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The impacts of ocean acidification on marine trace gases and the implications for atmospheric chemistry and climate

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Surface ocean biogeochemistry and photochemistry regulate ocean–atmosphere fluxes of trace gases critical for Earth’s atmospheric chemistry and climate. The oceanic processes governing these fluxes are often sensitive to the changes in ocean pH (or $pCO_2$) accompanying ocean acidification (OA), with potential for future climate feedbacks. Here, we review current understanding (from observational, experimental and model studies) on the impact of OA on marine sources of key climate-active trace gases, including dimethyl sulfide (DMS), nitrous oxide (N$_2$O), ammonia and halocarbons. We focus on DMS, for which available information is considerably greater than for other trace gases. We highlight OA-sensitive regions such as polar oceans and upwelling systems, and discuss the combined effect of multiple climate stressors (ocean warming and deoxygenation) on trace gas fluxes. To unravel the biological mechanisms responsible for trace gas production, and to detect adaptation, we propose combining process rate measurements of trace gases with longer term experiments using both model organisms in the laboratory and natural planktonic communities in the field. Future ocean observations of trace gases should be routinely accompanied by measurements of two components of the carbonate system to improve our understanding of how in situ carbonate chemistry influences trace gas production. Together, this will lead to improvements in current process model capabilities and more reliable predictions of future global marine trace gas fluxes.

1. Introduction

The interface between the ocean and the atmosphere is a crucial boundary of the Earth system. It controls not only the exchange of substances which influence the chemistry of the atmosphere and our climate, but also the transfer of essential elements vital for human health and ecosystem functioning, from the ocean to the land. The Earth system is currently facing unprecedented changes in global biogeochemical and physical processes, driven by human emissions of greenhouse gases [1]. In this review, we focus on one such change, ocean acidification (OA), and assess its impact upon the production of marine trace gases and resulting feedbacks to the atmosphere. We discuss the role of marine trace gases in the Earth system’s chemistry and climate, and provide an overview of the state-of-the-knowledge of the marine trace gas response to OA derived from both experimental and modelling studies. In addition, we consider regions especially sensitive to OA, and discuss the effects of other environmental changes, such as rising temperatures and ocean deoxygenation, on the production and emission of marine trace gases.

2. Marine trace gases

The surface ocean is a key source of a variety of trace gases, which flux to the atmosphere and play critical roles in the Earth’s biogeochemical cycles, and strongly influence the chemistry of its atmosphere and its radiative budget. These include greenhouse gases, such as carbon dioxide (CO$_2$), nitrous oxide (N$_2$O) and methane (CH$_4$), that have relatively well-understood effects on
global radiative forcing and atmospheric chemistry [2]. In addition, the ocean releases a range of biogenic volatile organic compounds (BVOCs), containing carbon, sulfur, nitrogen and halogens (figure 1). The transfer of these compounds from ocean to land via the atmosphere represents a key step in the global cycling of essential elements that provide benefits to ecosystem function and human health [3]. Furthermore, the atmospheric oxidation products of some trace gases, such as dimethyl sulfide (DMS), methylamines and a variety of BVOCs, can impact upon marine aerosols, thereby influencing cloud-related processes and global radiative forcing [4–8]. Other marine trace gases, including halocarbons and oxygenated volatile organic compounds (OVOCs), produce highly reactive atmospheric radicals that readily destroy protective stratospheric ozone (O\(_3\)), and drive the rapid cycling of tropospheric photo-oxidants and O\(_3\) with implications for coastal air quality [9–12]. Biogenic marine trace gases are directly produced by micro- and macro-algae, and by prokaryotic microbes [13–15]. They are also released from sediments [16], seafloor seeps [17,18], as a result of bacterial degradation of precursor compounds [19–22], and via reactions between organic matter, sunlight and O\(_3\) [23] (figure 1). Whereas the sources and sinks of CO\(_2\), N\(_2\)O and DMS, are reasonably well established, others remain poorly understood. Given the critical role marine trace gases play in atmospheric chemistry and climate-related processes, it is important to consider the influence of global environmental change on their oceanic flux, and associated feedbacks to climate.

3. CO\(_2\)-driven ocean acidification

Anthropogenic CO\(_2\) emissions from burning fossil fuels and land-use change are currently the primary driver of global climate change [24]. Atmospheric CO\(_2\) concentrations have steadily risen over the last 150 years and are now higher than at any time during at least the last 800 000 years of Earth’s history [25,26]. This rise directly results in increased oceanic CO\(_2\) absorption
In addition to increased hydrogen ion concentration (H\(^+\)), and hence decreased pH, associated chemical changes include increased concentrations of bicarbonate ions (HCO\(_3^-\)) and reductions in carbonate ions (CO\(_3^{2-}\)) [27,28]. Globally, a decrease in surface ocean pH of approximately 0.1 units has already occurred relative to preindustrial times, with a projected fall of a further approximately 0.3 units by 2100 under high-emission scenarios [1,29,30]. Sustained ocean observations from seven globally distributed time-series stations, including the northerly Iceland and Irminger Seas, the subtropical Bermuda Atlantic Time-series Study (BATS) and the tropical Hawaiian Ocean Time-series (HOT) show a 0.013–0.025 pH unit per decade decline since the 1980s [31]. This rate of change to ocean biogeochemistry is rapid on geological timescales and is probably unprecedented in the last 300 million years of Earth’s history [32]. Most OA research has focused on potential effects on calcifying organisms (e.g. [33–37]) and other ecologically and economically important species [38–41]. To date, there has been little assessment of OA impacts upon marine biogeochemical cycles that potentially involve changes in the production of marine trace gases and associated feedbacks on atmospheric chemistry and climate [42–45]. Given the documented sensitivity of marine microbes to a variety of environmental factors (e.g. seawater chemistry, temperature; [46]), there is clearly potential for climatically significant future change in the production and emissions of trace gases. Ultimately, OA will act in concert with global warming and modify the physical–chemical environment of pelagic and benthic communities. This in turn is expected to trigger changes in species composition of micro- and macroalgal communities [47–51]. Shifts in the community composition, or any increase in stress arising from OA, may alter the production of trace gases and the geographical pattern of trace gas emissions.

4. Biological processes drive trace gas production

We focus on biogenic marine trace gases (excluding CO\(_2\)) that are directly and/or indirectly produced by bacteria, phytoplankton and seaweeds, as well as trace gases produced by reactions involving dissolved organic matter (DOM). Given the known and predicted effects of OA on biological processes [52], it is likely that the net production of biogenic trace gases (including both production and loss processes) may be influenced by OA. First, we provide a general overview of the potential effects of OA on biological processes related to marine trace gas production, while later we discuss specific trace gases in greater detail.

Marine bacteria drive essential biogeochemical processes, including organic matter decomposition, elemental cycling and nutrient regeneration, and are key players in the production and consumption of marine trace gases [53,54] (figure 1). While we have a reasonable understanding of the influence of temperature and organic matter availability on bacterial processes, the role of OA is less well constrained [46,55]. The majority of studies imply that bacterial processes are seemingly resilient to reductions in pH due to the rapid microbial cell division and the associated adaptive potential that this affords, but our mechanistic understanding of this, and how it may influence trace gas production, is limited [56–58]. Recent work has shown that pH-induced changes in bacterial physiological processes affect cellular energy allocation, thereby influencing fluxes of carbon and energy in microbial systems [59]. In addition, OA may lead to increased exudation of dissolved organic carbon by phytoplankton, altering substrate availability and form [60] and enhancing extracellular enzyme activity [61–63]. Such changes have the potential to indirectly influence trace gas levels by altering the availability of precursors in the dissolved organic carbon pool, and by influencing the rate of bacterial processes that both produce and consume trace gases, and that ultimately result in their net production.

Many marine photosynthesizers, including single-celled phytoplankton and macrophytes such as seaweeds, are a direct source of marine trace gases, including DMS [64] and certain halocarbons, such as methyl iodide (CH\(_3\)) and bromoform (CHBr\(_3\)) [65,66]. DMS production by algal cells may be a means to release excess carbon and sulfur via a metabolic overflow mechanism [20,67], or alternatively, it may provide protection against physiological stress [68,69].
Similarly, halocarbon production has been attributed to relief from stressors including light-induced [70], oxidative [71] or mechanical [13] stress. In general, the biosynthesis of trace gases and their precursors is poorly understood, but is likely to vary substantially between species. For example, intracellular concentrations of dimethylsulfoniopropionate (DMSP), the algal precursor for DMS, have been shown to vary by up to three orders of magnitude across phytoplankton groups within a population [72].

Shifts in phytoplankton community composition in response to OA might alter global DMS emissions. For example, coccolithophores, a relatively well studied and ubiquitous group of marine calcifying haptophytes, possess high levels of intracellular DMSP, with blooms of this species associated with the release of vast quantities of DMS [64]. Their fate under OA has been the target of numerous experimental studies and field surveys over the past 20 years. However, the direction and magnitude of the response to OA varies substantially [33,73–75] and is still poorly understood. This underlines our still limited process understanding, which continues to hinder our ability to anticipate changes in DMS emissions under future climate change.

5. Experimental evidence: exploring effects of ocean acidification on marine trace gases

Our knowledge of the effects of OA on marine trace gas production stems from the results of a suite of experimental approaches, summarized in table 1. At the simplest level, on an experimental spectrum of complexity, are incubations with single-species algal cultures (less than 11, 2–3 replicates and 7–40 days). This is an approach which, given the reduced complexity, serves as a means to establish baseline concepts and identify the most sensitive or relevant physiological processes and mechanisms for trace gas production. Studies have considered ambient CO₂ versus one high CO₂ treatment: 370–395 µatm compared with 750–1000 µatm, corresponding to pH treatments of 8.1–8.3 (ambient) and 8.0–7.7 (High CO₂) (table 1). Of greater complexity and closer to actual ocean conditions are in situ mesocosm experiments, essentially giant ‘test tubes’ that allow large-scale, field-based, community-level assessments of the effects of OA on natural surface ocean communities (2400–75 000 l, 1–3 replicates and 25–35 days). Mesocosms provide an understanding of the net effects on the whole community response to OA, in many cases investigated under conditions of high productivity and growth associated with a phytoplankton bloom. Earlier experiments considered two triplicated CO₂ treatments (‘Ambient’ CO₂ 300–400 µatm, pH 8.2–8.1 versus High CO₂ 700–900 µatm, pH 7.9–7.8) [61,77,84,87] (table 1). Later experiments considered a wider range of CO₂/pH treatments (175–3000 µatm, pH 8.3–7.3) using a gradient of treatments levels across up to nine mesocosm enclosures, and allowing linear relationships between CO₂/pH and response parameters to be determined [79,82,83,86,88] (table 1). The observed OA effects on trace gases may reflect a combination of stress responses including acclimation, population re-structuring and associated adaptation [89]. Shipboard microcosm experiments are a useful tool to bridge the gap between complex mesocosm experiments and simple culture experiments (5–10 l, 3–12 replicates and 4–10 days). Conducting multiple short-term experiments over extensive spatial scales enables both the physiological effects of OA to be assessed, as well as the spatial variability in responses of surface ocean communities to future OA scenarios. The experimental design, involving relatively small incubation volumes of 5–10 l, allows multiple CO₂ treatments to be considered. For 18 experiments performed over a range of temperate and polar waters, Hopkins & Archer [80] and Hopkins et al. [90] used four CO₂ treatments in triplicate (Mid: 533.4 ± 40.0 µatm, pH 7.9 ± 0.03, High: 673.8 ± 82.2 µatm, pH 7.8 ± 0.1, High+: 841.5 ± 128.2 µatm, pH 7.8 ± 0.1, High++: 1484.0 ± 104.0 µatm, pH 7.5) and an ambient control (320.2 ± 38.3, pH 8.1 ± 0.1). Hussherr et al. [81] adopted a different approach by exposing phytoplankton communities within six single incubations to a pH gradient, from 509 µatm (pH 7.94) to 3296 µatm (pH 7.16) (table 1).

In the following section, we use information from all three types of experiments to consider the impacts of OA on trace gas production from the cellular to the community level, and in table 2,
we provide an overview of the types of response to OA that may result in changes in trace gas production.

To place the data in the context of key ocean–atmosphere linkages, we group the trace gases according to their primary atmospheric roles: those that primarily influence aerosols and cloud albedo, those that play a key role in regulating oxidative capacity and those that exhibit direct radiative effects, while recognizing that some trace gases possess multiple atmospheric roles. A large proportion of the following discussion focuses on DMS, since the amount of information available for this trace gas currently dwarfs the available information for all others. Furthermore, research into the net production of DMS in the surface oceans has been prominent within the fields of marine biogeochemistry and sea–air interactions for more than three decades due to the global significance of its role in climatic and atmospheric processes. The significance of oceanic DMS production and emission and its potential role in influencing global climate and atmospheric chemistry was highlighted in the 1980s by the seminal publication of Charlson et al. [4]; this spurred more than three decades of intensive investigation into the marine biogeochemistry, air–sea interactions and climate impacts of oceanic DMS.

6. Atmospheric role: aerosols and albedo

(a) Dimethyl sulfide and dimethylsulfoniopropionate

DMS is produced via enzymatic breakdown of the algal and bacterial secondary metabolite DMSP [20,93]. The release of intracellular DMSP into the surrounding seawater, and its subsequent and rapid conversion to DMS, is triggered by a number of processes including the active exudation of DMSP from living cells, and cell lysis during senescence, viral attack or grazing [20]. Most of the resulting DMS undergoes rapid processing in seawater via both bacterial (50–88%) and photochemical (8–34%) pathways [94]. The remaining DMS, which amounts to around 4–16% of total production, ventilates from the surface ocean to the lower atmosphere [95].

Upon entering the atmosphere, DMS undergoes rapid oxidation to species including SO$_2$, H$_2$SO$_4$, methanesulfonic acid (MSA) and dimethylsulfoxide (DMSO), thereby contributing to aerosol formation and growth and to atmospheric acidity [4,96]. When simulated at a global level, annually averaged DMS-derived aerosol radiative forcing at the top of the atmosphere has been estimated to have a climate-cooling effect of between $-1.69$ W m$^{-2}$ [97] and $-1.79$ W m$^{-2}$ [98]. For context, greenhouse gas radiative forcing for 1750–2011 was estimated at $+1.83$ W m$^{-2}$ for CO$_2$, $+0.61$ W m$^{-2}$ for CH$_4$ and $+0.17$ W m$^{-2}$ for N$_2$O [99]. DMS is also a major source of cloud condensation nuclei (CCN) via its rapid gas-phase oxidation to sulfuric acid (H$_2$SO$_4$), which influences the radiative properties of clouds, both microscopically via cloud droplet number concentration and effective radius, and at the larger scale by influencing cloud abundance, albedo and lifetime [100–102]. DMS also plays a significant role in the atmospheric oxidation pathways of other key trace gases, including isoprene, ammonia and halocarbons [103–105].

The response of DMS to increasing ocean acidity has been studied at several levels of biological and environmental complexity. At a fundamental level, DMS and DMSP concentration changes have been monitored in single-species cultures of algae exposed to relatively short-term (7–25 days) variations in OA. These studies report the response in two seaweed species, Ulva lactuca and U. clathrata, three strains of the well-studied DMSP-producer Emiliania huxleyi and two species of diatom [76–78,88,106]. The results of these studies, each following different experimental approaches, indicate a variety of physiological responses to short-term OA exposure between different species and strains of algae (electronic supplementary material, table S1). Such experiments exclude key bacterially mediated processes and do not consider how the activity of micro- and mesozooplankton (grazers) may be affected by OA. This illustrates the challenge of predicting the response of natural communities on the basis of the response of single species, and emphasizes the need for more in-depth understanding of the physiological roles of DMSP and DMS.
Table 1. Overview of experimental methods employed in studies of the effects of OA on marine trace gases.

| experimental technique | vol. (l) | number of replicates | duration (days) | key studies | CO₂ and pH treatments | method of acidification | What can it tell us? | strengths | weaknesses |
|------------------------|---------|----------------------|-----------------|-------------|---------------------|------------------------|----------------------|-----------|------------|
| single-species algal cultures | <1 | 2–3 | 7–40 | [76–79] | CO₂ (µatm) | pH | aeration with CO₂-enriched air, pH-stat to maintain constant DIC and pH | batch cultures: acclimated/physiological response to OA | useful tool for establishing baseline concepts | reduced complexity compared with natural populations | determinates direct response on trace gas production by phytoplankton isolates (if axenic) | high duplication/reproducibility | do not simulate complex natural systems | elimination of extracellular (bacterial) processes that may be key control on trace gas production |

| shipboard microcosm experiments | 5–10 | up to 12 | 4–10 | [80] | [81] | Av. of 18 expts: 320.2 ± 38.3 533.4 ± 40.0 673.8 ± 82.2 841.5 ± 128.2 1484.0 ± 104.0 5 treatments and control over range: 509–3296 | addition of strong acid/base, e.g. HCl/NaHCO₃ | physiological response and extent of the variability in response/plasticity between communities level of sensitivity to OA/high CO₂ | extensive spatial coverage natural gradients in carbonate chemistry, temperature, nutrients multiple short-term identical experiments on complex natural communities results in large, highly replicated, statistically robust data sets | short-term physiological response: representative? | bottle effects rapid acidification |

(Continued.)
Table 1. Continued.

| experimental technique | vol. (l) | number of replicates | duration (days) | key studies | CO$_2$ (µatm) | pH | method of acidification | What can it tell us? | strengths | weaknesses |
|------------------------|---------|----------------------|-----------------|-------------|--------------|----|------------------------|----------------------|-----------|------------|
| mesocosm experiments   | 2400–75 000 | 1–3                 | 25–35           |             |              |    |                        |                      |           | limited by number of experimental replicates difficult to test logistically challenging (physically and financially) minimal geographical coverage |
|                        | [82]    | 175–1085             | 8.3–7.6         | aeration with | CO$_2$-enriched | whole community response during bloom conditions (light and temperature) + large volume longer timescale = improved realism of representation of surface ocean towards a whole community, adaptive response |
|                        | [83]    | 400–1252             | 8.1–7.6         |             | air, or addition of |            |                        |                      |           |            |
|                        | [77]    | ambient              | 8.2 versus      |             |             |    |                        |                      |           |            |
|                        | [84]    | versus 700           | 7.8             |             |             |    |                        |                      |           |            |
|                        | [85]    | 300 versus           | 8.1 versus      |             |             |    |                        |                      |           |            |
|                        | [61]    | 780                  | 7.8             |             |              |    |                        |                      |           |            |
|                        | [86]    | 175–1085             | 8.3–7.6         |             |              |    |                        |                      |           |            |
|                        | [87]    | 400 versus           | no data         |             |              |    |                        |                      |           |            |
|                        | [79]    | 900                  | no data         |             |              |    |                        |                      |           |            |
|                        | [88]    | 160–830              | 8.1 versus      |             |              |    |                        |                      |           |            |
|                        |         | 350 versus           | 7.9             |             |              |    |                        |                      |           |            |
|                        |         | 700                  | 8.1–7.3         |             |              |    |                        |                      |           |            |
|                        |         | 280–3000             | 7.9–7.5         |             |              |    |                        |                      |           |            |
|                        |         | 330–1166             |                 |             |              |    |                        |                      |           |            |
Table 2. Overview of types of response to OA relevant to trace gas production and cycling.

| type of response to ocean acidification | description/example                                                                 | Relevance to which trace gases? |
|----------------------------------------|-------------------------------------------------------------------------------------|---------------------------------|
| direct chemical                        | effect of OA on chemical processes/equilibria that regulate trace gas production    | NH$_3$ methyl amines            |
|                                        | e.g. pH-induced shift from NH$_3$ to NH$_4^+$ leads to reduced NH$_3$ emissions      |                                 |
|                                        | [91]                                                                                |                                 |
| direct biogeochemical                  | effect of OA on biogeochemical processes that regulate trace gas production         | N$_2$O NH$_3$                    |
|                                        | e.g. pH-induced reduction in NH$_3$ leads to reduced nitrification and reduced N$_2$O production [92] |                                 |
| direct biological                      | effect of OA on organism-level processes that regulate trace gas production         | DMS(P) halocarbons CO isoprene  |
|                                        | e.g. pH-induced reduction in calcification in coccolithophores, leads to reduced abundance and reduced DMS emissions [79] |                                 |
| indirect biological                    | effect of OA on availability/type of organic substrates that regulate trace gas production | halocarbons OVOCs CO OCS, CS$_2$ isoprene |
|                                        | e.g. pH-induced increase in DOC exudation by phytoplankton enhances substrate/precursor availability [60] which may affect trace gas production |                                 |
| community level                        | effect of OA on community-level processes/community structure that regulate trace gas production and cycling | DMS(P) halocarbons N$_2$O OVOCs CO OCS, CS$_2$ isoprene |
|                                        | e.g. high CO$_2$(aq)-induced community-level shift towards dinoflagellates with low CO$_2$(aq) affinity and increased DMS(P) producing ability [82] |                                 |

At a more complex community level, the DMS response to OA has also been assessed using shipboard microcosms [80,81,90] (table 1). Such experiments test the resilience of natural communities to abrupt manipulation of carbonate chemistry, generally in controlled conditions using simulated light and temperature regimes. Of the 18 microcosm experiments reported in the meta-analysis by Hopkins et al. [90], all 11 experiments performed in the temperate waters of the northwest European shelf showed consistent large and significant increases in DMS in response to OA. By contrast, the seven experiments carried out in Arctic and Southern Ocean waters exhibited minimal OA influence on net DMS production. The discrepancy in DMS response between shipboard microcosms in temperate versus polar waters was hypothesized to be a product of variable levels of sensitivity of the respective communities to changes in the mean state of
carbonate chemistry [107]. In comparison to the well-buffered temperate waters of the northwest European shelf, polar waters are poorly buffered with respect to the addition of CO₂, resulting in a naturally large variability in carbonate chemistry [108]. Furthermore, the polar oceans experience unique biogeochemical processes, such as sea-ice formation and melt, iron-stimulated ice-edge blooms and under-ice organic matter respiration that also contribute to large natural variability in carbonate chemistry [109–111]. Thus, relative to the temperate communities, polar communities may be adapted to, and may be able to tolerate, large variations in carbonate chemistry, as reflected in the low sensitivity of DMS production to OA in polar waters. Of course, this hypothesis may not be universally applicable. A further 9 day microcosm study in polar waters performed during the summer [81] illustrated a substantial decrease in DMS concentrations with increased CO₂ and a less substantial but significant decrease in particulate DMSP concentrations. Such contrasting results may be unsurprising, given that the complexity of the DMS response to OA and the influence of a multitude of factors. For example, beyond the latitudinal variability in carbonate chemistry discussed in Richier et al. [107], the Arctic Ocean itself possesses high regional carbonate chemistry variability [112], related to sea-ice formation and the input of riverine and meltwater. Furthermore, spatial or seasonal differences in phytoplankton community composition, as well as the associated variability in physiological response (e.g. DMSP synthesis), could result in contrasting DMS responses to OA [77,80]. Alternatively, the overall DMS response may be associated with distinctive impacts on the transformation of DMSP via zooplankton grazing [61,86] or bacterial activity [79,81,82]. Given the wide variability in plankton community composition, activity and turnover rates, this emphasizes the need to consider both spatial and seasonal contexts when evaluating the sensitivity of DMS production to OA.

The majority of studies on the influence of OA on DMS have considered the response in large mesocosm enclosures (approx. 24–55 m³), incubated under close to natural environmental conditions. Such experiments incorporate the response of complex natural planktonic communities and the multiple processes that control the concentration of DMS in surface waters. Figure 2 provides an overview of nine mesocosm experiments carried out by different research groups in five Northern Hemisphere locations, ranging from arctic to subtropical latitudes and covering early summer to winter seasons. When DMS concentrations are integrated over the duration of each experiment, the difference in concentration between pCO₂ treatments of approximately 350 µatm versus approximately 750 µatm varied from +26% to −42%, with seven of the nine experiments showing decreased DMS concentrations with increased acidity (figure 2; electronic supplementary material, table S2). These pCO₂ levels approximate, respectively, the average ambient conditions at the time of experiments and the twofold increase that could occur by 2100 according to the Intergovernmental Panel on Climate Change (IPCC) Representative Concentration Pathways RCP6.0 emissions scenario [113].

Predominantly, a decrease in concentrations of DMS in response to OA in mesocosm experiments has been observed, which suggests that a dominant control on DMS net production is affected by a change in carbonate chemistry or H⁺ concentration. DMSP production by phytoplankton is highly species-specific and several studies have demonstrated correlations between phytoplankton community composition and DMSP concentrations that may have influenced the response of DMS production to OA [77,79,82,83,86] (table 2). On several occasions, specific aspects of DMS production have been examined: direct measurements of the rates of DMSP synthesis [82], grazing rates on phytoplankton that may enhance conversion of DMSP to DMS [61,86], the in vitro activity of DMSP-lyase enzymes that convert DMSP to DMS [86] and rates of bacterial metabolism of DMSP and conversion to DMS [83]. Nonetheless, disentangling the complex processes responsible for the observed changes in DMS remains challenging.

The variety of ways in which data from the nine large-scale mesocosm experiments have been interpreted also creates a further challenge when attempting to unify the DMS response to OA. Time-integration of DMS concentrations across the full duration of each experiment (figure 2) is a start towards consistent interpretation but a more in-depth approach is required to fully compare experiments and extract a comprehensive evaluation. As an example, a large significant
increase in DMS in relation to increased CO₂ was reported by Kim et al. [61] for their experiment in Korean waters—strongly in contrast to all other studies, and which was attributed to an OA-induced enhancement of dinoflagellate grazers. Such differences highlight the complex responses that can play out within the bounds of a mesocosm experiment, making a general, broad-brush interpretation of results challenging.

The effects of both increased temperature and CO₂ on natural communities have been investigated in three experiments using similar sized mesocosms (2.4–2.6 m³), either in situ [61,86] or located in controlled-environment containers [114]. In two cases, a reduction in time-integrated DMS with increased CO₂ was observed but the elevated temperature had contrasting effects: in the warmer Korean coastal waters (16–19°C), a 2°C elevation in temperature appeared to
decrease net DMS production slightly. Conversely, in the cooler Gulf of St Lawrence, enhanced seawater temperature (10 versus 15°C) led to an increased rate of growth of phytoplankton and bacteria, resulting in elevated DMSP and DMS concentrations. By contrast, Kim et al. [61] saw large grazing-induced increases in DMS under both high CO₂ and high CO₂ + 3°C, with the greatest increase in the high CO₂ treatment. These contrasting results could reflect differences in the level of adaptation of the respective communities to natural temperature and CO₂ variability, but also highlights the challenges involved in disentangling the complex processes that result in net DMS production.

Finally, it should be recognized that clear discrepancies have arisen in the DMS response to OA between different experimental techniques, which may make interpretation of the overall response challenging. For example, the results of shipboard microcosms contrast strongly with those from mesocosms. However, interpretation of the data can be facilitated by an understanding of the strengths and weaknesses of each technique, and the specific hypotheses each technique is designed to address. Each approach provides valuable information on how OA may influence DMS production in the future. Microcosm experiments are necessarily short term (less than 10 days), so the response to OA is considered to reflect the physiological plasticity of the community, i.e. how well they are adapted to rapidly changing carbonate chemistry, but may not fully capture the effects of shifts in community composition. By contrast, mesocosm experiments are generally much longer (typically approx. 30 days), allowing multigenerational OA-induced changes in taxonomy and community structure to affect DMS concentrations. The microcosm approach may aid in the identification of OA-sensitive regions in terms of DMS production. On the other hand, mesocosm experiments provide some information on how community composition shifts in response to OA may affect the processes controlling the cycle of DMSP and DMS and hence determine their concentrations.

(b) Nitrogen species

(i) Ammonia, methylated amines, alkyl nitrates

Oceanic emissions of the soluble trace gas ammonia (NH₃) play a role in marine aerosol formation, and the related ammonium ion (NH₄⁺) provides an inorganic nutrient fundamental to phytoplankton productivity in the surface ocean [104,115]. Both NH₃ and its organic analogues, the methylated amines (RₙNH(3−n)), are directly affected by changing pH due to their capacity to accept protons (i.e. as bases). A decrease in seawater pH will result in a shift in NH₃:NH₄⁺ equilibrium towards NH₄⁺, and the projected decline in ocean pH from 8.1 to 7.8 by the year 2100 is estimated to reduce the NH₃ concentration by 50% [91], decreasing the availability of gas-phase NH₃ for ocean–atmosphere gas transfer (table 2). As the recycling of NH₃ between the sea and atmosphere is considered to be a major component of the cycling of nitrogen in the marine atmosphere [116], OA has the potential to have a major impact on marine aerosol chemistry over the open ocean, including feedbacks on atmospheric acidity and iron solubilization (e.g. [117,118]) and on particle formation (e.g. [119]).

Recent studies have suggested that marine nitrification of ammonium to nitrate may be significantly inhibited under OA (e.g. [120]) in line with a shift in NH₃:NH₄⁺ speciation towards NH₄⁺. Nitrogen fixation, an important source of new nitrogen to ocean ecosystems will potentially be enhanced under high CO₂ conditions ([121] and references therein). A recent meta-analysis of OA studies suggests a decrease of 29% in nitrification, and corresponding increase in nitrogen fixation of 29%, by 2100 under the ‘business as usual’ emissions scenario [45]. In addition, there is some evidence that NH₄⁺ uptake by diatoms may be suppressed by OA [122].

Given the complex controls on NH₄⁺ concentration in the marine environment, it is currently uncertain whether OA will lead to higher NH₄⁺ concentrations and thus lower ammonia emissions. However, it should be considered whether the simple chemistry that results in a pH-induced shift in NH₃:NH₄⁺ equilibrium could on its own alter seawater NH₃ concentrations.
enough to influence the sea–air exchange of NH$_3$ and amines. Further studies considering the
direct effects of OA on the production or consumption of NH$_3$, amines or the atmospherically
important alkyl nitrates are required.

(ii) Terpenes

The marine terpenes (isoprene C$_5$H$_8$ and monoterpenes –C$_{10}$) occur ubiquitously in the marine
environment and have the potential to significantly influence climate via the production of
secondary organic aerosol (SOA) [123–125]. There is some recent evidence that OA may induce
changes in terpene production by macroalgae, although the direction of response is uncertain and
may vary between species, and so requires further investigation [126].

7. Atmospheric role: oxidative capacity

(a) Halocarbons

The surface ocean is a key source of short-lived brominated and iodinated organic compounds
(halocarbons) to the atmosphere. Marine emissions of halocarbons, dominated by bromoform
(CHBr$_3$), dibromomethane (CH$_2$Br$_2$) and methyl iodide (CH$_3$I) [127], originate from a range
of biological and photochemical processes. These include direct biosynthesis by bacteria (e.g.
[128]), phytoplankton (e.g. [70]) and macroalgae (e.g. [13]), and indirect production via reactions
between DOM and light [129,130] and/or ozone [23]. Upon entering the atmosphere, halocarbons
are rapidly oxidized, yielding short atmospheric lifetimes of less than half a year [131,132], and
releasing highly reactive halogen radicals (e.g. I, IO, Br, BrO). These radicals exert an important
control on tropospheric ozone [10,133–135], and contribute to the production of new particles and
CCN with the potential to influence climate [136].

As marine production of halocarbons is governed by biological processes and the availability
of biological substrates (table 2), OA is expected to impact upon their production, with potential
feedbacks on atmospheric and climatic processes [137]. However, mesocosm studies have found
no obvious effect of high $p$CO$_2$ and OA on the emission of CHBr$_3$ or CH$_2$Br$_2$ (e.g. Norwegian fjord
[84]), Arctic (Spitsbergen) [85], and brackish waters (Baltic Sea, [88]). By contrast, concentrations
of CH$_3$I were significantly reduced (by up to 67%) under high $p$CO$_2$ conditions during and after a
phytoplankton bloom in temperate waters [84], while in the Baltic Sea, no response was observed
[88] (see the electronic supplementary material, table S3 for a summary). Given that these limited
studies report conflicting relationships between OA and halocarbon production by surface ocean
communities, this is an area that requires further investigation.

A single study [138] has also considered the effects of OA on halocarbon production by
tropical seaweeds. Seaweeds are important localized sources of halocarbons [13,14]. In the tropics,
biogenic halocarbons contribute disproportionately to stratospheric halogen concentrations and
ozone cycling via deep tropical atmospheric convection [139]. Furthermore, seaweed farming is
a growing industry in the tropics, so the importance of halocarbon emissions to the atmosphere
may increase in the future [138]. Mithoo-Singh et al. [138] assessed the response of halocarbon
production by five tropical seaweed species to four OA treatments (pH 7.8, 7.6, 7.4, 7.2) relative
to ambient (pH 8.0). In general, lower pH resulted in higher halocarbon emission rates, with the
effect greatest at the lowest pH treatments (7.4, 7.2). Some resilience within the tested seaweeds
to the less severe pH treatments (7.8, 7.6) was apparent, which may result from a degree of
adaptation to variation in pH which occurs naturally in coastal waters [140]. However, this should
not be taken to represent a linear response given that pH is the $\log_{10}$ of the H$^+$ concentration.
Hence, a greater effect could be expected from the difference between the two lower pH values
(7.4, 7.2) than between pH 7.8 and 8.0.
(b) Oxygenated volatile organic compounds

The small and simple oxygenated VOCs (OVOCs) include methanol, ethanol, propanol, acetaldehyde and acetone. Although predominantly emitted from terrestrial ecosystems [141,142], the oceans play a role as both a source and sink of OVOCs [143–149]. OVOCs affect the oxidative capacity of the troposphere by influencing the ozone budget, consuming hydroxyl (OH) radicals and creating hydrogen oxide (HO\textsubscript{x}) radicals [150,151]. Although understanding is limited, marine production of OVOCs is linked to biological processes [146,147,152]. For example, acetone and acetaldehyde are thought to be principally produced by photochemical reactions involving the humic component of chromophoric dissolved organic matter (CDOM) [153–156] with up to 68% of gross acetaldehyde production and up to 100% of gross acetone production via this route [147]. Therefore, any OA-induced effect on CDOM characteristics or availability may impact upon the production of these compounds (table 2). Some methanol production may occur via release from living or senescing algal cells [152,157–160], so any OA effects on algal processes could affect the subsequent production of methanol. More research into these gases is required if we are to increase our understanding of the effects of OA on their net production and fluxes.

8. Atmospheric role: direct radiative effects

(a) Nitrous oxide (N\textsubscript{2}O)

The ocean accounts for approximately one third of natural global emissions of the trace gas N\textsubscript{2}O [161,162]. N\textsubscript{2}O has the third largest radiative forcing of the anthropogenic greenhouse gases (approx. 300 x CO\textsubscript{2} on a molecule per molecule basis) on a global basis [2], and is also a dominant ozone-depleting substance in the stratosphere [163]. It is produced primarily via nitrification in the open ocean, as a by-product of the oxidation of ammonium (NH\textsubscript{4}+) to nitrite (NO\textsubscript{2}−). N\textsubscript{2}O is also produced as a by-product of the reductive denitrification pathway in hypoxic and suboxic environments such as oxygen minimum zones and sediments, where O\textsubscript{2} concentrations are sufficient to inhibit N\textsubscript{2}O consumption by nitrous oxide reductase enzymes [18,164].

Although there are few studies on the influence of OA on N\textsubscript{2}O, there is greater insight into its impact on the primary source process of nitrification [165]. Huesemann et al. [166] identified a reduction in nitrification rate by up to 90% at pH 6.5, relative to ambient pH, with a linear rate decline across this pH range. Similarly, ammonium oxidation (the first stage of nitrification) decreased to near-complete inhibition at pH 6.5 in experiments using surface waters from the English Channel [167,168]. Although these results are compelling, it should be noted that both studies used lower pH levels than that projected for the next century. Nevertheless, using a more conservative and relevant pH range from 8.09 to 7.42, Beman et al. [120] showed unequivocal evidence of an inhibitory effect of OA on nitrification at locations in the Pacific and Atlantic.

Conversely, Fulweiler et al. [170] found that nitrification rates increased with decreasing natural gradients of pH in Narraganset Bay, which they attributed in part to changes in the microbial community in response to competition for NH\textsubscript{4}+/NH\textsubscript{3}. This is consistent with other suggestions that nitrification may be influenced by OA directly via altered microbial physiology or community composition, or indirectly by changes in the supply of organic material [171]. Hutchins et al. [172] speculated that increasing levels of CO\textsubscript{2} may lead to an increase in
autotrophic nitrification rates via a CO₂ fertilization effect, although this has not been observed in the open ocean. Changes in microbial community composition and abundance in response to OA have been reported [173], particularly for ammonium oxidizing bacteria (AOB) relative to ammonium oxidizing archaea (AOA). Whereas AOB and AOA are ubiquitous and both produce N₂O [174,175], AOA are considered to be the principal nitrifying organisms [176,177], and so an OA-induced shift to AOB may alter marine N₂O production. However, metabolic flexibility may provide some degree of adaptability, with continued growth by coastal AOA reported at a pH of less than 6 [177].

At a lower pH the N₂O:N₂ yield of denitrification increases in other environments [178], yet the limited studies in marine systems to date suggest no overall significant effect of OA on denitrification [45]. However, as nitrification and denitrification are coupled in coastal sediments, an OA-induced reduction in nitrification rate may reduce nitrate availability for denitrification leading to a net decrease in N₂O production by both processes [45], although this has yet to be confirmed.

The limited evidence to date suggests that nitrification and associated N₂O production may decrease in the future in response to OA with potential implications for the global marine N₂O source. In a meta-analysis, Wannicke et al. [45] concluded that OA might reduce nitrification by 29 ± 10%, consistent with the observed reduction of 3–44% reported by Beman et al. [120]. This equates to a decrease in global N₂O production for the next 2–3 decades of 0.06–0.83 Tg N yr⁻¹, which is comparable with current global N₂O production from fossil fuel combustion and industrial processes (0.7 Tg N yr⁻¹). On the assumption that 50% of the global ocean N₂O source is produced by nitrification [162], Rees et al. [92] projected comparable, albeit slightly lower, reductions in oceanic N₂O production. Consequently, the evidence to date suggests the influence of OA may have a small negative feedback on climate change via a reduction in radiative forcing attributed to marine N₂O emissions.

(b) Methane

Methane (CH₄) is a long-lived atmospheric trace gas, which acts as a potent greenhouse gas in the troposphere with a radiative forcing effect, on a molecule per molecule basis, of approximately 25 × CO₂ [179,180]. The ocean plays a minor role in the present-day global CH₄ budget of the atmosphere [181], contributing a maximum of 10% of the global CH₄ burden [182]. Marine CH₄ sources are, however, not well constrained, owing to a paucity of observations [183]. Coastal environments including estuaries could account for approximately 75% of the marine source [184], and coastal upwelling areas are also strong sources [185]. Despite the uncertainty regarding the source of CH₄ in the surface open ocean [186], there is a potential for direct impacts of OA on CH₄ production, via two recently identified methane production pathways, involving DMSP [187] and methane phosphonate [188,189]. CH₄ production and consumption mechanisms could also be indirectly impacted by OA, for example, via OA-induced changes in transparent exopolymer particles (TEP) and particle formation [60] that influence methanogenesis in anoxic microsites [190]. CH₄ production did not show any OA effects in two studies, but both are currently unpublished: one used Arctic microcosms (A Rees 2019, personal communication) and another used coastal mesocosms (F Deans and C Law 2019, personal communication).

(c) Carbon monoxide and carbonyl sulfide

The surface ocean is a net source of carbon monoxide (CO), produced via both microbial and abiotic processes, and removed by microbes, mixing and gas exchange [167,168,191,192]. In the atmosphere, CO is a greenhouse gas with a radiative forcing effect of ~2 × CO₂ (on a molecule for molecule basis). Furthermore, CO indirectly affects the climate by out-competing CH₄ in reactions with tropospheric OH radicals, resulting in enhanced concentrations of this far more potent greenhouse gas (CH₄ ~25 × CO₂, see Methane) [193]. Although there are still large uncertainties over the size of the oceanic source of CO, it is likely to be controlled by the quality and quantity
of available CDOM [167,168]. Thus, OA effects on bacterial or phytoplankton processes (that determine the CDOM pool) may alter CO production.

Carbonyl sulfide (OCS) is the most abundant sulfur-containing trace gas in the atmosphere, with marine emissions contributing significantly to the total global budget [194]. OCS is produced in the surface ocean via reactions between UV radiation and CDOM [194,195]. It enters the atmosphere directly via oceanic emission, and indirectly via the oxidation of DMS and carbon disulfide (CS$_2$) in the atmosphere [196–198]. OCS is both a climate-warming greenhouse gas and a climate-cooling aerosol precursor, with the two opposing radiative effects currently in near-balance [199]. However, future changes in the magnitude of sources and sinks may upset that balance [200]. For example, OA may change the oceanic source of OCS via either altering direct emissions from the ocean and wetlands or indirectly via changes in emissions of its precursor gases DMS and CS$_2$.

9. Cold and naturally carbonated: trace gas emissions from ocean acidification-sensitive regions

(a) Polar oceans

Although OA is a global phenomenon, it is progressing with the greatest speed in regions of the ocean that have naturally high dissolved inorganic carbon (DIC) levels and low alkalinity such as high latitude waters of the Southern Ocean and Arctic [28]. In the Arctic Ocean, OA is also accompanied by sea-ice melt water, glacial runoff and river discharge, as well as enhanced terrestrial organic carbon loading, thawing permafrost, gas hydrate destabilization and anthropogenic pollution, which might all further accelerate OA [201–204]. The surface waters of the Arctic Ocean could see a 185% increase in hydrogen ion concentration (ΔpH = −0.45) and basin-wide undersaturation in aragonite ($\Omega_{\text{arag}}<1$) by the end of this century [201,205,206], although with high regional heterogeneity. Based on simulations using the RCP8.5 scenario with the highest concentrations of atmospheric CO$_2$, Popova et al. [112] suggest that the central Arctic, Canadian Arctic Archipelago and Baffin Bay present the greatest rates of acidification and carbonate saturation decline as a result of melting sea ice. By contrast, areas affected by Atlantic inflow including the Greenland Sea and outer shelves of the Barents, Kara and Laptev seas, see minimal decreases in pH and carbonate saturation because diminishing ice cover leads to greater vertical mixing and primary production. OA in the Southern Ocean is primarily driven by the oceanic uptake of anthropogenic CO$_2$ in combination with the naturally strong winter upwelling of DIC-rich, low-alkalinity subsurface waters [207]. Regions of the Southern Ocean already experience sporadic short-term aragonite undersaturation events, the spatial extent and duration of which are expected to accelerate within the next 15–20 years under high CO$_2$ emission scenarios (RCP 8.5) [208].

The polar regions are important for the production of aerosol precursors, such as DMS, that influence CCN production and radiative forcing. In the summertime Arctic atmosphere, marine DMS-derived aerosols significantly contribute to new particle formation events that may influence cloud processes and Arctic atmospheric albedo [7,8,209,210]. The Southern Ocean is a globally important DMS source—regions north of the sub-Antarctic front contribute approximately 15% to global DMS emissions [211], making a significant contribution to DMS-driven secondary aerosol formation [6], and contributing 6–10 W m$^{-2}$ to reflected short wavelength radiation—comparable with the forcing by anthropogenic aerosols in the Northern Hemisphere [212]. Thus, any climate change-induced modification to DMS emissions from polar regions could influence radiative forcing at both regional and global scales. The modelling studies by Six et al. [43] and Schwinger et al. [44] indeed show a significant radiative forcing and surface temperature increase due to OA-induced reductions in polar DMS production under high emissions pathways (e.g. a 0.86 W m$^{-2}$ reduction in reflected short wave radiation south of 40°S in the study of Schwinger et al.).
A small number of experimental studies report the effects of OA on DMS in polar waters. In a mesocosm experiment in Kongsfjorden, Svalbard Archipelago (78°N), a 35% decrease in DMS at 750 µatm was attributed to a decrease in bacterial DMSP-to-DMS yields [82]. Similarly, in a microcosm experiment conducted in Baffin Bay, Canadian Archipelago (71°N), a 25% decrease in DMS at approximately 1500 µatm was found, attributed to an OA-related increase in sulfur demand by the bacterial assemblage [81]. These limited results suggest that the net production of DMS during the productive summer season in the Arctic could decrease via bacterioplankton-mediated processes with ongoing OA. In contrast to these previous experiments, in a series of shipboard microcosm experiments in the Arctic and Southern Ocean surface waters, little biological effect and minimal DMS response to OA was observed [90], suggesting a high level of resilience to a changing carbonate chemistry environment within the sampled communities. This agrees with previous evidence that polar microbial communities may be adapted to a changing carbonate chemistry environment as they experience strong natural fluctuations in pH (over the range 7.5–8.3) over diurnal/seasonal and local/regional scales [112,213–215]. However, it cannot be excluded that variations in community responses may be linked to differences in experimental approaches used, as previously described (see section on ‘Reconciling differences within and between experimental techniques’).

(b) Eastern boundary upwelling systems

Eastern Boundary Upwelling Systems (EBUS) are considered particularly susceptible to OA, given the combined effects of their naturally high DIC concentrations and enhanced uptake of anthropogenic CO₂ [216]. Characteristic examples of EBUS include the California and Peru/Humboldt EBUS in the Pacific, and the Canary and Benguela EBUS in the Atlantic. Cold DIC-rich subsurface waters are upwelled to the surface layer by trade wind forcing at seasonal and interannual timescales, lowering the pH of surface waters relative to open ocean surface waters [216]. This enhances the rates of OA within such systems relative to the global surface ocean. Recent data from the California Upwelling System, using a proxy record of fossil foraminifera calcification response, have shown a 35% decrease in [CO₃²⁻] and a drop in pH of 0.21 units since pre-industrial times, which exceeds the global mean decline by a factor of two [217]. Due to the high decomposition rate of organic matter as well as the input of equatorial low-O₂ water masses, oxygen minimum zones (OMZs) exist in these coastal areas affecting the regional oxygen, nitrogen, carbon and sulfur cycles through nitrification, denitrification, anammox and sulfate reduction processes and influencing local trace gas production. EBUS are considered to be ‘hot spots’ for emissions of greenhouse gases (N₂O, CH₄) and reactive species such as DMS and H₂S [18,218–224].

The addition of anthropogenic CO₂ into the already corrosive waters of EBUS could rapidly push these systems closer to critical thresholds, such as aragonite undersaturation [216]. Indeed, pH values as low as 7.6 and 7.7 have been measured in the Californian upwelling, accompanied by a shoaling of the aragonite saturation horizon by about 50 m since preindustrial times [216]. This leads to periodic upwelling of corrosive waters during the summer months [216,225,226], with the potential to impact upon ecologically and economically important species [216,225,227,228]. Despite the observational work that has been undertaken, little experimental work has been conducted on the implications of OA in EBUSs (e.g. [229]). Thus, the OA impact on trace gas production in these regions is still highly uncertain, but potentially in line with the responses observed in other regions, as described above.

10. Ocean acidification, warming and deoxygenation: the multi-stressor effects on marine trace gases

OA is not occurring in isolation to other global environmental changes. In addition to having taken up approximately 28% of the excess anthropogenic CO₂ since 1750, the ocean has also
absorbed approximately 93% of the excess heat over the past 45 years [30]. Both processes profoundly modify the physical and chemical environment experienced by marine organisms. Warming enhances biological rates [230,231] and decreases the solubility of gases, resulting in decreasing global ocean oxygen inventories [29]. Warming and freshening enhances surface ocean stratification [232], which in turn decreases mixed layer depth and reduces the entrainment of nutrients into the euphotic layer, while resulting in higher levels of irradiance experienced by organisms [29]. This alleviates light limitation at high latitudes, but enhances nutrient limitation at low- to mid-latitudes. Reductions in nutrient entrainment may be compensated for by the atmospheric deposition of anthropogenic aerosols, which itself could be countered by future improvements in air quality standards [233]. Thus, CO₂-driven changes in seawater carbonate chemistry occur simultaneously with warming, deoxygenation, localized freshening of the ocean and changes to nutrient dynamics [234,235]. Numerous modelling studies have addressed future changes in marine ecosystems and biogeochemistry in response to these drivers, either in isolation or combined [29,49,236], yet very few have focused on trace gas emissions and OA-related feedbacks to the Earth system [42–44].

(a) Changing dimethyl sulfide emissions in response to ocean acidification: earth system feedbacks

Although experimental data provide useful information on the potential future DMS response to OA, these data become most powerful when included in an Earth System Model (electronic supplementary material) to facilitate upscaling and estimation of feedbacks of projected changes in DMS emissions on future climate (figure 3). So far, two studies have used electronic supplementary material to provide evidence for a potential positive climate feedback arising from pH-sensitivity of DMS production [43,44]. At the end of the current century, the electronic supplementary material showed major pH-induced reductions in DMS production for areas of high biological production, such as the upwelling equatorial Pacific and other EBUS, the eddy-driven upwelling in the Southern Ocean around 40°S and the subpolar biome in the North Atlantic [43,44]. Both studies revealed a subsequent significant radiative forcing and surface warming in response to the decreased DMS flux to the atmosphere and subsequent changes in aerosol and cloud properties. Schwinger et al. [44] used a fully interactive model, able to simulate a range of feedbacks: they found a global linear relationship between pH-induced changes of DMS sea–air fluxes and a transient surface temperature change of $-0.041^\circ\text{C} \, \text{TgS}^{-1} \, \text{yr}^{-1}$, driven by reductions in global DMS emissions. These model experiments were conducted with a high-emission scenario (RCP8.5) as a baseline, leading to average surface pH reductions of 0.44 and 0.73 units in 2100 and 2200, respectively. The corresponding reduction of DMS fluxes (assuming the ‘medium’ pH-sensitivity of DMS production of [43]) is 4 Tg S yr$^{-1}$ (17%) in 2100 and 7.3 Tg S yr$^{-1}$ (31%) in 2200. The simulated additional surface warming has a north–south gradient with much stronger surface warming in the Southern Hemisphere due to the larger area covered by ocean (figure 3).

Both models described here are parametrized using the empirical relationship between pH and DMS observed in a number of mesocosm studies [77,82,84,87], while recognizing that the level of understanding of the DMS response to OA within these experiments is limited. It should also be noted that these data consider OA as a single stressor, with a complete lack of information for other key climate stressors such as ocean warming. Furthermore, our interpretation of the DMS response between mesocosm studies is confounded by inconsistencies in composition and physiological status of starting communities and experimental set-up (e.g. volume of seawater, method of acidification, inorganic nutrient additions, inclusion/exclusion of higher trophic levels, light and UV cycles, mixing, wall effects/cleaning) that make it difficult to draw direct comparisons (see discussion below). To increase the accuracy of model outcomes and facilitate a better understanding of the future feedbacks and climatic effects, improved comparison and integration of all DMS data from mesocosm experiments are required. For example, normalizing
Figure 3. Outputs from the fully interactive Earth system model run from Schwinger et al. [44]. The top row of panels show DMS sea–air flux (a), sulfate aerosol burden (d) and surface temperature (g) in the reference simulation (no sensitivity of DMS fluxes to OA). The corresponding changes in a simulation assuming a decrease of DMS production with increasing pH are shown in the middle row of panels (b,e,h) and zonal mean changes are depicted in the bottom row (c,f,i). The grey shaded area in the zonal mean plots gives the range of natural variability (defined as the standard deviation of the zonal mean found in the control run). (Online version in colour.)

for differences in experimental design, community structure and carbonate chemistry dynamics could lead towards a more accurate empirical relationship between pH and DMS. Where the DMS representation in a model is detailed enough, e.g. [237], it would be beneficial to include the effects of OA on the processes controlling DMS production in the surface ocean, for example, using data from short term, small-scale experiments, such as shipboard microcosms (e.g. [80,81,90]).

Finally, there is a large gap in our understanding of the response of net DMS production to other climate stressors in community-level experiments, in particular the response to increasing temperatures. Three mesocosm experiments to date have considered the combined effects of OA and temperature [61,86,114]. Our understanding of the multi-stressor response of DMS, and other trace gases, would be improved with a greater understanding of such processes. For example, Dani & Loreto [22] hypothesize that phytoplankton isoprene emitters favour warmer latitudes, as opposed to cold water-favouring DMS emitters. This may imply that as warmer waters extend towards higher latitudes with climate change, there could be an increase in geographical extent of oceanic isoprene producers to the detriment of DMS producers. However, further work is now required to fill such gaps in our understanding.

(b) Marine nitrogen cycle Earth system feedbacks

Changes to the future ocean source of N\textsubscript{2}O have been evaluated as a direct consequence of global warming-driven changes in ocean circulation and productivity [238] and combined with anthropogenic nitrogen deposition [239]. Both studies report a decrease in N\textsubscript{2}O emissions by 2100 of between 4–12% [238] and 24% [239]. The decrease results from both a net reduction in N\textsubscript{2}O production and an increase in N\textsubscript{2}O storage driven by enhanced stratification and reduced N\textsubscript{2}O sea–air flux. The reduction in net N\textsubscript{2}O production in both studies is largely driven by reduced primary production and export production resulting in decreased water column nitrification, changes in ocean circulation in response to global warming and atmospheric N\textsubscript{2}O
concentrations, combined with expansions of OMZs and associated increases in water column denitrification [238,239]. On millennial timescales under sustained anthropogenic climate forcing, Battaglia & Joos [240] project increases in N₂O production of 21% due to deoxygenation and elevated remineralization fluxes. Under steady-state conditions, these millennial increases in N₂O emissions are shown to cause a small positive climate feedback (0.004 W m⁻¹ K⁻¹ for RCP8.5, [240]).

However, given the limited evidence for direct effects of OA on N₂O production, the model studies assessing the future evolution of N₂O emissions have not, so far, included the effects of OA. Similarly, effects of deoxygenation on N₂O production in a warming ocean remain underexplored, largely due to persistent biases in the climatological representation of OMZs and in reproducing the expansion of low oxygen waters in electronic supplementary material [241,242]. Modelling studies that apply parametrizations based on mesocosm studies to describe the effect of OA on the stoichiometry of organic matter [243] have shown that OA can exacerbate ocean deoxygenation via enhanced C: N ratios in organic matter [244,245]. The higher C: N ratio in organic matter would constitute a negative feedback on atmospheric CO₂ through the strengthening of the biological pump. However, the enhanced O₂ utilization during remineralization would promote the production of N₂O, a positive feedback to the Earth radiative budget, which would offset the first one. The change in the stoichiometry of organic matter in response to OA remains, however, to be confirmed by further studies [246,247]. In summary, current model projections suggest a future decline in global marine N₂O emissions, and small negative feedback to climate change. However, these analyses do not account for the influences of OA and have limited capability to assess key influences on the marine nitrogen cycle such as deoxygenation. Elucidating these influences will require a combination of improved process knowledge and incorporation of this into more representative biogeochemical process models.

(c) Key areas of future research on multi-stressors

While models are vital to exploring responses of trace gas emissions to multiple stressors, the development of adequate parameterization depends on experimentally evidenced process understanding. Figure 4 summarizes our knowledge of the anticipated direct and indirect effects of multiple stressors on trace gas production. These stressors operate at global scales, including warming and acidification, and at regional scales, such as in coastal waters and polar regions. Figure 4 also indicates the inferred trace gas response (increased or decreased production) to each stressor, although many of these are based on limited observations. Whereas some stressors, such as eutrophication, are considered to have a primarily stimulatory effect on trace gas production, others can have both positive and negative impacts. For example, warming may stimulate trace gas production by enhancing metabolic rates and reducing oxygen availability, but may also reduce phytoplankton diversity potentially reducing production of taxa-specific trace gases such as DMS and halocarbons. From the perspective of individual trace gases, the production of some, such as methane, may increase in response to most stressors, whereas the majority of trace gases may show increased or decreased production depending on the stressor. Perturbation experiments on marine ecosystems that assess multiple stressors are still rare, and consequently, there is little information as to how they influence trace gas production. Although the overriding trend in marine multiple stressor studies is synergistic, relative to rates of the individual stressors [248], multiple stressors with opposing impacts may cancel each other out, or alternatively one may dominate. It is recommended that future studies of trace gas production consider the impact of multiple stressors [249] using region-specific projections for climate variables (see [250]).

11. Conclusion

The potential for marine trace gas emissions to influence and impact atmospheric chemistry and climate are substantial. The changes in net production of some trace gases such as DMS and N₂O, indicated in OA studies and models, point to potentially large and globally significant
Figure 4. Summary of our knowledge on multiple stressors and their anticipated direct and indirect effects on trace gas production. Coloured arrows represent known/anticipated trace gas response (red, increase; blue, decrease; green, no net change), and black arrows describe the direction of change of the related process. HABs, harmful algal blooms; TEP, transparent exopolymer particles; DON, dissolved organic nitrogen. (Online version in colour.)
modifications to sea–air fluxes. This could lead to either warming (e.g. lower DMS emissions) or cooling (e.g. lower N$_2$O emissions) effects on climate. Where data for other trace gases are scant, we cannot yet be confident in the direction of change, but we can have greater certainty that there is the potential for impacts on net production, and so chemistry and climate, with global-scale effects.

However, relative to other aspects of marine biological and ecological research, the field of marine trace gas production is under studied. Even our understanding of the basics could be improved, such as the processes driving production and cycling within the surface ocean. These knowledge gaps can limit our ability to design appropriate experiments or to interpret findings in the context of OA. Furthermore, even where some data are available, the limited mechanistic representation of biological and biogeochemical processes in electronic supplementary material limits the predictive capability of future trace gas production and emissions, and related climate effects. Inconsistencies in the effect of OA on trace gas production result from the complexity of trace gas cycling, with the involvement of multiple production and loss processes (e.g. phytoplankton species composition, bacterial processes and grazing activities). Further complications arise when the potential for both direct and indirect effects on trace gas production is considered, and as with other aspects of OA research, the indirect effects are more challenging to pin down (figure 4). Finally, interpretation of experimental data and projections in terms of atmospheric chemistry and climate are complicated by trace gas sensitivity to other climate change stressors (warming, deoxygenation and eutrophication), some of which may be more important determinants of production and emissions.

Of course, understanding the biological mechanisms (and their regulation) will be crucial for interpreting the trace gases response to OA, using both model organisms in the laboratory and natural communities within field experiments, and addressing the current shortcomings requires an improved experimental approach. Although short-term OA studies provide useful information on the physiological plasticity of surface ocean communities and associated trace gas production, and existing levels of adaptation to fluctuations in carbonate chemistry, such experiments cannot accommodate the potential for evolutionary adaptation of planktonic communities (e.g. [251–254]). Therefore, it would be beneficial to carry out longer term experimental studies, encompassing multiple generations, in order to detect adaptation of planktonic communities to OA and other climate change stressors. Such adaptation in phytoplankton becomes evident after only a few hundred generations, representative of a period of approximately 6–12 months (e.g. [251]). Parallel measurements of process rates and standing stocks of trace gases would provide greater insight into the role of OA in influencing trace gas production. However, the implementation of long-term experiments of this kind are likely to be limited to culture conditions using isolated strains, and thus at the expense of other important ecological and biogeochemical interactions (see [52]). Ecological level experiments will still involve a trade-off in terms of duration and number of generations, but will continue to provide important information on the role of species interactions and succession on trace gas production. Both experimental approaches could integrate multiple stressors, thus closing some of the gaps in our understanding of the trace gas response to climate change. An enriched experimental understanding could be complemented by improved surface ocean measurements. To this end, we recommend that future surface ocean trace gas measurements are accompanied by quantification of at least two components of the carbonate system, so that global databases can be used to relate spatial variability in trace gas concentrations to variations in surface ocean pH. This would greatly increase our understanding of the influence of the carbonate system, including the physical and biogeochemical processes that control pH, on trace gas concentrations in the surface oceans.

To provide reliable projections of future marine emissions of climate-relevant gases, studies will need to characterize and quantify the nature of the adaptation and/or resilience of diverse trace gas-producing communities. The complexities of these investigations are compounded by the multitude of environmental changes, including OA, that affect these communities. Such studies would be complex in design and implementation, and require community-wide
collaborative efforts, involving researchers from multiple disciplines and significant levels of financial investment (see [52,249,255]). However, given success, this would improve our understanding of the longer term effects of OA on the biological and biogeochemical processes involved in trace gas production, and build an improved mechanistic representation of these processes into models. This would be a much-needed improvement on the current use of empirical relationships and could lead to a step change in our predictive capability.

Data accessibility. This article does not contain any additional data.

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References

1. Ciais P et al. 2014 Carbon and other biogeochemical cycles. Climate change 2013: the physical science basis. In Contribution of working group I to the fith assessment report of the intergovernamental panel on climate change, pp. 465–570. Cambridge, UK: Cambridge University Press.

2. Myhre G et al. 2013 Climate change 2013: the physical science basis. In Contribution of working group I to the fith assessment report of the intergovernamental panel on climate change (eds K Tignor, SK Allen, J Boschung, A Nauels, Y Xia, V Bex, PM Midgley). Cambridge, UK: Cambridge University Press.

3. Fuge R, Johnson CC. 2015 Iodine and human health, the role of environmental geochemistry and diet, a review. Appl. Geochem. 63, 282–302. (doi:10.1016/j.apgeochem.2015.09.013)

4. Charlson RJ, Lovelock JE, Andreae MO, Warren SG. 1987 Oceanic phytoplankton, atmospheric sulphur, cloud albedo and climate. Nature 326, 655–661. (doi:10.1038/326655a0)

5. Andreae MO, Crutzen PJ. 1997 Atmospheric aerosols: biogeochemical sources and role in atmospheric chemistry. Science 276, 1052–1058. (doi:10.1126/science.276.5315.1052)

6. Korhonen H, Carslaw KS, Spracklen DV, Mann GW, Woodhouse MT. 2008 Influence of oceanic dimethyl sulfide emissions on cloud condensation nuclei concentrations and
seasonality over the remote Southern Hemisphere oceans: a global model study. J. Geophys. Res.-Atmos. 113, 16. (doi:10.1029/2007JD009718
7. Leaitch WR et al. 2013 Dimethyl sulfide control of the clean summertime Arctic aerosol and cloud. Elementa: Sci. Anthropocene 1, 00017.
8. Willis MD et al. 2016 Growth of nucleation mode particles in the summertime Arctic: a case study. Atmos. Chem. Phys. 16, 7663–7679. (doi:10.5194/acp-16-7663-2016)
9. Carpenter L. 2003 Iodine in the marine boundary layer. Chem. Rev. 103, 4953–4962. (doi:10.12952/journal.elementa.000017)
10. Read KA et al. 2008 Extensive halogen-mediated ozone destruction over the tropical Atlantic. Nature 453, 1232–1235. (doi:10.1038/nature07035)
11. Carpenter L, MacDonald SM, Shaw MD, Kumar R, Saunders RW, Parthipan R, Wilson J, Plane JM. 2013 Atmospheric iodine levels influenced by sea surface emissions of inorganic iodine. Nat. Geosci. 6, 108–111. (doi:10.1038/ngeo1687)
12. Hossaini R, Chipperfield MP, Montzka SA, Rap A, Dhomse S, Feng W. 2015 Efficiency of short-lived halogens at influencing climate through depletion of stratospheric ozone. Nat. Geosci. 8, 186–190. (doi:10.1038/ngeo2363)
13. Nightingale PD, Malin G, Liss PS. 1995 Production of chloroform and other low-molecular weight halocarbons by some species of macroalgae. Limnol. Oceanogr. 40, 680–689. (doi:10.4319/lo.1995.40.4.0680)
14. Keng FS-L, Phang S-M, Rahman NA, Leedham EC, Hughes C, Robinson AD, Harris NR, Pyle JA, Sturges WT. 2013 Volatile halocarbon emissions by three tropical brown seaweeds under different irradiances. J. Appl. Phycol. 25, 1377–1386. (doi:10.1007/s10811-013-9990-x)
15. Leedham E, Hughes C, Keng F, Phang S-M, Malin G, Sturges W. 2013 Emission of atmospherically significant halocarbons by naturally occurring and farmed tropical macroalgae. Biogeosciences 10, 3615–3633. (doi:10.5194/bg-10-3615-2013)
16. Shaw SL, Gantt B, Meskhidze N. 2010 Production and emissions of marine isoprene and monoterpenes: a review. Adv. Meteorol. 2010, Article ID 408696. (doi:10.1155/2010/408696)
17. Shakhova N, Semiletov I. 2007 Methane release and coastal environment in the East Siberian Arctic shelf. J. Mar. Syst. 66, 227–243. (doi:10.1016/j.jmarsys.2006.06.006)
18. Naqvi S, Bange H, Farías L, Monteiro P, Scranton M, Zhang J. 2010 Marine hypoxia/anoxia as a source of CH4 and N2O. Biogeosciences 7, 2159–2190. (doi:10.5194/bg-7-2159-2010)
19. Laturnus F. 1996 Volatile halocarbons released from Arctic macroalgae. Mar. Chem. 55, 359–366. (doi:10.1016/S0304-4203(97)89401-7)
20. Stefels J, Steinke M, Turner S, Malin G, Belviso S. 2007 Environmental constraints on the production and removal of the climatically active gas dimethylsulphide (DMS) and implications for ecosystem modelling. Biogeochemistry 83, 245–275. (doi:10.1007/s10533-007-9091-5)
21. Hughes C, Sun S. 2016 Light and brominating activity in two species of marine diatom. Mar. Chem. 181, 1–9. (doi:10.1016/j.marchem.2016.02.003)
22. Dani KGS, Loreto F. 2017 Trade-off between dimethyl sulfide and isoprene emissions from marine phytoplankton. Trends Plant Sci. 22, 361–372. (doi:10.1016/j.tplants.2017.01.006)
23. Martino M, Mills GP, Woeltjen J, Liss PS. 2009 A new source of volatile organoiodine compounds in surface seawater. Geophys. Res. Lett. 36, L01609. (doi:10.1029/2008GL036334)
24. Bindoff NL et al. 2013 Detection and attribution of climate change: from global to regional (Chapter 10). In Climate Change 2013: The Physical Science Basis. IPCC Working Group I Contribution to AR5. Cambridge, UK: Cambridge University Press.
25. Lüthi D et al. 2008 High-resolution carbon dioxide concentration record 650,000–800,000 years before present. Nature 453, 379–382. (doi:10.1038/nature06949)
26. Willeit M, Ganopolski A, Calov R, Broykin V. 2019 Mid-Pleistocene transition in glacial cycles explained by declining CO2 and regolith removal. Sci. Adv. 5, eaav7337. (doi:10.1126/sciadv.aav7337)
27. Caldeira K, Wickett ME. 2003 Anthropogenic carbon and ocean pH. Nature 425, 365. (doi:10.1038/425365a)
28. Orr JC et al. 2005 Anthropogenic ocean acidification over the twenty-first century and its impact on calcifying organisms. Nature 437, 681–686. (doi:10.1038/nature04095)
29. Bopp L et al. 2013 Multiple stressors of ocean ecosystems in the 21st century: projections with CMIP5 models. Biogeosciences 10, 6225–6245. (doi:10.5194/bg-10-6225-2013)
30. Gattuso J-P et al. 2015 Contrasting futures for ocean and society from different anthropogenic CO2 emissions scenarios. Science 349, aac4722.
31. Bates NR et al. 2014 A time-series view of changing surface ocean chemistry due to ocean uptake of anthropogenic CO2 and ocean acidification. Oceanography 27, 126–141. (doi:10.5670/oceanog.2014.16)
32. Hönisch B et al. 2012 The geological record of ocean acidification. Science 335, 1058–1063. (doi:10.1126/science.1208277)
33. Riebesell U, Zondervan I, Rost B, Tortell PD, Zeebe RE, Morel FMM. 2000 Reduced calcification of marine plankton in response to increased atmospheric CO2. Nature 407, 364–367. (doi:10.1038/35030078)
34. Hoegh-Guldberg O et al. 2007 Coral reefs under rapid climate change and ocean acidification. Science 318, 1737–1742. (doi:10.1126/science.1152509)
35. Kroeker KJ, Kordas RL, Crim RN, Singh GG. 2010 Meta-analysis reveals negative yet variable effects of ocean acidification on marine organisms. Ecol. Lett. 13, 1419–1434. (doi:10.1111/j.1461-0248.2010.01518.x)
36. Gangstø R, Joos F, Gehlen M. 2011 Sensitivity of pelagic calcification to ocean acidification. Biogeoosciences 8, 433–458. (doi:10.5194/bg-8-433-2011)
37. Bach LT, Riebesell U, Gutowska MA, Federwisch L, Schulz KG. 2015 A unifying concept of coccolithophore sensitivity to changing carbonate chemistry embedded in an ecological framework. Prog. Oceanogr. 135, 125–138. (doi:10.1016/j.pocean.2015.04.012)
38. Munday PL, Dixson DL, Donelson JM, Jones GP, Pratchett MS, Devitsina GV, Døving KB. 2009 Ocean acidification impairs olfactory discrimination and homing ability of a marine fish. Proc. Natl Acad. Sci. USA 106, 1848–1852. (doi:10.1073/pnas.0809996106)
39. Munday PL, Dixson DL, McCormick MI, Meekan M, Ferrari MC, Chivers DP. 2010 Replenishment of fish populations is threatened by ocean acidification. Proc. Natl Acad. Sci. USA 107, 12 930–12 934. (doi:10.1073/pnas.1004519107)
40. Nagelkerken I, Russell BD, Gillanders BM, Connell SD. 2016 Ocean acidification alters fish populations indirectly through habitat modification. Nat. Clim. Change 6, 89–93. (doi:10.1038/nclimate2757)
41. Lemasson AJ, Fletcher S, Hall-Spencer JM, Knights AM. 2017 Linking the biological impacts of ocean acidification on oysters to changes in ecosystem services: a review. J. Exp. Mar. Biol. Ecol. 492, 49–62. (doi:10.1016/j.jembe.2017.01.019)
42. Gehlen M, Gruber N, Gangstø R, Bopp L, Oschlies A. 2011 Biogeochemical consequences of ocean acidification and feedbacks to the earth system. In Ocean acidification (eds JP Gattuso, I. Hansson), vol. 1, pp. 230–248. Oxford, UK: Oxford University Press
43. Six KD, Kloster S, Ilyina T, Archer SD, Zhang K, Maier-Reimer E. 2013 Global warming amplified by reduced sulphur fluxes as a result of ocean acidification. Nat. Clim. Change 3, 975. (doi:10.1038/nclimate1981)
44. Schwinger J, Tipputra J, Goris N, Six KD, Kirkevag A, Seland O, Heinze C, Ilyina T. 2017 Amplification of global warming through pH dependence of DMS production simulated with a fully coupled Earth system model. Biogeosciences 14, 3633–3648. (doi:10.5194/bg-14-3633-2017)
45. Wannicke N, Frey C, Law CS, Voss M. 2018 The response of the marine nitrogen cycle to ocean acidification. Glob. Change Biol. 24, 5031–5043. (doi:10.1111/gcb.14424)
46. Hutchins DA, Fu F. 2017 Microorganisms and ocean global change. Nat. Microbiol. 2, 17058. (doi:10.1038/nmicrobiol.2017.58)
47. Porzio L, Buia MC, Hall-Spencer JM. 2011 Effects of ocean acidification on macroalgal communities. J. Exp. Mar. Biol. Ecol. 400, 278–287. (doi:10.1016/j.jembe.2011.02.011)
48. Brussaard CPD, Noordeloos AAM, Witte H, Collenteur MCJ, Schulz K, Ludwig A, Riebesell U. 2013 Arctic microbial community dynamics influenced by elevated CO2 levels. Biogeosciences 10, 719–731. (doi:10.5194/bg-10-719-2013)
49. Dutkiewicz S, Morris JJ, Follows MJ, Scott J, Levitan O, Dyhrman ST, Berman-Frank I. 2015 Impact of ocean acidification on the structure of future phytoplankton communities. Nat. Clim. Change 5, 1002–1006. (doi:10.1038/nclimate2722)
50. Schulz KG et al. 2017 Phytoplankton blooms at increasing levels of atmospheric carbon dioxide: experimental evidence for negative effects on prymnesiophytes and positive on small picoeukaryotes. Front. Mar. Sci. 4, 64. (doi:10.3389/fmars.2017.00064)
51. Sett S, Schulz KG, Bach LT, Riebesell U. 2018 Shift towards larger diatoms in a natural phytoplankton assemblage under combined high-CO2 and warming conditions. *J. Plankton Res.* 40, 391–406. (doi:10.1093/plankt/fby018)

52. Riebesell U, Gattuso J-P. 2015 Lessons learned from ocean acidification research. *Nat. Clim. Change* 5, 12–14. (doi:10.1038/nclimate2456)

53. Falkowski PG, Fenchel T, Delong EF. 2008 The microbial engines that drive Earth’s biogeochemical cycles. *Science* 320, 1034–1039. (doi:10.1126/science.1153213)

54. Kolb S, Horn MA, Murrell JC, Knief C. 2017. Editorial: The impact of microorganisms on consumption of atmospheric trace gases. *Front. Microbiol.* 8, 1856. (doi:10.3389/fmicb.2017.01856)

55. Joint I, Doney SC, Karl DM. 2011 Will ocean acidification affect marine microbes? *ISME J.* 5, 1. (doi:10.1038/ismej.2010.79)

56. Oliver AE, Newbold LK, Whiteley AS, van der Gast CJ. 2014 Marine bacterial communities are resistant to elevated carbon dioxide levels. *Environ. Microbiol. Rep.* 6, 574–582. (doi:10.1111/1758-2229.12159)

57. Baltar F, Palovaara J, Vila-Costa M, Salazar G, Calvo E, Pelejero C, Marrasé C, Gasol JM, Pinhassi J. 2015 Response of rare, common and abundant bacterioplankton to anthropogenic perturbations in a Mediterranean coastal site. *FEMS Microbiol. Ecol.* 91, fiv058. (doi:10.1093/femsec/fiv058)

58. Hornick T, Bach LT, Crawford KJ, Spilling K, Achterberg EP, Brussaard CPD, Riebesell U, Grossart HP. 2017 Ocean acidification impacts bacteria-phytoplankton coupling at low-nutrient conditions. *Biogeosciences* 14, 1–15. (doi:10.5194/bg-14-1-2017)

59. Bunse C et al. 2016 Response of marine bacterioplankton pH homeostasis gene expression to elevated CO2. *Nat. Clim. Change* 6, 483–487. (doi:10.1038/nclimate2914)

60. Engel A, Piontek J, Grossart H-P, Riebesell U, Schulz KG, Sperling M. 2014 Impact of CO2 enrichment on organic matter dynamics during nutrient induced coastal phytoplankton blooms. *J. Plankton Res.* 36, 641–657. (doi:10.1093/plankt/fbt125)

61. Kim JM et al. 2010 Enhanced production of oceanic dimethylsulfide resulting from CO2-induced grazing activity in a high CO2 world. *Environ. Sci. Technol.* 44, 8140–8143. (doi:10.1021/es102028k)

62. Burrell TJ, Maas EW, Hulston DA, Law CS. 2017 Variable response to warming and ocean acidification by bacterial processes in different plankton communities. *Aquat. Microb. Ecol.* 79, 49–62. (doi:10.3354/ame01819)

63. Sunda WG, Hardison R, Kiene RP, Bucciarelli E, Harada H. 2007 The effect of nitrogen limitation on cellular DMSP and DMS release in marine phytoplankton: climate feedback implications. *Aquat. Sci.* 69, 341–351. (doi:10.1007/s00027-007-0887-0)

64. Leedham EEC, Phang SM, Sturges WT, Malin G. 2015 The effect of desiccation on the emission of volatile bromocarbons from two common temperate macroalgae. *Biogeosciences* 12, 387–398. (doi:10.5194/bg-12-387-2015)

65. Tokarczyk R, Moore RM. 1994 Production of volatile organohalogens by phytoplankton cultures. *Geophys. Res. Lett.* 21, 285–288. (doi:10.1029/94GL00009)

66. Sunda W, Kieber DJ, Kiene RP, Bucciarelli E, Harada H. 2002 An antioxidant function for DMSP and DMS in marine algae. *Nature* 418, 2849–2854. (doi:10.1038/nature00851)

67. Archer SD, Tarraan GA, Stephens JA, Butcher LJ, Kimmance SA. 2011 Combining cell sorting with gas chromatography to determine phytoplankton group-specific intracellular dimethylsulphoniopropionate. *Aquat. Microb. Ecol.* 62, 109–121. (doi:10.3354/ame01464)
73. Buitenhuis ET, De Baar HJW, Veldhuis MJW. 1999 Photosynthesis and calcification by *Emiliania huxleyi* (Prymnesiophyceae) as a function of inorganic carbon species. *J. Phycol.* **35**, 949–959. (doi:10.1046/j.1529-8817.1999.355094.x)

74. Iglesias-Rodriguez MD *et al.* 2008 Phytoplankton calcification in a high-CO₂ world. *Science* **320**, 336–340. (doi:10.1126/science.1154122)

75. Meyer J, Riebesell U. 2015 Reviews and syntheses: responses of coccolithophores to ocean acidification: a meta-analysis. *Biogeosciences* **12**, 1671. (doi:10.5194/bg-12-1671-2015)

76. Arnold HE, Kerrison P, Steinke M. 2013 Interacting effects of ocean acidification and warming on growth and DMS production in the haptophyte coccolithophore *Emiliania huxleyi*. *Glob. Change Biol.* **19**, 1007–1016. (doi:10.1111/gcb.12105)

77. Avgoustidi V, Nightingale PD, Joint IR, Steinke M, Turner SM, Hopkins FE, Liss PS. 2012 Decreased marine dimethyl sulfide production under elevated CO₂ levels in mesocosm and in vitro studies. *Environ. Chem.* **9**, 399–404. (doi:10.1071/EN111125)

78. Spielmeyer A, Pohnert G. 2012 Influence of temperature and elevated carbon dioxide on the production of dimethylsulfiniopropionate and glycine betaine by marine phytoplankton. *Mar. Environ. Res.* **73**, 62–69. (doi:10.1016/j.marenvres.2011.11.002)

79. Webb AL, Malin G, Hopkins FE, Ho KL, Riebesell U, Schulz KG, Larsen A, Liss PS. 2015 Ocean acidification has different effects on the production of dimethylsulfide and dimethylsulfonpropionate measured in cultures of *Emiliania huxleyi* and a mesocosm study: a comparison of laboratory monocultures and community interactions. *Environ. Chem.* **13**, 314–329. (doi:10.1071/en14268)

80. Hopkins FE, Archer SD. 2014 Consistent increase in dimethyl sulfide (DMS) in response to high CO₂ in five shipboard bioassays from contrasting NW European waters. *Biogeosciences* **11**, 4925–4940. (doi:10.5194/bg-11-4925-2014)

81. Hushherr R *et al.* 2017 Impact of ocean acidification on Arctic phytoplankton blooms and dimethyl sulfide concentration under simulated ice-free and under-ice conditions. *Biogeosciences* **14**, 2407. (doi:10.5194/bg-14-2407-2017)

82. Archer SD, Kimmance SA, Stephens JA, Hopkins FE, Bellerby RGJ, Schulz KG, Piontek J, Engel A. 2013 Contrasting responses of DMS and DMSP to ocean acidification in Arctic waters. *Biogeosciences* **10**, 1893–1908. (doi:10.5194/bg-10-1893-2013)

83. Archer SD, Suffrian K, Posman KM, Bach LT, Matrai PA, Countway PD, Ludwig A, Riebesell U. 2018 Processes that contribute to decreased dimethyl sulfide production in response to ocean acidification in subtropical waters. *Front. Mar. Sci.* **5**, 245. (doi:10.3389/fmars.2018.00245)

84. Hopkins FE, Turner SM, Nightingale PD, Steinke M, Liss PS. 2010 Ocean acidification and marine biogenic trace gas production. *Proc. Natl Acad. Sci. USA* **107**, 760–765. (doi:10.1073/pnas.0907163107)

85. Hopkins FE, Kimmance SA, Stephens JA, Bellerby RGJ, Brussaard CP, Czerny J, Schulz KG, Archer SD. 2013 Response of halocarbons to ocean acidification in the Arctic. *Biogeosciences* **10**, 2331–2345.

86. Park K-T *et al.* 2014 Direct linkage between dimethyl sulfide production and microzooplankton grazing, resulting from prey composition change under high partial pressure of carbon dioxide conditions. *Environ. Sci. Technol.* **48**, 4750–4756. (doi:10.1021/es403355h)

87. Vogt M, Steinke M, Turner S, Paulino A, Meyerhöfer M, Riebesell U, LeQuéré C, Liss P. 2008 Dynamics of dimethylsulphonio propionate and dimethylsulphide under different CO₂ concentrations during a mesocosm experiment. *Biogeosciences* **5**, 407–419. (doi:10.5194/bg-5-407-2008)

88. Webb AL *et al.* 2016 Effect of ocean acidification and elevated fCO₂ on trace gas production by a Baltic Sea summer phytoplankton community. *Biogeosciences* **13**, 4595–4613. (doi:10.5194/bg-13-4595-2016)

89. Bach LT *et al.* 2016 Influence of ocean acidification on a natural winter-to-summer plankton succession: first insights from a long-term mesocosm study draw attention to periods of low nutrient concentrations. *PLoS ONE* **11**, e0159068. (doi:10.1371/journal.pone.0159068)

90. Hopkins FE, Archer SD, Stephens JA, Moore CM, Richier S, Cripps GL, Nightingale PD. 2020 A meta-analysis of microcosm experiments shows that dimethyl sulfide (DMS) production in polar waters is insensitive to ocean acidification. *Biogeosciences* **17**, 1–24. (doi:10.5194/bg-17-163-2020)
91. Wyatt NJ, Kitidis V, Woodward EMS, Rees AP, Widdicombe S, Lohan M. 2010 Effects of high CO2 on the fixed nitrogen inventory of the Western English Channel. J. Plankton Res. 32, 631–641. (doi:10.1093/plankt/fbp140)

92. Rees AP, Brown IJ, Jayakumar A, Ward BB. 2016 The inhibition of N2O production by ocean acidification in cold temperate and polar waters. Deep Sea Res. Part II 127, 93–101. (doi:10.1016/j.dsr2.2015.12.006)

93. Curson ARJ et al. 2018 DSYB catalyses the key step of dimethylsulfoniopropionate biosynthesis in many phytoplankton. Nat. Microbiol. 4, 540–542.

94. Galí M, Simó R, Vilà-Costa M, Ruiz-González C, Gasol JM, Matrai P. 2013 Diel patterns of oceanic dimethylsulfide (DMS) cycling: microbial and physical drivers. Global Biogeochem. Cycles 27, 620–636. (doi:10.1002/gbc.20047)

95. Galí M, Simó R. 2015 A meta-analysis of oceanic DMS and DMSP cycling processes: disentangling the summer paradox. Global Biogeochem. Cycles 29, 496–515. (doi:10.1002/2014GB004940)

96. Enami S, Sakamoto Y, Hara K, Osada K, Hoffmann MR, Colussi AJ. 2016 ‘Sizing’ heterogeneous chemistry in the conversion of gaseous dimethyl sulfide to atmospheric particles. Environ. Sci. Technol. 50, 1834–1843. (doi:10.1021/acs.est.5b05337)

97. Fiddes SL, Woodhouse MT, Nicholls Z, Lane TP, Schofield R. 2018 Cloud, precipitation and radiation responses to large perturbations in global dimethyl sulfide. Atmos. Chem. Phys. 18, 10 177–10 198. (doi:10.5194/acp-18-10177-2018)

98. Mahajan AS, Fadnavis S, Thomas MA, Pozzoli L, Gupta S, Royer S-J, Saiz-Lopez A, Simó R. 2015 Quantifying the impacts of an updated global dimethyl sulfide climatology on microwaves and aerosol radiative forcing. J. Geophys. Res.: Atmos. 120, 2524–2536. (doi:10.1002/2014JD022687)

99. Etminan M, Myhre G, Highwood E, Shine K. 2016 Radiative forcing of carbon dioxide, methane, and nitrous oxide: a significant revision of the methane radiative forcing. Geophys. Res. Lett. 43, 614–623. (doi:10.1002/2016GL071930)

100. Brooks SD, Thornton DC. 2018 Marine aerosols and clouds. Annu. Rev. marine science 10, 289–313. (doi:10.1146/annurev-marine-121916-063148)

101. Galí M, Levasseur M, Devred E, Simó R, Babin M. 2018 Sea-surface dimethylsulfide (DMS) concentration from satellite data at global and regional scales. Biogeosciences 15, 3497–3519. (doi:10.5194/bg-15-3497-2018)

102. Sanchez KJ et al. 2018 Substantial seasonal contribution of observed biogenic sulfate particles to cloud condensation nuclei. Sci. Rep. 8, 1–14. (doi:10.1038/s41598-017-17765-5)

103. von Glasow R, Crutzen PJ. 2004 Model study of multiphase DMS oxidation with a focus on halogens. Atmos. Chem. Phys. 4, 589–608. (doi:10.5194/acp-4-589-2004)

104. Johnson M, Bell T. 2008 Coupling between dimethylsulfide emissions and the ocean-atmosphere exchange of ammonia. Environ. Chem. 5, 259–267. (doi:10.1071/EN08030)

105. Chen T, Jang M. 2012 Secondary organic aerosol formation from photooxidation of a mixture of dimethyl sulfide and isoprene. Atmos. Environ. 46, 271–278. (doi:10.1016/j.atmosenv.2011.09.082)

106. Kerrison P, Suggett DJ, Hepburn LJ, Steinke M. 2012 Effect of elevated pCO2 on the production of dimethylsulfinopropionate (DMSP) and dimethylsulphide (DMS) in two species of Ulva (Chlorophyceae). Biogeochemistry 110, 5–16. (doi:10.1007/s10533-012-9707-2)

107. Richier S, Achterberg EP, Humphreys MP, Poulton AJ, Suggett DJ, Tyrrell T, Moore CM. 2018 Geographical CO2 sensitivity of phytoplankton correlates with ocean buffer capacity. Glob. Change Biol. 24, 4438–4452. (doi:10.1111/gcb.14324)

108. Sabine CL et al. 2004 The oceanic sink for anthropogenic CO2. Science 305, 367–371. (doi:10.1126/science.1079403)

109. Lewis CN, Brown KA, Edwards LA, Cooper G, Findlay HS. 2013 Sensitivity to ocean acidification parallels natural pCO2 gradients experienced by Arctic copepods under winter sea ice. Proc. Natl Acad. Sci. USA 110, E4960–E4967. (doi:10.1073/pnas.1315162110)

110. Thoisen C, Riisgaard K, Lundholm N, Nielsen TG, Hansen PJ. 2015 Effect of acidification on an Arctic phytoplankton community from Disko Bay, West Greenland. Mar. Ecol. Prog. Ser. 520, 21–34. (doi:10.3354/meps11123)

111. Tynan E et al. 2016 Physical and biogeochemical controls on the variability in surface pH and calcium carbonate saturation states in the Atlantic sectors of the Arctic and Southern Oceans. Deep Sea Res. Part II 127, 7–27. (doi:10.1016/j.dsr2.2016.01.001)
112. Popova E, Yool A, Aksenov Y, Coward AC, Anderson TR. 2014 Regional variability of acidification in the Arctic: a sea of contrasts. *Biogeosciences* **11**, 293–308. (doi:10.5194/bg-11-293-2014)
113. Pachauri RK et al. 2014 Climate change 2014: synthesis report. In Contribution of working groups I, II and III to the fifth assessment report of the intergovernmental panel on climate change. lpc.
114. Bénard R et al. 2019 Contrasting effects of acidification and warming on dimethylsulfide concentrations during a temperate estuarine fall bloom mesocosm experiment. *Biogeosciences* **16**, 1167–1185. (doi:10.5194/bg-16-1167-2019)
115. Paulot F, Jacob DJ, Johnson MT, Bell TG, Baker AR, Keene WC, Lima ID, Doney SC, Stock CA. 2015 Global oceanic emission of ammonia: constraints from seawater and atmospheric observations. *Global Biogeochem. Cycles* **29**, 1165–1178. (doi:10.1002/2015GB005106)
116. Jickells T et al. 2017 A reevaluation of the magnitude and impacts of anthropogenic atmospheric nitrogen inputs on the ocean. *Global Biogeochem. Cycles* **31**, 289–305.
117. Quinn PK, Charlson RJ, Zoller WH. 1987 Ammonia, the dominant base in the remote marine troposphere: a review. *Tellus B* **39**, 413–425. (doi:10.3402/tellusb.v39i5.15359)
118. Johnson MT et al. 2008 Field observations of the ocean-atmosphere exchange of ammonia: fundamental importance of temperature as revealed by a comparison of high and low latitudes. *Global Biogeochem. Cycles* **22**, GB1019. (doi:10.1029/2007GB003039)
119. Kulmala M, Petäjä T, Ehn M, Thornton J, Sipilä M, Worsnop D, Kerminen V-M. 2014 Chemistry of atmospheric nucleation: on the recent advances on precursor characterization and atmospheric cluster composition in connection with atmospheric new particle formation. *Annu. Rev. Phys. Chem.* **65**, 21–37. (doi:10.1146/annurev-physchem-040412-110014)
120. Beman JM, Chow C-E, King AL, Feng Y, Fuhrman JA, Andersson A, Bates NR, Popp BN, Hutchins DA. 2011 Global declines in oceanic nitrification rates as a consequence of ocean acidification. *Proc. Natl Acad. Sci. USA* **108**, 208–213. (doi:10.1073/pnas.1011053108)
121. Das S, Mangwani N. 2015 Ocean acidification and marine microorganisms: responses and consequences. *Oceanologica* **57**, 349–361. (doi:10.1016/j.oceano.2015.07.003)
122. Gu X, Li K, Pang K, Ma Y, Wang X. 2017 Effects of pH on the growth and NH4-N uptake of *Skeletonema costatum* and *Nitzschia closterium*. *Mar. Pollut. Bull.* **124**, 946–952. (doi:10.1016/j.marpolbul.2017.01.048)
123. Hackenberg SC et al. 2017 Basin-scale observations of monoterpenes in the Arctic and Atlantic Oceans. *Environ. Sci. Technol.* **51**, 10 449–10 458. (doi:10.1021/acs.est.7b02240)
124. Kim MJ, Novak GA, Zoerb MC, Yang M, Blomquist BW, Huebert BJ, Cappa CD, Bertram TH. 2017 Air-sea exchange of biogenic volatile organic compounds and the impact on aerosol particle size distributions. *Geophys. Res. Lett.* **44**, 3887–3896. (doi:10.1002/2017GL072975)
125. Montzka S et al. 2011 Scientific Assessment of Ozone Depletion: 2010, Global Ozone Research and Monitoring Project-Report No. 52. Geneva, Switzerland: World Meteorological Organization.
126. Del Monaco C, Hay ME, Garrell P, Mumby PJ, Diaz-Pulido G. 2017 Effects of ocean acidification on the potency of macroalgal allelopathy to a common coral. *Sci. Rep.* **7**, 41053. (doi:10.1038/srep41053)
127. Hegglin MI, D作出 painful details about the corrupted data. -snow et al. 2011 Contribution of very short-lived organic substances to stratospheric chlorine and bromine in the tropics—a case study. *Atmos. Chem. Phys.* **8**, 7325–7334. (doi:10.5194/acp-8-7325-2008)
133. Solomon S, Garcia RR, Ravishankara AR. 1994 On the role of iodine in ozone depletion. *J. Geophys. Res.* 99(D10), 20 491–20 499. (doi:10.1029/94JD02028)

134. Davis D, Crawford J, Liu S, McKeen S, Bandy A, Thornton D, Rowland F, Blake D. 1996 Potential impact of iodine on tropospheric levels of ozone and other critical oxidants. *J. Geophys. Res.* 101, 2135–2147. (doi:10.1029/95JD02727)

135. Glasov RV, Sander R. 2002 Modelling halogen chemistry in the marine boundary layer 1. Cloud-free MBL. *J. Geophys. Res.* 107, 942. (doi:10.1029/2001JD000942)

136. Makela JM et al. 2002 Biogenic iodine emissions and identification of end-products in coastal ultrafine particles during nucleation bursts. *J. Geophys. Res.* 107(D19), 8110. (doi:10.1029/2001JD000580)

137. Lim Y-K, Phang S-M, Abdul Rahman N, Sturges WT, Malin G. 2017 Halocarbon emissions from marine phytoplankton and climate change. *Int. J. Environ. Sci. Technol.* 14, 1355–1370. (doi:10.1007/s13762-016-1219-5)

138. Mithoo-Singh PK, Keng FS-L, Phang S-M, Elvidge ECL, Sturges WT, Malin G, Rahman NA. 2017 Halocarbon emissions by selected tropical seaweeds: species-specific and compound-specific responses under changing pH. *PeerJ* 5, e2918. (doi:10.7717/peerj.2918)

139. Jacob DJ et al. 2014 Update on ozone-depleting substances (ODSs) and other gases of interest to the Montreal protocol. In *Scientific assessment of ozone depletion*, pp. 1.1–1.101.

140. Fassbender AJ, Sabine CL, Feifel KM. 2016 Consideration of coastal carbonate chemistry in understanding biological calcification. *Geophys. Res. Lett.* 43, 4467–4476. (doi:10.1002/2016GL068860)

141. Jacob DJ, Field BD, Jin EM, Bey I, Li Q, Logan JA, Yantosca RM, Singh HB. 2002 Atmospheric budget of acetone. *J. Geophys. Res. Atmos.* 107, ACH 5-1–ACH 5-17. (doi:10.1029/2001JD000694)

142. Jacob DJ et al. 2005 Global budget of methanol: constraints from atmospheric observations. *J. Geophys. Res. Atmos.* 110(D8), n/a–n/a. (doi:10.1029/2004JD005172)

143. Marandino CA, de Bruyn WJ, Miller SD, Prather MJ, Saltzman ES. 2005 Oceanic uptake and the global atmospheric acetone budget. *Geophys. Res. Lett.* 32, 15. (doi:10.7717/peerj.2918)

144. Beale R, Liss PS, Nightingale PD. 2010 First oceanic measurements of ethanol and propanol. *Geophys. Res. Lett.* 37, L24607. (doi:10.1029/2010GL045534)

145. Dixon JL, Beale R, Nightingale PD. 2010 Microbial methanol uptake in northeast Atlantic waters. *ISME J.* 5, 704–716. (doi:10.1038/ismej.2010.169)

146. Beale R, Dixon JL, Arnold SR, Liss PS, Nightingale PD. 2013 Methanol, acetaldehyde, and acetone in the surface waters of the Atlantic Ocean. *J. Geophys. Res. Oceans* 118, 5412–5425. (doi:10.1002/jgrc.203285)

147. Dixon JL, Beale R, Nightingale PD. 2013 Production of methanol, acetaldehyde, and acetone in the surface waters of the Atlantic Ocean. *J. Geophys. Res. Oceans* 118, 5412–5425. (doi:10.1002/jgrc.203285)

148. Yang M, Beale R, Smyth T, Blomquist B. 2013 Measurements of OVOC fluxes by eddy covariance using a proton-transfer-reaction mass spectrometer—method development at a coastal site. *Atmos. Chem. Phys.* 13, 6165–6184. (doi:10.5194/acp-13-6165-2013)

149. Schlundt C, Tegtmeier S, Lennartz ST, Bracher A, Cheah W, Krüger K, Quack B, Marandino CA. 2017 Oxygenated volatile organic carbon in the western Pacific convective centre: ocean cycling, air–sea gas exchange and atmospheric transport. *Atmos. Chem. Phys.* 17, 10 837–10 854. (doi:10.5194/acp-17-10837-2017)

150. Carpenter LJ, Archer SD, Beale R. 2012 Ocean-atmosphere trace gas exchange. *Chem. Soc. Rev.* 41, 6473–6506. (doi:10.1039/c2cs35121h)

151. Carpenter LJ, Nightingale PD. 2015 Chemistry and release of gases from the surface ocean. *Chem. Rev.* 115, 4015–4034. (doi:10.1021/cr5007123)

152. Beale R, Dixon JL, Smyth TJ, Nightingale PD. 2015 Annual study of oxygenated volatile organic compounds in UK shelf waters. *Mar. Chem.* 171, 96–106. (doi:10.1016/j.marchem.2015.02.013)

153. Mopper K, Stahovec WL. 1986 Sources and sinks of low molecular weight organic carbonyl compounds in seawater. *Mar. Chem.* 19, 305–321. (doi:10.1016/0304-4203(86)90052-6)

154. Kieber RJ, Zhou X, Mopper K. 1990 Formation of carbonyl compounds from UV-induced photodegradation of humic substances in natural waters: fate of riverine carbon in the sea. *Limnol. Oceanogr.* 35, 1503–1515. (doi:10.4319/lo.1990.35.7.1503)
155. Zhou X, Mopper K. 1997 Photochemical production of low-molecular-weight carbonyl compounds in seawater and surface microlayer and their air-sea exchange. *Mar. Chem.* **56**, 201–213. (doi:10.1016/S0304-4203(96)00076-X)

156. de Bruyn WJ, Clark CD, Pagel L, Takehara C. 2011 Photochemical production of formaldehyde, acetaldehyde and acetone from chromophoric dissolved organic matter in coastal waters. *J. Photochem. Photobiol., A* **226**, 16–22. (doi:10.1016/j.jphotochem.2011.10.002)

157. Sieburth JM, Keller MD. 1989 Methylaminotrophic bacteria in xenic nanoalgal cultures: incidence, significance, and role of methylated algal osmoprotectants. *Biol. Oceanogr.* **6**, 383–395.

158. Nightingale PD. 1991 *Low molecular weight halocarbons in seawater*. Norwich, UK: School of environmental sciences, University of East Anglia.

159. Heikes BG *et al.* 2002 Atmospheric methanol budget and ocean implication. *Global Biogeochem. Cycles** **16**, 80-1–80-13. (doi:10.1029/2002GB001895)

160. Dixon J, Beale R, Nightingale P. 2011 Rapid biological oxidation of methanol in the tropical Atlantic: significance as a microbial carbon source. *Biogeosciences** **8**, 2707–2716. (doi:10.5194/bg-8-2707-2011)

161. Nevison CD, Weiss RF, Erickson DJ. 1995 Global oceanic emissions of nitrous oxide. *J. Geophys. Res. Oceans** **100**, 15 809–15 820. (doi:10.1029/95JC00684)

162. Menon S *et al.* 2007 Couplings between changes in the climate system and biogeochemistry. Berkeley, CA: Ernest Orlando Lawrence Berkeley National Laboratory.

163. Ravishankara AR, Daniel JS, Portmann RW. 2009 Nitrous oxide (N2O): the dominant ozone-depleting substance emitted in the 21st century. *Science** **326**, 123–125. (doi:10.1126/science.1176985)

164. Babbin AR, Bianchi D, Jayakumar A, Ward BB. 2015 Rapid nitrous oxide cycling in the suboxic ocean. *Science** **348**, 1127–1129. (doi:10.1126/science.aaa8380)

165. Freing A, Wallace DW, Bange HW. 2012 Global oceanic production of nitrous oxide. *Phil. Trans. R. Soc. B** **367**, 1245–1255. (doi:10.1098/rstb.2011.0360)

166. Kitidis V, Laverock B, McNeill LC, Beesley A, Cummings D, Tait K, Osborn MA, Widdicombe S. 2011 Impact of ocean acidification on benthic and water column ammonia oxidation. *Geophys. Res. Lett.* **38**, n/a–n/a. (doi:10.1029/2011GL049095)

167. Kitidis V, Tilstone GH, Smyth TJ, Torres R, Law CS. 2011 Carbon monoxide emission from a Mauritanian upwelling filament. *Mar. Chem** **127**, 123–133. (doi:10.1016/j.marchem.2011.08.004)

168. Clark DR, Brown IJ, Rees AP, Somerfield PJ, Miller PI. 2014 The influence of ocean acidification on nitrogen regeneration and nitrous oxide production in the northwest European shelf sea. *Biogeosciences** **11**, 4985–5005. (doi:10.5194/bg-11-4985-2014)

169. Fulweiler RW, Emery HE, Heiss EM, Berounsky VM. 2011 Assessing the role of pH in determining water column nitrification rates in a coastal system. *Estuaries Coasts** **34**, 1095. (doi:10.1007/s12237-011-9432-4)

170. Codispoti LA. 2010 Interesting times for marine N2O. *Science** **327**, 1339–1340. (doi:10.1126/science.1184945)

171. Hutchins DA, Mulholland MR, Fu F. 2009 Nutrient cycles and marine microbes in a CO2-enriched ocean. *Oceanography** **22**, 128–145. (doi:10.5670/oceanog.2009.103)

172. Bowen JL, Kearns PJ, Holcomb M, Ward BB. 2013 Acidification alters the composition of ammonia-oxidizing microbial assemblages in marine mesocosms. *Mar. Ecol. Prog. Ser.** **492**, 1–8. (doi:10.3354/meps10526)

173. Santoro AE, Buchwald C, McIlvin MR, Casciotti KL. 2011 Isotopic signature of N2O produced by marine ammonia-oxidizing archaea. *Science** **333**, 1282–1285. (doi:10.1126/science.1208239)

174. Loescher CR, Kock A, Koenneke M, LaRoche J, Bange HW, Schmitz RA. 2012 Production of oceanic nitrous oxide by ammonia-oxidizing archaea. *Biogeosciences** **9**, 2419–2429. (doi:10.5194/bg-9-2419-2012)

175. Wuchter C *et al.* 2006 Archaeal nitrification in the ocean. *Proc. Natl Acad. Sci. USA** **103**, 12317–12322. (doi:10.1073/pnas.0600756103)
177. Qin W et al. 2014 Marine ammonia-oxidizing archaeal isolates display obligate mixotrophy and wide ecotypic variation. Proc. Natl Acad. Sci. USA 111, 12504–12509. (doi:10.1073/pnas.1324115111)
178. Pan Y, Ye L, Ni B-J, Yuan Z. 2012 Effect of pH on N₂O reduction and accumulation during denitrification by methanol utilizing denitrifiers. Water Res. 46, 4832–4840. (doi:10.1016/j.watres.2012.06.003)
179. Stocker TF et al. 2013 IPCC, 2013: climate change 2013: the physical science basis. In Contribution of working group I to the fifth assessment report of the intergovernmental panel on climate change. Cambridge, UK: Cambridge University Press.
180. WMO. 2016 The global climate in 2011–2016.
181. Wuebbles DJ, Hayhoe K. 2002 Atmospheric methane and global change. Earth Sci. Rev. 57, 177–210. (doi:10.1016/S0012-8252(01)00062-9)
182. Grunwald M, Dellwig O, Beck M, Dippner JW, Freund JA, Kohlmeier C, Schnetger B, Brumsack H-J. 2009 Methane in the southern North Sea: sources, spatial distribution and budgets. Estuar. Coast. Shelf Sci. 81, 445–456. (doi:10.1016/j.ecss.2008.11.021)
183. Forster G, Upstill-Goddard RC, Gist N, Robinson C, Uher G, Woodward EMS. 2009 Nitrous oxide and methane in the Atlantic Ocean between 50 N and 52 S: latitudinal distribution and sea-to-air flux. Deep Sea Res. Part II 56, 964–976. (doi:10.1016/j.dsr2.2008.12.002)
184. Upstill-Goddard RC, Barnes J. 2016 Methane emissions from UK estuaries: re-evaluating the estuarine source of tropospheric methane from Europe. Mar. Chem. 180, 14–23. (doi:10.1016/j.marchem.2016.01.010)
185. Kock A, Gebhardt S, Bange HW. 2008 Methane emissions from the upwelling area off Mauritania (NW Africa). Biogeosciences 5, 1119–1125. (doi:10.5194/bg-5-1119-2008)
186. Kiene RP. 1991 Production and consumption of methane in aquatic systems. In Microbial production and consumption of greenhouse gases: methane, nitrogen oxides, and halomethanes (eds JE Rogers, WB Whitman), pp. 111–146. Washington, DC: American Society for Microbiology.
187. Conte L, Szopa S, Séférian R, Bopp L. 2019 The oceanic cycle of carbon monoxide and its emissions to the atmosphere. Biogeosciences 16, 881–902. (doi:10.5194/bg-16-881-2019)
188. Uher G, Andreae MO. 1997 Photochemical production of carbonyl sulfide in North Sea water: a process study. Limnol. Oceanogr. 42, 432–442. (doi:10.4319/lo.1997.42.3.0432)
189. Watts SF. 2000 The mass budgets of carbonyl sulfide, dimethyl sulfide, carbon disulfide and hydrogen sulfide. Atmos. Environ. 34, 761–779. (doi:10.1016/S1352-2310(99)00342-8)
190. Kettle A, Kuhn U, Von Hobel M, Kesselmeier J, Andreae M. 2002 Global budget of atmospheric carbonyl sulfide: temporal and spatial variations of the dominant sources and sinks. J. Geophys. Res.: Atmos. 107(D22), 4658. (doi:10.1029/2001JD001290)
191. Bruhl C, Leliieveld J, Crutzen P, Tost H. 2012 The role of carbonyl sulphide as a source of stratospheric sulphate aerosol and its impact on climate. Atmos. Chem. Phys. 12, 1239–1253. (doi:10.5194/acp-12-1239-2012)
192. Lennartz ST et al. 2017 Direct oceanic emissions unlikely to account for the missing source of atmospheric carbonyl sulfide. Atmos. Chem. Phys. 17, 385–402. (doi:10.5194/acp-17-385-2017)
201. Steinacher M, Joos F, Frollicher TL, Plattner GK, Doney SC. 2009 Imminent ocean acidification in the Arctic projected with the NCAR global coupled carbon cycle-climate model. Biogeosciences 6, 515–533. (doi:10.5194/bg-6-515-2009)

202. Comiso JC. 2012 Large decadal decline of the Arctic multiyear ice cover. J. Climate 25, 1176–1193. (doi:10.1175/JCLI-D-11-00113.1)

203. Shadwick E, Trull T, Thomas H, Gibson J. 2013 Vulnerability of polar oceans to anthropogenic acidification: comparison of Arctic and Antarctic seasonal cycles. Sci. Rep. 3, 2339. (doi:10.1038/srep02339)

204. Semiletov I et al. 2016 Acidification of East Siberian Arctic Shelf waters through addition of freshwater and terrestrial carbon. Nat. Geosci. 9, 361–365. (doi:10.1038/ngeo2695)

205. Doney SC, Fabry VJ, Feely RA, Kleypas JA. 2009 Ocean acidification: the other CO2 problem. Annu. Rev. Mar. Sci. 1, 169–192. (doi:10.1146/annurev.marine.010908.163834)

206. Fabry VJ, McClintock JB, Mathis JT, Grebmeier JM. 2009 Ocean acidification at high latitudes: the bellweather. Oceanography 22, 160. (doi:10.5670/oceanog.2009.105)

207. McNeil BI, Matear R. 2008 Southern Ocean acidification: a tipping point at 450-ppm atmospheric CO2. Proc. Natl Acad. Sci. USA 105, 18860–18864. (doi:10.1073/pnas.0806318105)

208. Hauri C, Friedrich T, Timmermann A. 2016 Abrupt onset and prolongation of aragonite undersaturation events in the Southern Ocean. Nat. Clim. Change 6, 172–176. (doi:10.1038/nclimate2844)

209. Chang RYW et al. 2011 Aerosol composition and sources in the central Arctic Ocean during ASCOS. Atmos. Chem. Phys. 11, 10619–10636. (doi:10.5194/acp-11-10619-2011)

210. Collins DB et al. 2017 Frequent ultrafine particle formation and growth in the Canadian Arctic marine environment. Atmos. Chem. Phys. 17, 13119. (doi:10.5194/acp-17-13119-2017)

211. Jarníková T, Tortell PD. 2016 Towards a revised climatology of summertime dimethylsulfide concentrations and sea–air fluxes in the Southern Ocean. Environ. Chem. 13, 364–378. (doi:10.1071/EN14272)

212. McCoy DT, Burrows SM, Wood R, Grosvenor DP, Elliott SM, Ma P-L, Rasch PJ, Hartmann DL. 2015 Natural aerosols explain seasonal and spatial patterns of Southern Ocean cloud albedo. Sci. Adv. 1, e1500157. (doi:10.1126/sciadv.1500157)

213. ACIA. 2005 ACIA: Arctic climate impact assessment. Cambridge, UK: Cambridge University Press.

214. Kapsenberg L, Kelley AL, Shaw EC, Martz TR, Hofmann GE. 2015 Near-shore Antarctic pH variability has implications for the design of ocean acidification experiments. Sci. Rep. 5, 9638. (doi:10.1038/srep09638)

215. Hoppe CJM, Wolf KKE, Schuback N, Tortell PD, Rost B. 2018 Compensation of ocean acidification effects in Arctic phytoplankton assemblages. Nat. Clim. Change 8, 529–533. (doi:10.1038/s41558-018-0142-9)

216. Feely RA, Sabine CL, Hernandez-Ayon JM, Ianson D, Hales B. 2008 Evidence for upwelling of corrosive ‘acidified’ water onto the continental shelf. Science 320, 1490–1492. (doi:10.1126/science.1155676)

217. Osborne EB, Thunell RC, Gruber N, Feely RA, Benitez-Nelson CR. 2020 Decadal variability in twentieth-century ocean acidification in the California Current Ecosystem. Nat. Geosci. 13, 43–49. (doi:10.1038/s41561-019-0499-z)

218. Weeks SJ, Currie B, Bakun A. 2002 Massive emissions of toxic gas in the Atlantic. Nature 415, 493–494. (doi:10.1038/415493b)

219. Nevison CD, Lueker TJ, Weiss RF. 2004 Quantifying the nitrous oxide source from coastal upwelling. Global Biogeochem. Cycles 18, GB1018. (doi:10.1029/2003GB002110)

220. Gutknecht E et al. 2013 Coupled physical/biogeochemical modeling including O2-dependent processes in the Eastern Boundary Upwelling Systems: application in the Benguela. Biogeosciences 10, 3559–3591. (doi:10.5194/bg-10-3559-2013)

221. Law CS et al. 2013 Evolving research directions in surface ocean–lower atmosphere (SOLAS) science. Environ. Chem. 10, 1–16. (doi:10.1071/EN12159)

222. Arevalo-Martínez DL, Kock A, Löscher C, Schmitz RA, Bange HW. 2015 Massive nitrous oxide emissions from the tropical South Pacific Ocean. Nat. Geosci. 8, 530–533. (doi:10.1038/ngeo2469)

223. Löscher C et al. 2016 Water column biogeochemistry of oxygen minimum zones in the eastern tropical North Atlantic and eastern tropical South Pacific oceans. Biogeosciences (BG) 13, 3585–3606. (doi:10.5194/bg-13-3585-2016)
224. Ohde T, Dadou I. 2018 Seasonal and annual variability of coastal sulphur plumes in the northern Benguela upwelling system. *PLoS ONE* 13, e0192140. (doi:10.1371/journal.pone.0192140)

225. Feely RA et al. 2016 Chemical and biological impacts of ocean acidification along the west coast of North America. *Estuar. Coast. Shelf Sci.* 183, 260–270. (doi:10.1016/j.ecss.2016.08.043)

226. Feely RA, Okazaki RR, Cai W-J, Bednarsek N, Alin SR, Byrne RH, Fassbender A. 2018 The combined effects of acidification and hypoxia on pH and aragonite saturation in the coastal waters of the California current ecosystem and the northern Gulf of Mexico. *Cont. Shelf Res.* 152, 50–60. (doi:10.1016/j.csr.2017.11.002)

227. Bednarsek N, Feely R, Beck M, Glippa O, Kanerva M, Engström-Öst J. 2018 El Niño-related thermal stress coupled with upwelling-related ocean acidification negatively impacts cellular to population-level responses in pteropods along the California Current System with implications for increased bioenergetic costs. *Front. Mar. Sci.* 5, 486. (doi:10.3389/fmars.2018.00486)

228. Fennel K et al. 2019 Advancing marine biogeochemical and ecosystem reanalyses and forecasts as tools for monitoring and managing ecosystem health. *Front. Mar. Sci.* 6, 89. (doi:10.3389/fmars.2019.00089)

229. Frame CH, Lau E, Nolan IV EJ, Goepfert TJ, Lehmann MF. 2017 Acidification enhances hybrid N₂O production associated with aquatic ammonia-oxidizing microorganisms. *Front. Microbiol.* 7, 2104. (doi:10.3389/fmicb.2016.02104)

230. López-Urrutia Á, San Martin E, Harris RP, Irigoien X. 2006 Scaling the metabolic balance of the oceans. *Proc. Natl Acad. Sci. USA* 103, 8739–8744. (doi:10.1073/pnas.0601137103)

231. O’Connor MI, Piehler MF, Leech DM, Anton A, Bruno JF. 2009 Warming and resource availability shift food web structure and metabolism. *PLoS Biol.* 7, e1000178. (doi:10.1371/journal.pbio.1000178)

232. Capotondi A, Alexander MA, Bond NA, Curchitser EN, Scott JD. 2012 Enhanced upper ocean stratification with climate change in the CMIP3 models. *J. Geophys. Res. Oceans* 117, C04031. (doi:10.1029/2011JC007409)

233. Wang R et al. 2015 Influence of anthropogenic aerosol deposition on the relationship between oceanic productivity and warming. *Geophys. Res. Lett.* 42, 10745–10754. (doi:10.1002/2015GL066753)

234. Gruber N. 2011 Warming up, turning sour, losing breath: ocean biogeochemistry under global change. *Phil. Trans. R. Soc. A* 369, 1980–1996. (doi:10.1098/rsta.2011.0003)

235. Rheuban JE, Doney SC, McCorkle DC, Jakuba RW. 2019 Quantifying the effects of nutrient enrichment and freshwater mixing on coastal ocean acidification’. *J. Geophys. Res. Oceans* 124, 9085–9100. (doi:10.1029/2019JC015556)

236. Barton AD, Irwin AJ, Finkel ZV, Stock CA. 2016 Anthropogenic climate change drives shift and shuffle in North Atlantic phytoplankton communities. *Proc. Natl Acad. Sci. USA* 113, 2964–2969. (doi:10.1073/pnas.1519080113)

237. Pollimene L, Archer SD, Butenschön M, Allen JJ. 2012 A mechanistic explanation of the Sargasso Sea DMS ‘summer paradox’. *Biogeochemistry* 110, 243–255. (doi:10.1007/s10533-011-9674-z)

238. Martinez-Rey J, Bopp L, Gehlen M, Tagliabue A, Gruber N. 2015 Projections of oceanic N₂O emissions in the 21st century using the IPSL Earth system model. *Biogosciences* 12, 4133–4148. (doi:10.5194/bg-12-4133-2015)

239. Landolfi A, Somes C, Koeve W, Zamora LM, Oschlies A. 2017 Oceanic nitrogen cycling and N₂O flux perturbations in the Anthropocene. *Global Biogeochem. Cycles* 31, 1236–1255. (doi:10.1002/2017gb005633)

240. Battaglia G, Joos F. 2018 Marine N₂O emissions from nitrification and denitrification constrained by modern observations and projected in multimillennial global warming simulations. *Global Biogeochem. Cycles* 32, 92–121. (doi:10.1002/2017GB005671)

241. Andrews OD, Bindoff NL, Halloran PR, Ilyina T, Le Quéré C. 2013 Detecting an external influence on recent changes in oceanic oxygen using an optimal fingerprinting method. *Biogosciences* 10, 1799–1813. (doi:10.5194/bg-10-1799-2013)

242. Oschlies A, Duteil O, Getzlaff J, Koeve W, Landolfi A, Schmidtko S. 2017 Patterns of deoxygenation: sensitivity to natural and anthropogenic drivers. *Phil. Trans. R. Soc. A* 375, 20160325. (doi:10.1098/rsta.2016.0325)

243. Riebesell U et al. 2007 Enhanced biological carbon consumption in a high CO₂ ocean. *Nature* 450, 545–548. (doi:10.1038/nature06267)
244. Oschlies A, Schulz KG, Riebesell U, Schmittner A. 2008 Simulated 21st century’s increase in oceanic suboxia by CO2-enhanced biotic carbon export. *Global Biogeochem. Cycles* **22**, GB4008. (doi:10.1029/2007GB003147)

245. Andrews O, Buitenhuis E, Le Quéré C, Suntharalingam P. 2017 Biogeochemical modelling of dissolved oxygen in a changing ocean. *Phil. Trans. R. Soc. A* **375**, 20160328. (doi:10.1098/rsta.2016.0328)

246. Thingstad TF *et al.* 2008 Counterintuitive carbon-to-nutrient coupling in an Arctic pelagic ecosystem. *Nature* **455**, 387–390. (doi:10.1038/nature07235)

247. Tagliabue A, Bopp L, Gehlen M. 2011 The response of marine carbon and nutrient cycles to ocean acidification: large uncertainties related to phytoplankton physiological assumptions. *Global Biogeochem. Cycles* **25**, GB3017. (doi:10.1029/2010GB003929)

248. Gunderson AR, Armstrong EJ, Stillman JH. 2016 Multiple stressors in a changing world: the need for an improved perspective on physiological responses to the dynamic marine environment. *Annu. Rev. Mar. Sci.* **8**, 357–378. (doi:10.1146/annurev-marine-122414-033953)

249. Boyd PW *et al.* 2018 Experimental strategies to assess the biological ramifications of multiple drivers of global ocean change—a review. *Glob. Change Biol.* **24**, 2239–2261. (doi:10.1111/gcb.14102)

250. Boyd PW, Lennartz ST, Glover DM, Doney SC. 2015 Biological ramifications of climate-change-mediated oceanic multi-stressors. *Nat. Clim. Change* **5**, 71–79. (doi:10.1038/nclimate2441)

251. Lohbeck KT, Riebesell U, Reusch TB. 2012 Adaptive evolution of a key phytoplankton species to ocean acidification. *Nat. Geosci.* **5**, 346. (doi:10.1038/ngeo1441)

252. Lohbeck KT, Riebesell U, Reusch TB. 2014 Gene expression changes in the coccolithophore *Emiliania huxleyi* after 500 generations of selection to ocean acidification. *Proc. R. Soc. B* **281**, 20140003. (doi:10.1098/rspb.2014.0003)

253. Pančić M, Hansen PJ, Tammilehto A, Lundholm N. 2015 Resilience to temperature and pH changes in a future climate change scenario in six strains of the polar diatom *Fragilariopsis cylindrus*. *Biogeosciences* **12**, 4235–4244. (doi:10.5194/bg-12-4235-2015)

254. Wang Y, Zhang R, Zheng Q, Deng Y, Van Nostrand JD, Zhou J, Jiao N. 2016 Bacterioplankton community resilience to ocean acidification: evidence from microbial network analysis. *ICES J. Mar. Sci.* **73**, 865–875. (doi:10.1093/icesjms/fsv187)

255. Riebesell U, Fabry VJ, Hansson L, Gattuso J-P. 2011 *Guide to best practices for ocean acidification research and data reporting*, 258 pp. Luxembourg, Europe: Office for Official Publications of the European Communities.