

Balancing of Ocean Acidification by Superoxide Redox Chemistry?

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cean acidification is typically caused by production of carbonic acid (H_2CO_3) through the dissolution of increasing atmospheric $CO₂$, which adds to $CO₂$ plus DIC (dissolved CO_2 , H_2CO_3 , HCO_3^- , and CO_3^2) produced in seawater by several processes including biological ones (primary production followed by respiration).1−³ Acidification can decrease the saturation states of carbonate minerals, which may considerably endanger the dynamics[, st](#page-1-0)ructure and biodiversity of coral reefs and other marine calcifying organisms.^{2,3} The consequence is the decline in the early development stages of shellfish, coral reefs, or other marine calcifiers, [wit](#page-1-0)h impacts on fertilization, sexual reproduction, cleavage, larval settlement, survival and growth, finally causing a substantial population decline.² Decline in shellfish or coral reefs, which form the foundation of marine ecosystems, would markedly affect the whole foo[d](#page-1-0) webs and marine population dynamics.²

Despite the apparent straightforwardness of such a scenario, the actual [d](#page-1-0)etermination of the impact of atmospheric $CO₂$ on seawater pH is a difficult task. This happens primarily because of the relatively small pH changes, unless very long time spans are taken into account. Moreover, pH modifications can also take place at a local scale because of several possible confounding factors, 1 of which we will show the potential role of superoxide chemistry. Among additional causes for seawater pH variatio[n](#page-1-0), one is the acidification connected with eutrophication phenomena that have increased worldwide since the last few decades. Indeed, changes in land-use practices can induce the release of high amounts of nutrients and terrestrial organic matter (OM) to coastal seawaters, including dissolved OM (DOM) and particulate OM (POM).^{1,2} The transformation of organic P and N into phosphate and nitrate because of microbial or photochemical proces[ses](#page-1-0) can cause a significant decrease of the alkalinity of coastal seawater and, therefore, a decrease in pH.² Another possibility (although limited in space and time) is the elevated production of $CO₂$ and DIC during harmful alga[l](#page-1-0) blooms, which can significantly alter the pH of seawater as long as they are operational.¹ Eutrophication phenomena and algal blooms can both be enhanced by global warming, which causes an increase [of](#page-1-0) surface seawater temperature and leads to a longer summer stratification period. In warm, sunlit surface seawater the photochemical and biological processes that are involved in the degradation of DOM and POM can be strongly enhanced, leading to increased production of $CO₂$ and of anionic species that decrease the alkalinity of seawater.² The third possible issue is connected with atmospheric acid rain, most notably i[n](#page-1-0)volving $HNO₃$ and $H₂SO₄$ that can directly lower the seawater $pH²$ All such processes are usually limited to the coastal areas that are most affected by eutrophication phenomena, [w](#page-1-0)hile acid rains are not expected to have a comparatively important impact on the pH of the open ocean.

Within this context, an additional factor of potential importance is represented by the chemistry of superoxide. The radical ion O_2^- is produced by various sources including extracellular generation by heterotrophic bacteria that are commonly detected in lakes, soil, hydrothermal vents, marine sediments, estuaries, and oceans.⁴ Rates of superoxide production normalized to the proportion of metabolically active cells are fou[n](#page-1-0)d to vary between 0.02 ± 0.02 amol cell⁻¹ hour⁻¹ (mean \pm standard error) and 19.4 \pm 5.2 amol cell⁻¹ hour^{-1.4} Such production rates could lead to an alteration of . seawater pH because of H^+ consumption upon dismutation of supero[xi](#page-1-0)de $(2 O_2^{-\bullet} + 2H^+ \rightarrow H_2O_2 + O_2)$. The latter process is catalyzed by redox-active metals such as $Cu(I/II)$ and $Fe(II/$ III) and possibly (but still controversially) by organic compounds with redox-capable moieties.⁵ With a biological generation rate of O_2 ^{-•} prudentially assumed to be 0.2 pmol dm⁻³ s⁻¹, which is only 10% of that rep[or](#page-1-0)ted by Diaz et al.,⁴ based on superoxide dismutation alone one can have a H⁺ consumption rate of ~6 μ mol [d](#page-1-0)m⁻³ year⁻¹. A simplified seawater model shows that the process would be able to compensate for an acidification rate of $0.1-0.3$ pH units year⁻¹

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(Figure 1), which is far higher than the observed acidification rate. The dismutation of superoxide has thus potential to

Figure 1. Equilibrium pH of seawater as a function of the concentrations of bicarbonate and added H⁺. .

significantly impact the pH of seawater, but it is clear that before drawing any firm conclusion over this issue it is mandatory to carry out a complete assessment of the charge balance of the whole process. The complete charge balance involves: (i) O_2 ^{-•} generation *via* both (photo)chemical and biological processes, in particular because the initial production of HO_2^{\bullet} would compensate for the H⁺ consumption in $O_2^{\bullet-\bullet}$ dismutation; (ii) the relative role of O_2 ^{-•} dismutation vs. scavenging upon oxidation $(O_2^{-\bullet} \rightarrow O_2 + e^-)$ or reduction $(O_2^{-\bullet} + e^- + 2 H^+ \rightarrow H_2O_2)$ reactions, which entail a different number of exchanged H^+ ions; and (iii) the possible involvement of H^+ in the semireactions of the redox couples (metal species and complexes, organic compounds and so on) reacting with O_2 ^{-•} and related oxygen species. A full understanding of all the processes (both major and secondary ones) of O_2 ^{-•} production and scavenging is vital to get insight into seawater acidification as well as redox state. Indeed, although the H⁺ balance of HO_2^{\bullet} generation and $O_2^{\bullet-\bullet}$ dismutation closes to zero, a small local prevalence of additional processes such as superoxide oxidation or reduction could modify seawater pH, because the chemistry of superoxide is potentially able to exchange much more H^+ ions than those involved in dissolution of atmospheric $CO₂$.

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Notes

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