensitivity and the rate of ocean acidification: future impacts,**
2 **and implications for experimental design**

3 Matthew P. Humphreys. Ocean and Earth Science, University of Southampton, Waterfront
4 Campus, European Way, Southampton SO14 3ZH, UK.

5 Email: m.p.humphreys@soton.ac.uk. Tel.: +442380593641.

6 **Abstract**

7 The global mean surface temperature and partial pressure of carbon dioxide (CO₂) are
8 increasing both in the atmosphere and ocean. Oceanic CO₂ uptake causes a decline in pH
9 called ocean acidification (OA), which also alters other biologically-important carbonate
10 system variables such as carbonate mineral saturation states. Here, we discuss how a
11 ‘temperature buffering’ effect chemically links the rates of warming and OA at a more
12 fundamental level than is often appreciated, meaning that seawater warming could mitigate
13 some of the adverse biological impacts of OA. In a global mean sense, the rate of warming
14 relative to the CO₂ increase can be quantified by the climate sensitivity, the exact value of
15 which is uncertain. It may initially appear that a greater climate sensitivity would therefore
16 reduce the negative influence of OA. However, the dependence of the rate of CO₂ increase
17 on the climate sensitivity could enhance, nullify or even reverse the temperature buffering
18 effect, depending upon the future trajectory of anthropogenic CO₂ emissions. Regional
19 deviations from the global mean seawater temperature and CO₂ uptake trends could
20 modulate local responses to OA. For example, mitigation of OA impacts through
21 temperature buffering could be particularly effective in the Arctic Ocean, where the
22 surface seawater warming rate is greater than the global mean, and the aqueous CO₂
23 concentration might increase more slowly than elsewhere. Some carbonate system
24 variables are more strongly affected than others, highlighting the need to develop a

25 mechanistic understanding of precisely which variables are important to each
26 biogeochemical process. Temperature buffering of the marine carbonate system should be
27 taken into account when designing experiments to determine marine species and
28 ecosystem responses to warming and OA, in order that their results accurately reflect
29 future conditions, and therefore can generate realistic predictions when applied to Earth
30 system models.

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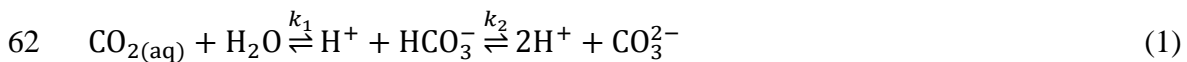
32 **Introduction**

33 The partial pressure of carbon dioxide in the atmosphere ($p\text{CO}_2$) is currently increasing
34 due to anthropogenic CO_2 emissions and land use change. Climate change is the most
35 well-known consequence, which has regionally varied impacts, along with an increase in
36 the Earth's global mean surface temperature (IPCC, 2013). About a quarter of each year's
37 CO_2 emissions dissolve into the oceans (Le Quéré *et al.*, 2009), where they react to
38 decrease the oceanic pH in a process commonly known as ocean acidification (OA)
39 (Gattuso and Hansson, 2011). Although this oceanic CO_2 uptake is beneficial from a
40 climatic perspective, OA may have consequences of its own for marine species and
41 ecosystems (Doney *et al.*, 2009), and for calcifiers in particular. Evidence from the
42 geological record suggests that previous hyperthermal events that were accompanied by
43 OA are often associated with declines in biogenic calcification and calcium carbonate
44 (CaCO_3) preservation, along with major extinction events for marine calcifiers (Hönisch *et*
45 *al.*, 2012). These past events are not perfect analogues for the present transient because
46 their rates of change were far slower, so recent research has focussed on identifying
47 biological responses to OA and warming in extant species (Hendriks *et al.*, 2010; Harvey
48 *et al.*, 2013).

49 A wide range of approaches have been taken to better understand the effects of OA and
50 warming (and other stressors) on biological systems. Recently there has been a drive
51 towards multi-stressor experiments, as it has been recognised that organisms under stress
52 from changes in one variable are likely to be less resilient to changes in another (Dupont
53 and Pörtner, 2013; Riebesell and Gattuso, 2015). For example, a species might be able to
54 tolerate a small change in temperature or a low level of OA, but not both together. Multi-
55 stressor experiments may also be more appropriate as future changes in seawater chemistry
56 and temperature will occur simultaneously.

57 **Temperature and carbonate chemistry**

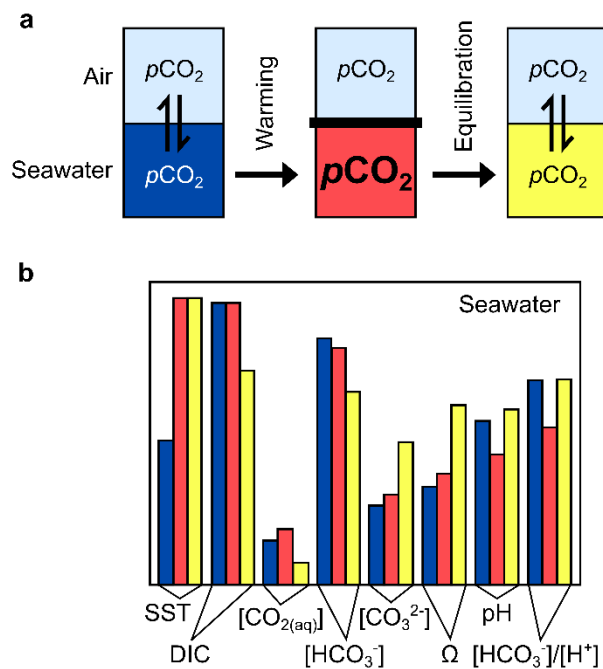
58 An additional complication that is often overlooked is that OA is directly regulated by
59 seawater temperature at a fundamental chemical level. After dissolving into seawater,
60 aqueous CO₂ (CO_{2(aq)}) reacts to form bicarbonate (HCO₃⁻) and carbonate (CO₃²⁻) ions, and
61 releases protons (H⁺):



63 These reactions continue in dynamic equilibrium, and the relative concentration of each
64 chemical species can be determined from the equilibrium constants k_1 and k_2 . These
65 constants have been theoretically (Millero and Pierrot, 1998) and empirically (e.g.
66 Mehrbach *et al.*, 1973) shown to be temperature-dependent. Temperature changes
67 therefore result in shifting of these equilibria, and redistributions of the concentrations of
68 these chemical species. This process is effectively instantaneous from an OA perspective,
69 as the equilibration timescale for the reactions in (1) is on the order of a few seconds or
70 faster (Eigen, 1964; Johnson, 1982).

71 We can perform a few simple calculations to investigate the potential consequences of this
 72 temperature dependency. First, consider a parcel of seawater that is cut off from CO₂
 73 exchange with the air. Warming this seawater will decrease its pH and [HCO₃⁻] while
 74 increasing its [CO_{2(aq)}] and [CO₃²⁻] (Fig. 1; the square brackets indicate the concentration
 75 of the enclosed species in micromoles per kilogram of seawater). These changes occur
 76 entirely as a result of re-equilibration of the reactions in (1) following temperature-induced
 77 changes to k_1 and k_2 . What are the consequences for OA? The answer depends on which
 78 variables are biologically important. Considering pH in isolation, it initially appears that its
 79 decline following warming constitutes an enhancement of OA. However, further
 80 investigation of other variables and an extension of the timescale for this thought
 81 experiment both suggest that we might more correctly consider it to be the opposite, in a
 82 biological context: temperature buffering of OA impacts.

83



84

85 Figure 1. Schematic illustration of warming and CO₂ equilibration thought experiments.
86 (a) Initially, a parcel of seawater (blue) is in equilibrium with the air. After warming, and
87 with no air-sea CO₂ exchange, the seawater pCO₂ has increased (red). The seawater then
88 re-equilibrates back to the pCO₂ of the air (yellow). (b) Changes in key variables during
89 each step of the thought experiments. The colours of the bars correspond to the equivalent
90 steps in Fig. 1a. The vertical axis scale is different for each variable, and the relative sizes
91 of the bars are schematic rather than accurate.

92

93 It is not fully understood which variables are of direct, mechanistic importance to marine
94 organisms, but many studies have focussed on CaCO₃ mineral saturation states (Ω),
95 particularly in the context of calcification (e.g. Langdon *et al.*, 2000; Riebesell *et al.*, 2000;
96 Orr *et al.*, 2005). These can be expressed as follows:

$$97 \quad \Omega = \frac{[\text{Ca}^{2+}][\text{CO}_3^{2-}]}{k_{\text{sp}}} \quad (2)$$

98 where k_{sp} is the ‘solubility product’ of the CaCO₃ mineral (usually calcite or aragonite).

99 Returning to our simple thought experiment, the increase in [CO₃²⁻] driven by warming
100 will increase Ω , which is considered to be biologically beneficial. The k_{sp} is also a function
101 of temperature (Mucci, 1983), but its effect on Ω is an order of magnitude smaller than that
102 of the simultaneous temperature-driven changes in [CO₃²⁻]. The overall effect of warming
103 alone is in an increase in both $\Omega_{\text{aragonite}}$ and Ω_{calcite} , even in this hypothetical situation where
104 there is no sea-air CO₂ flux (Fig. 1).

105 Of course, the assumption of zero air-sea CO₂ flux is unlikely to hold true in the surface
106 ocean, which is particularly important for the biological impacts of OA. As already
107 mentioned, warming increases [CO_{2(aq)}] relative to the total dissolved inorganic carbon

108 (DIC, the sum of the concentrations of $\text{CO}_{2(\text{aq})}$, HCO_3^- and CO_3^{2-}). This means that after
109 warming, the seawater $p\text{CO}_2$ ($p\text{CO}_2^{\text{sw}}$) is greater, which could drive a net sea-to-air CO_2
110 flux. The solubility of CO_2 is also inversely proportional to temperature (Weiss, 1974), so
111 warming drives a further increase in the $p\text{CO}_2^{\text{sw}}$ through temperature control of the
112 equilibrium constant k_0 :



114 This equilibrium is slower than those in (1), taking on the order of a year for a typical
115 surface ocean mixed layer (Zeebe and Wolf-Gladrow, 2001). Nevertheless, on the multi-
116 decadal timescale of OA we could assume that the global mean surface ocean will remain
117 roughly in equilibrium with the atmosphere, and at a similar level of disequilibrium
118 regionally (Takahashi *et al.*, 2009). Extending the previous thought experiment by
119 allowing the seawater CO_2 to re-equilibrate with a constant atmospheric value shows that
120 the solubility effect enhances temperature buffering of the OA impact on Ω further still. In
121 this case, warming results in a loss of DIC, and this combined with the repartitioning of the
122 carbonate species in (1) leads to decreasing $[\text{HCO}_3^-]$ and $[\text{CO}_{2(\text{aq})}]$, whilst pH and $[\text{CO}_3^{2-}]$
123 both increase (Fig. 1). The $[\text{CO}_3^{2-}]$ increase is an order of magnitude greater than in the
124 previous calculation where air-sea CO_2 exchange was restricted, so its effect on $\Omega_{\text{aragonite}}$
125 and Ω_{calcite} is proportionally greater (Fig. 1). Seawater warming in itself therefore acts to
126 buffer the OA impact on carbonate mineral saturation states.

127 In addition (or alternatively) to Ω , the ratio of $[\text{HCO}_3^-]$ to $[\text{H}^+]$ has been proposed as being
128 of biological importance, as it quantifies the balance between a growth substrate (HCO_3^-)
129 and an inhibitor (H^+) (e.g. Bach, 2015; Cyronak *et al.*, 2016). Applying the same thought
130 experiments to this compound variable as we did to Ω reveals a quite different result.

131 Warming in the absence of air-sea gas exchange results in a significant decline of

132 $[\text{HCO}_3^-]/[\text{H}^+]$ – this is OA enhancement, rather than buffering (Fig. 1). Taking into account
133 CO_2 solubility and permitting total equilibration with a constant atmosphere, we find that
134 temperature has little influence on $[\text{HCO}_3^-]/[\text{H}^+]$ at a present-day $p\text{CO}_2^{\text{sw}}$ of 400 μatm . The
135 potential for temperature buffering of OA impacts on $[\text{HCO}_3^-]/[\text{H}^+]$ is much smaller than
136 for Ω . We expect the patterns of future Ω and $[\text{HCO}_3^-]/[\text{H}^+]$ change to be different from
137 each other (Bach, 2015), particularly in hydrographically-complex continental shelf seas
138 (Fassbender *et al.*, 2016). This underlines the importance of establishing which variable is
139 mechanistically more important to marine organisms in order to accurately predict their
140 future response to OA. Either way, this uncertainty relates primarily to biological CaCO_3
141 formation; Ω is a key control on its dissolution (Morse *et al.*, 2007), so its strong
142 temperature buffering could still be important in some contexts .

143 These thought experiments illustrate the possible chemical responses to warming in the
144 two endmember situations: total and zero CO_2 equilibration with a constant atmosphere.
145 The real-world response should fall somewhere within this range, depending upon the air-
146 sea gas exchange rate. We do not provide exact values of the changes because they will
147 vary depending upon local conditions (e.g. total alkalinity and salinity), but the overall
148 patterns that we have described should be ubiquitous.

149 **Climate sensitivity**

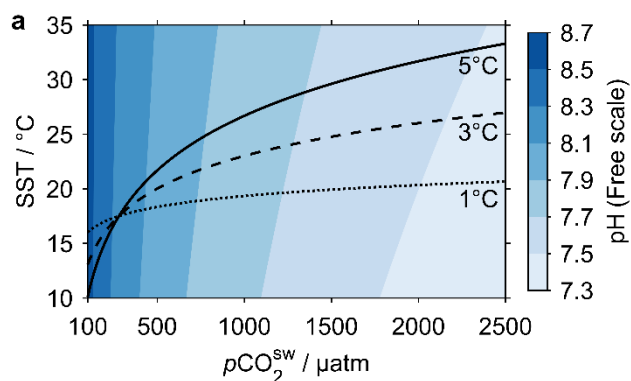
150 One further interaction has been excluded from our thought experiments so far: in the real
151 world, global mean sea surface temperature (SST) and $p\text{CO}_2^{\text{sw}}$ are increasing together. The
152 atmospheric $p\text{CO}_2$ is commonly linked to the global mean surface temperature through the
153 concept of climate sensitivity (or response) (e.g. Otto *et al.*, 2013; Marvel *et al.*, 2016).
154 The equilibrium climate sensitivity (ECS) is the change in the global mean near-surface air
155 temperature resulting from a doubling of atmospheric CO_2 , after the climate system has

156 been given time to re-equilibrate. Another definition of climate sensitivity that may be
157 more relevant to the present anthropogenic CO₂ increase is the transient climate response
158 (TCR), which is the warming that would occur after 70 years if the atmospheric CO₂
159 concentration rose by 1% per year throughout that period. The TCR is thought to be lower
160 than the ECS because of the relatively slow rate of ocean heat uptake (Winton *et al.*,
161 2009). In either case, the higher the ECS or TCR, the greater the increase in temperature
162 for a given increase in CO₂. Recent estimates suggest that the TCR is likely between 1.0
163 and 2.5°C, while the ECS is likely in the range from 1.5 to 4.5°C (Stocker *et al.*, 2013).
164 Over recent decades, the rate of global mean surface $p\text{CO}_2^{\text{sw}}$ increase has been similar to
165 that of atmospheric $p\text{CO}_2$ growth (Takahashi *et al.*, 2009), while SST has increased at
166 about 70-80% of the global mean surface temperature rate (Hartmann *et al.*, 2013). This
167 suggests that these climate sensitivity values should be decreased by 20-30% for our
168 purposes, assuming that these trends continue into the future.

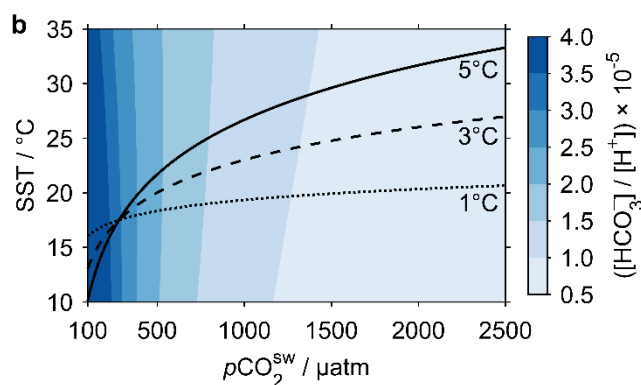
169 The climate sensitivity framework allows us to investigate how important temperature
170 buffering could be in a global mean sense, because a given ECS or TCR value defines a
171 specific trajectory through SST- $p\text{CO}_2^{\text{sw}}$ phase space. Assuming a constant total alkalinity
172 (TA), we can therefore calculate the future global mean state of the carbonate system as
173 $p\text{CO}_2^{\text{sw}}$ increases (Fig. 2). For pH (Fig. 2a) and $[\text{HCO}_3^-]/[\text{H}^+]$ (Fig. 2b), temperature
174 buffering is negligible at present day $p\text{CO}_2^{\text{sw}}$; the pH (or $[\text{HCO}_3^-]/[\text{H}^+]$) value is similar at
175 any given $p\text{CO}_2^{\text{sw}}$, regardless of the climate sensitivity trajectory. It increases at higher
176 $p\text{CO}_2^{\text{sw}}$, although it does not become important until very high $p\text{CO}_2^{\text{sw}}$ values are reached,
177 predicted beyond the year 2100. For Ω , temperature buffering is apparent throughout the
178 $p\text{CO}_2^{\text{sw}}$ range (Fig. 2c); the greater the climate sensitivity, the smaller the change in Ω for
179 a given increase in $p\text{CO}_2^{\text{sw}}$. In this simplistic ‘constant atmosphere’ scenario, a greater

180 climate sensitivity could therefore be considered beneficial for mitigating the biological
 181 impacts of OA.

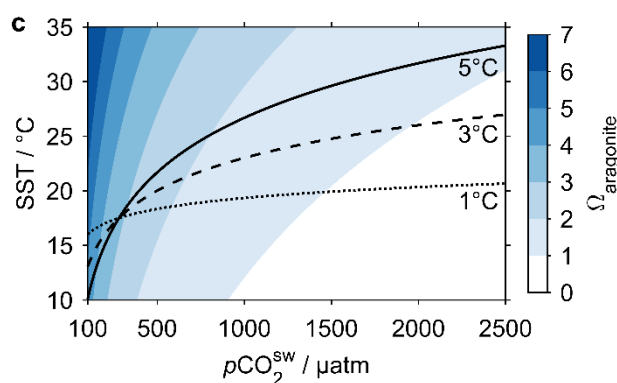
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185 Figure 2. The relationship between sea surface temperature, seawater $p\text{CO}_2$ and (a) pH, (b)
 186 bicarbonate ion to ‘proton’ ratio and (c) aragonite saturation state. The black lines indicate
 187 the trajectories of global mean SST and $p\text{CO}_2^{\text{sw}}$ for climate sensitivities of 1, 3 and 5°C,
 188 beginning from preindustrial global averages of 17.5°C and 280 μatm (Ahn *et al.*, 2012).
 189 In (c), they intersect with the $\Omega_{\text{aragonite}} = 1$ contour at $p\text{CO}_2^{\text{sw}}$ values of approximately 1460,
 190 1950 and 2940 μatm respectively (the latter is beyond the right hand edge of the plot),

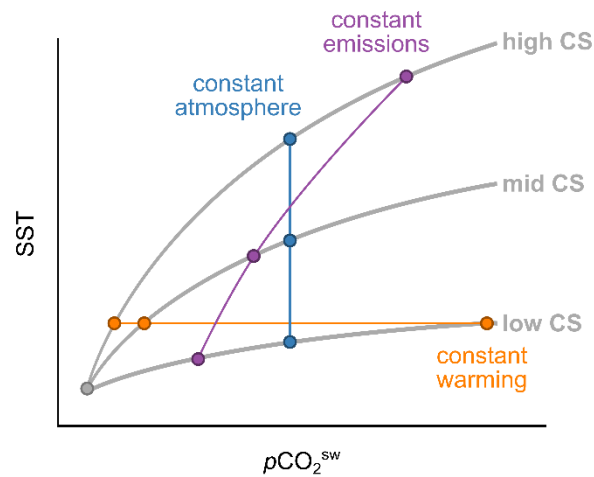
191 showing that the greater the climate sensitivity, the higher that $p\text{CO}_2^{\text{sw}}$ can rise before
192 aragonite becomes undersaturated. All CO_2 system variables were calculated using version
193 1.1 of the CO_2SYS program for MATLAB (van Heuven *et al.*, 2011), with dissociation
194 constants for carbonic acid following Lueker *et al.* (2000) and for bisulfate following
195 Dickson (1990), the boron:chlorinity of Lee *et al.* (2010). The contour fields were
196 calculated at a TA of $2250 \mu\text{mol kg}^{-1}$, practical salinity of 35, atmospheric pressure, and
197 zero silicate and phosphate.

198

199 However, this scenario assumes that the rate of $p\text{CO}_2^{\text{sw}}$ increase is independent of the
200 climate sensitivity, which is inaccurate. It effectively uses $p\text{CO}_2$ as a metric of passing
201 time rather than time itself, but it makes more sense to assess how quickly OA-related
202 changes might occur, rather than at what $p\text{CO}_2$ value they might occur. Warming of the
203 surface ocean decreases CO_2 solubility (Weiss, 1974), thus it increases $p\text{CO}_2^{\text{sw}}$ and reduces
204 the net air-sea CO_2 flux. Therefore under the same CO_2 emissions scenario, more CO_2
205 should remain in the atmosphere when the climate sensitivity is higher, so atmospheric
206 $p\text{CO}_2$ (and $p\text{CO}_2^{\text{sw}}$) will increase faster (Fig. 3). In terms of Fig. 2, this means that we
207 would move more quickly from left to right along the 5°C trajectory than along those for
208 lower climate sensitivities, thus opposing the temperature buffering. To approximately
209 quantify this effect, we considered a suite of emission-driven model simulations to the year
210 2100 (Fig. 12.36 from Collins *et al.*, 2013). These predicted a range of atmospheric $p\text{CO}_2$
211 values from about 800 to 1100 μatm , corresponding to global mean surface temperature
212 increase of about 2.5 to 6.5°C . Assuming that the highest temperature rise corresponds to
213 the highest $p\text{CO}_2$ increase and equivalently for the lowest, the $\Omega_{\text{aragonite}}$ would be about
214 0.16 units lower in the warmer case (calculated at the same conditions used in Fig. 1). For
215 comparison, if these two endmember levels of warming occurred at a constant,

216 intermediate $p\text{CO}_2$ value, then $\Omega_{\text{aragonite}}$ would be 0.24 units higher in the warmer case. In
 217 this ‘constant emissions’ scenario, temperature buffering of global mean Ω could therefore
 218 be overcompensated for by the faster $p\text{CO}_2^{\text{sw}}$ increase, thus enhancing the adverse changes
 219 to the marine carbonate system.

220



221

222 Figure 3. Schematic of total changes in sea surface temperature and CO_2 partial pressure
 223 after a set amount of time from the initial condition indicated by the grey circle in the
 224 bottom left corner, for different climate sensitivities (CS) and future emissions scenarios:
 225 ‘constant atmosphere’ assumes that the $p\text{CO}_2^{\text{sw}}$ increase is independent of the CS;
 226 ‘constant emissions’ assumes that anthropogenic CO_2 emissions will continue along a set
 227 trajectory, independent of the CS; and ‘constant warming’ assumes that emissions are
 228 decreased in response to warming such that a set temperature threshold is not exceeded.

229

230 The constant emissions scenario is more realistic than the original constant atmosphere
 231 scenario, but it still may not be an appropriate guide to the future. It assumes that the
 232 anthropogenic CO_2 emission rate is itself independent of the climate sensitivity, which

233 may not be true. Under a higher climate sensitivity, temperature increases and climatic
234 changes will occur more quickly, thus enhancing societal and political pressure to reduce
235 CO₂ emissions. Indeed, the recent Paris Agreement under the United Nations Framework
236 Convention on Climate Change commits to limiting the global mean surface temperature
237 increase to “well below 2°C above pre-industrial levels.” This warming threshold could be
238 approached more quickly if the climate sensitivity is greater, theoretically triggering a
239 greater and swifter reduction in anthropogenic CO₂ emissions, and leading to a ‘constant
240 warming’ scenario (Fig. 3). In this case, we would move more slowly from left to right
241 along the higher climate sensitivity trajectories on Fig. 2. This would be beneficial for all
242 of the variables in Fig. 2, although the reduction of OA impacts is a consequence of
243 decreased emissions, rather than being a direct effect of temperature buffering.

244 In summary, although a warmer temperature at a given $p\text{CO}_2^{\text{sw}}$ will lead to a more
245 biologically-favourable carbonate system, we cannot be certain whether a higher climate
246 sensitivity would have a chemical benefit without first knowing which path we will take
247 on the spectrum between the constant emissions and constant warming scenarios.

248 **Arctic Ocean**

249 Although most oceanic regions do not have the same conditions as the global mean, the
250 same principles still apply: the trajectory through SST- $p\text{CO}_2^{\text{sw}}$ phase space determines the
251 magnitude of the impact of OA. If SST and $p\text{CO}_2^{\text{sw}}$ in any particular region follow a
252 different trajectory to the global mean, for example due to changing circulation and
253 currents, then that region could be said to have its own local climate response (LCR) that
254 controls the importance of temperature buffering therein.

255 A key region that deviates from the global mean is the Arctic Ocean. The Arctic Ocean is
256 presently an important sink for atmospheric CO₂, despite the limitations on air-sea gas

257 exchange imposed by seasonal sea-ice cover (Bates and Mathis, 2009). However, it has
258 experienced a significant decline in sea ice in recent years (Perovich and Richter-Menge,
259 2009; Perovich *et al.*, 2015), which has probably been mainly driven by atmospheric
260 warming (Screen and Simmonds, 2010). The albedo reduction associated with sea-ice loss
261 also acts as a positive feedback, further accelerating this warming (Perovich *et al.*, 2007).
262 Significant heat fluxes into the surface ocean have also been driven by inputs of relatively
263 warm water from the North Atlantic (Gerdes *et al.*, 2003) and turbulent vertical mixing
264 generated by tides (Rippeth *et al.*, 2015). As a result of these processes, the Arctic Ocean
265 has been warming faster than anywhere else on the planet, with surface SST in August
266 2015 reaching up to 4°C higher than the 1982-2010 mean (Timmermans and Proshutinsky,
267 2015). Consequently, the LCR of the Arctic Ocean should be greater than the global TCR,
268 as it has experienced more warming for a given increase in atmospheric $p\text{CO}_2$. However,
269 this enhanced warming is only half of the story.

270 Net primary production (NPP) in the Arctic Ocean rose by 30% between 1998 and 2012,
271 partly as a result of the reduced sea ice extent (Arrigo and van Dijken, 2015), and this
272 trend is likely to continue into the next few decades (Yool *et al.*, 2015). Indeed, 2015 saw
273 widespread high chlorophyll-*a* anomalies across the region (Frey *et al.*, 2015). There are
274 two ways in which this could act to further increase the Arctic LCR. Firstly, in an
275 extension of the well-known sea ice-albedo positive feedback (Perovich *et al.*, 2007),
276 seawater containing high chlorophyll-*a* and biomass has increased heat absorption relative
277 to oligotrophic seawater, so the enhanced NPP could further contribute to local warming
278 (Park *et al.*, 2015). Secondly, it may also decrease $p\text{CO}_2^{\text{sw}}$ through conversion of DIC into
279 organic matter, thus increasing the LCR even more.

280 From a chemical perspective, these factors (enhanced warming and mitigation of the
281 $p\text{CO}_2^{\text{sw}}$ increase) both increase the localised chemical impact of temperature change. In the

282 Arctic Ocean, the potential for temperature buffering to counteract the chemical
283 consequences of ocean acidification could therefore be greater than the global mean,
284 regardless of the climate sensitivity.

285 **Experimental design**

286 Several studies have noted that OA experiments need to become more complex, moving
287 from assessing single species responses to those of entire ecosystems, from single to
288 multiple stressors, and from short-term acclimative to long-term adaptive responses, as
289 well as assessing all life-cycle stages (Dupont and Pörtner, 2013; Riebesell and Gattuso,
290 2015). It has also been pointed out that as the surface ocean is frequently out of
291 equilibrium with atmospheric CO₂, it may not be appropriate to use global mean present
292 day atmospheric $p\text{CO}_2$ as a baseline for these studies, but rather to use conditions that are
293 more relevant to the natural environment (McElhany and Busch, 2013), and to take into
294 account natural co-variation of variables like SST and $p\text{CO}_2^{\text{sw}}$ (Reum *et al.*, 2015). In
295 addition, the interactions between carbonate chemistry and temperature discussed here
296 should also be considered in order to design experiments that simulate realistic future
297 ocean conditions.

298 For example, a single-stressor experiment that aims to change only temperature could also
299 unintentionally affect the carbonate system, while an experiment that alters the carbonate
300 system at constant temperature could effect greater chemical changes than are likely to
301 occur in the real world. For the former case, consider an experimental set-up of different
302 incubation containers that are kept at different temperatures, but all open to the same air.
303 The $p\text{CO}_2^{\text{sw}}$ would probably be similar in each container due to ongoing equilibration, so
304 the $\Omega_{\text{aragonite}}$ would be higher in the warmer containers (Fig. 1). If an $\Omega_{\text{aragonite}}$ decline
305 would be detrimental to the organism in question, then any negative impact of increased

306 temperature might be reduced somewhat by the inadvertent increase in $\Omega_{\text{aragonite}}$. This could
307 be countered by chemically manipulating the CO_2 system to ensure constant $\Omega_{\text{aragonite}}$ at the
308 different temperature conditions, but this manipulation would have knock-on effects for
309 other CO_2 system variables. It is well-established that all of the CO_2 system can be
310 calculated if any two of its variables are known (Zeebe and Wolf-Gladrow, 2001), but this
311 also means that only a maximum of two variables can have specific values selected as
312 manipulation targets. It may therefore be impossible to hold all of the CO_2 system
313 variables that might affect biological responses constant across a range of temperature
314 targets, rendering this possible only if the variable directly driving the biological response
315 is known. The simplest method to ensure realistic experimental results could therefore be
316 to increase both $p\text{CO}_2^{\text{sw}}$ and temperature together in line with the anticipated LCR. Indeed,
317 the results of a multi-stressor experiment that accurately reflected the future conditions in
318 the region of interest could be used in a predictive capacity without needing to definitively
319 establish which variables were mechanistically responsible for the observed biological
320 response.

321 **Using experimental results in models**

322 In any Earth system model that includes greenhouse warming and an accurate
323 representation of the marine carbonate system, temperature buffering of carbonate system
324 variables will already be taken into account (e.g. $\Omega_{\text{aragonite}}$ will automatically decrease more
325 slowly in regions where the SST increase is higher). The important step for future
326 experimental work is to ensure that the biological responses to OA are being measured in
327 appropriate conditions, in order to feed into models and accurately predict global-scale
328 responses.

329 For example, McNeil et al. (2004) used equations for the dependence of coral reef
330 calcification on temperature (Lough and Barnes, 2000) and $\Omega_{\text{aragonite}}$ (Langdon *et al.*, 2000)
331 to predict future changes in coral reef calcification in a coupled atmosphere-ice-ocean
332 carbon cycle model (Hirst *et al.*, 1996). These temperature- and $\Omega_{\text{aragonite}}$ -dependence
333 studies effectively considered these variables in isolation, which could cause problems for
334 the later model study. If we make the simplifying assumption that the temperature study
335 was carried out at constant TA and $p\text{CO}_2^{\text{sw}}$, then the $\Omega_{\text{aragonite}}$ would not have been constant
336 (Fig. 1). The temperature range considered by Lough and Barnes (2000) was about 23-
337 29°C. Assuming total equilibration at a $p\text{CO}_2$ of 400 μatm and a constant total alkalinity of
338 2250 $\mu\text{mol kg}^{-1}$, this corresponds to an increase in $\Omega_{\text{aragonite}}$ of 0.63 units. Using the
339 equation of Langdon et al. (2000), this increase in $\Omega_{\text{aragonite}}$ should correspond to an
340 increase in calcification rate by approximately 26.1 $\text{mmol-CaCO}_3 \text{ m}^{-2} \text{ day}^{-1}$ (assuming that
341 $\Omega_{\text{aragonite}}$ is the actual driver of calcification rate change, rather than $[\text{HCO}_3^-]/[\text{CO}_3^{2-}]$). This
342 could then be subtracted from the actual change in calcification from 23 to 29°C observed
343 by Lough and Barnes (2000) of 542 $\text{mmol-CaCO}_3 \text{ m}^{-2} \text{ day}^{-1}$ to arrive at the temperature-
344 only influence. This reduces the temperature-dependence of calcification by about 5%, so
345 McNeil et al. (2004) may have overestimated the warming-driven increase in calcification
346 by this amount. This relatively small adjustment would not alter the main conclusions
347 drawn by McNeil et al. (2004), but this might not be the case for other species or
348 ecosystems that are relatively less sensitive to warming and more responsive to $\Omega_{\text{aragonite}}$
349 changes. This uncertainty could be eliminated through proper consideration of temperature
350 buffering during the design of single- and multi-stressor experiments.

351 **Conclusion**

352 There is a temperature buffering effect through which the marine carbonate system is
353 altered in a fashion that is beneficial to biology upon seawater warming. The future rate of

354 warming relative to atmospheric CO₂ growth – the climate sensitivity – will therefore
355 modulate the chemical changes to the marine carbonate system caused by oceanic CO₂
356 uptake, but the global mean consequences will depend on the pattern of future CO₂
357 emissions. Different carbonate system variables (e.g. $\Omega_{\text{aragonite}}$ and $[\text{HCO}_3^-]/[\text{H}^+]$) will
358 respond differently from each other, so it is essential to further develop a mechanistic
359 understanding of precisely which variable influences each biogeochemical process of
360 interest. Some areas (e.g. the Arctic Ocean) are diverging significantly from the global
361 mean warming and $p\text{CO}_2^{\text{sw}}$ trends, causing the capacity for temperature buffering to
362 mitigate OA impacts to vary regionally. These principles should be considered when
363 designing experiments to assess the responses of marine species and ecosystems to
364 ongoing OA and warming, in order that the results of these experiments can be applied to
365 model simulations and return realistic predictions.

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