Consistent increase in dimethyl sulfide (DMS) in response to high CO$_2$ in five shipboard bioassays from contrasting NW European waters

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Abstract. The ubiquitous marine trace gas dimethyl sulfide (DMS) comprises the greatest natural source of sulfur to the atmosphere and is a key player in atmospheric chemistry and climate. We explore the short-term response of DMS production and cycling and that of its algal precursor dimethyl sulfoniopropionate (DMSP) to elevated carbon dioxide (CO$_2$) and ocean acidification (OA) in five 96 h shipboard bioassay experiments. Experiments were performed in June and July 2011, using water collected from contrasting sites in NW European waters (Outer Hebrides, Irish Sea, Bay of Biscay, North Sea). Concentrations of DMS and DMSP, alongside rates of DMSP synthesis and DMS production and consumption, were determined during all experiments for ambient CO$_2$ and three high-CO$_2$ treatments (550, 750, 1000 µatm). In general, the response to OA throughout this region showed little variation, despite encompassing a range of biological and biogeochemical conditions. We observed consistent and marked increases in DMS concentrations relative to ambient controls (110 % (28–223 %) at 550 µatm, 153 % (56–295 %) at 750 µatm and 225 % (79–413 %) at 1000 µatm), and decreases in DMSP concentrations (28 % (18–40 %) at 550 µatm, 44 % (18–64 %) at 750 µatm and 52 % (24–72 %) at 1000 µatm). Significant decreases in DMSP synthesis rate constants ($\mu$DMSP, d$^{-1}$) and DMSP production rates (nmol d$^{-1}$) were observed in two experiments (7–90 % decrease), whilst the response under high CO$_2$ from the remaining experiments was generally indistinguishable from ambient controls. Rates of bacterial DMS gross consumption and production gave weak and inconsistent responses to high CO$_2$. The variables and rates we report increase our understanding of the processes behind the response to OA. This could provide the opportunity to improve upon mesocosm-derived empirical modelling relationships and to move towards a mechanistic approach for predicting future DMS concentrations.

1 Introduction

Dimethyl sulfide (DMS) is a ubiquitous marine trace gas derived from the breakdown of the algal osmolyte $\beta$-dimethyl sulfoniopropionate (DMSP). A variety of phytoplankton species produce DMSP, with the majority of production attributed to prymnesiophytes, dinoflagellates and diatoms (Stefels, 2000). Reasons for algal biosynthesis of DMSP are thought to include its role as an overflow product for the regulation of carbon and sulfur metabolism, as an osmolyte or compatible solute, and as defence mechanism against microbes, viruses and zooplankton grazing. Finally, along with its breakdown products (DMS, acrylate, dimethyl sulfoxide (DMSO)), DMSP scavenges harmful hydroxyl radicals and other reactive oxygen species, potentially providing the cell with antioxidant protection (Sunda et al., 2002, 2007).

DMSP is released from phytoplankton cells into the dissolved phase by active exudation or when cells are lysed during grazing, viral attack or senescence (Stefels et al., 2007). Once in this phase, marine bacteria play a vital role in the fate of DMSP in the surface oceans. Indeed, most DMSP released from phytoplankton is either catabolised by bacteria to produce DMS (Todd et al., 2007, 2009), or
Demethylated/demethiolated to produce other key organosulfur compounds such as methanethiol (MeSH) (Kiene et al., 2000; Moran et al., 2012). The demethylation/demethiolation (de/de) pathway regularly dominates as it is considered more energetically advantageous than the cleavage pathway (Kiene et al., 2000; Simó et al., 2002; Moran et al., 2012). Thus, DMSP turnover often exceeds DMS production in the surface oceans. However, the factors regulating the switch between these two competing pathways are still poorly understood (Moran et al., 2012).

DMS is also subject to rapid biological consumption, a process thought to account for 50–80% of total DMS loss in the surface oceans and one which often dominates in bloom situations (Fabric et al., 1999; Simó, 2004). Photochemical loss of DMS competes with this biological loss pathway, comprising a comparable proportion of the total loss (Toole et al., 2006; Vila-Costa et al., 2008). Of the total DMSP produced in the oceans, only ~1% remains to undergo exchange to the atmosphere – however this flux, amounting to ~28 Tg S yr⁻¹, comprises approximately 50% of the total global natural S flux (Andreae, 1990; Länna et al., 2011). In the atmosphere, DMS is rapidly oxidised, forming oxidation products that contribute to the atmospheric aerosol burden, and can lead to the formation of and/or growth of particles (Charlson et al., 1987; Barnes et al., 2006). In addition, DMS-derived aerosols are highly effective at influencing cloud albedo, with implications for global radiative forcing (Rap et al., 2013).

Increasing atmospheric CO₂ levels and the corresponding oceanic uptake of excess CO₂ is resulting in changes to the carbonate chemistry of the surface oceans. This process, termed ocean acidification (OA), manifests itself as decreasing carbonate ion concentrations [CO₃²⁻], increasing hydrogen ion concentrations [H⁺] and a corresponding decrease in seawater pH (Caldeira and Wickett, 2003). Since industrialisation, mean surface ocean pH has fallen by 0.1 pH units. A further decrease of 0.4–0.5 pH units by 2100 is likely unless stringent global CO₂ emissions stabilisation is implemented (Caldeira and Wickett, 2003, 2005). Such rapid changes to seawater chemistry, potentially unprecedented in the last ~300 Ma, are likely to have serious repercussions for marine biological and biogeochemical processes (Raven et al., 2005; Hendriks et al., 2010; Liu et al., 2010; Riebesell and Tortell, 2011; Höӧnisch et al., 2012).

The majority of OA experiments that consider DMS/DMSP have been coastal temperate and subpolar mesocosms, with observations of bulk DMS and DMSP made over 3–4 weeks during a nutrient-induced springtime phytoplankton bloom (Wingenter et al., 2007; Vogt et al., 2008; Hopkins et al., 2010; Kim et al., 2010; Avgoustidi et al., 2012; Archer et al., 2013). Thus, available data are limited by lack of geographical and seasonal coverage. Decreases in DMS have been observed under predicted turn-of-the-century (year 2100) levels of CO₂, ranging from ~35% (Archer et al., 2013) to ~60% (Hopkins et al., 2010; Avgoustidi et al., 2012) relative to ambient controls. When similar levels of pH-driven changes in DMS concentrations are applied globally, the resultant changes in DMS emissions from the oceans may be sufficiently large to influence climate (Six et al., 2013). However, the pattern of decreasing DMS with decreased pH is not consistent amongst all mesocosm experiments. The study by Kim et al. (2010) reported a ~80% increase in DMS under high CO₂. In one case, little difference in concentrations between ambient and high CO₂ was observed (Vogt et al., 2008). The DMS response is similarly variable but of comparable orders of magnitude. Simple relationships have been derived with biological measurements (e.g. plankton community structure, pigments, bacterial abundance and rates) as drivers of the observed DMS and DMSP responses. Physiological effects at the cellular level are more difficult to detect. Broad community-level taxonomic shifts drive the DMS response because the altered conditions favour some species over others (Engel et al., 2008; Meakin and Wyman, 2011; Newbold et al., 2012; Brussaard et al., 2013). Besides the mesocosm experiments, one past study reported the response of natural communities using a shipboard continuous culture system (Lee et al., 2009); here, elevated CO₂ had no effect on concentrations of DMSP. There is still a fundamental lack of mechanistic and physiological understanding of the responses and limited information on regional variability.

Improved predictive capability of models is likely to require a better understanding of the mechanisms driving OA-induced changes in DMS biogeochemistry. Unialgal cultures potentially provide a useful insight into the specific physiological response of a single species to elevated CO₂. DMS decreased two- to tenfold in cultures of the coccolithophore Emiliania huxleyi under high CO₂ (Arnold et al., 2013; Avgoustidi et al., 2012), whereas intracellular DMSP responded differently, with both significant decreases (Avgoustidi et al., 2012) and increases (Arnold et al., 2013). Spilmeyer and Pohnert (2012) reported decreases in cellular DMSP in diatoms, but increases in non-calculating strains of E. huxleyi. How informative these results are with regard to the complex DMSP and DMS cycle in natural communities is questionable.

In an attempt to bridge the gap between the complexity of interpreting processes in traditional mesocosm experiments and the limited applicability of unialgal culture experiments to natural systems, this study was designed to assess the eco-physiological response of a variety of natural microbial communities from a wide geographic area to high CO₂. Detailed discussion of the choice of experimental design in the context of previous OA studies is given in this issue by Richier et al. (2014a). By determining rates of key processes, along with standing stock measurements of DMSP and DMS, we can consider a number of hypotheses. Firstly, increased CO₂ may stimulate primary productivity in phytoplankton communities (Riebesell and Tortell, 2011), resulting in enhanced DMSP synthesis rates and increased DMS concentrations.
(Archer et al., 2013). Secondly, elevated primary production may stimulate bacterial production by increasing the availability of organic substrates (Weinbauer et al., 2011; Engel et al., 2013; Piontek et al., 2013). This would create a greater bacterial demand for DMSP sulfur and stimulate the de/desulfurification pathway, whilst also resulting in an increase in bacterial DMS consumption. The overall effect would be decreased gross DMS production.

2 Materials and methods

2.1 Experimental bioassays

Five shipboard experimental bioassays (hereafter E01–E05) were undertaken during the UK Ocean Acidification Research Programme (UKOA) NW European shelf cruise aboard the RRS Discovery from 6 June to 10 July 2011. Full details of the sampling and experimental setup are given elsewhere in this issue by Richier et al. (2014a), and a general overview is given here; all data are available from the British Oceanographic Data Centre (Richier et al., 2014b). For bioassay water collection, a stainless steel conductivity–temperature–depth (CTD) rosette comprising twenty-four 20 L bottles was deployed, and all bottles were simultaneously fired at the near surface (5–12 m). Water collection commenced pre-dawn, at 02:00 GMT for E01–E04 and at 01:00 GMT for E05 due to its more northerly location and earlier sunrise time (locations and sample depths given in Fig. 1 and Table 1). Once the rosette was back on deck, the water was directly transferred into 4 L polycarbonate bottles, with no screening or filtration. Manipulations of the carbonate chemistry were achieved by additions of NaHCO$_3$ (1 M) and HCl (1 M) to attain the four target CO$_2$ levels (ambient, 550 µatm, 750 µatm, 1000 µatm). After first ensuring the absence of bubbles or headspace, the bottles were sealed with septa lids (high-density polyethylene (HDPE) with silicone and polytetrafluoroethylene (PTFE) septum) and placed in the incubation container. Bottles were incubated inside a custom designed temperature- and light-controlled shipping container, set to match the in situ water temperature at the time of collection. A constant light level (100 µE m$^{-2}$ s$^{-1}$) incorporating an 18–6 h light–dark cycle was provided by daylight simulation LED panels (Powerpak, UK). The daily light dose within the experimental bioassays was considered to be as close to representative as possible of that experienced in situ by the microbial community. Details of the integrated mixed layer irradiances at the time of sample collection for each experiment are given in Richier et al. (2014a). Each bottle belonged to a set of biological triplicates, and sacrificial sampling of bottles was performed at one of two time points (T1 at 48 h and T2 at 96 h), with three sets of biological triplicates for each time point to allow for the sampling requirements of the entire scientific party (3 × 3 bottles, × 2 time points, × 4 CO$_2$ treatments = 72 total). At each time point (48 h and 96 h), the relevant bottles were removed from the incubation container and sampled. Samples for carbonate chemistry measurements were made first to avoid gas exchange with ambient air, followed by sampling for DMS, DMSP and related parameters.

2.2 DMS and DMSP standing stocks

The volume of water required to fill all experimental bioassay bottles and to make key initial measurements (carbonate chemistry, nutrients, Chl $a$, photophysiology) was such that there was insufficient water for DMS-related initial measurements to be taken from the same CTD cast. Therefore, a second CTD cast was carried out after the bioassay cast to collect additional water for further initial measurements, at the same station and depth, 2–3 h after the primary bioassay cast. Carbonate chemistry parameters (dissolved inorganic carbon (DIC), total alkalinity (TA), pH, pCO$_2$; see Richier et al. (2014a) for details of methods) were determined from both 0 h bioassay bottles and the second CTD cast. Single values from bioassay 0 h bottles fell within the range of triplicate values from 0 h CTD casts (Table S1 in the Supplement), suggesting little or no change in water mass between CTD casts. It was from these additional CTD casts that all initial DMS and DMSP samples were taken. Samples were taken directly from the Niskin bottles using Tygon tubing and collected in 250 mL amber glass-stoppered bottles. The bottle was rinsed three times before being filled gently from the bottom and then allowed to overflow three times. Once full, the glass stopper was securely placed on the bottle, ensuring the presence of no headspace. Samples were kept in a cool box and analysed within 2 h.

Samples at experimental time points (48 h, 96 h) were taken directly from the bioassay bottles. After inverting the bioassay bottle three times to ensure resuspension of particulates, samples were siphoned from the bottles directly into 100 mL glass-stoppered bottles using 6 mm silicone tubing. The bottle was first rinsed then filled to the top, ensuring the presence of no bubbles or headspace.

Seawater DMS concentrations were determined by cryogenic purge and trap, followed by detection via gas chromatography with a pulsed flame photometric detector, as outlined in Archer et al. (2013). Total DMSP concentrations (DMSP$_t$) from the same sample bottle were fixed by the addition of 35 µl of 50 % H$_2$SO$_4$ to 7 mL of seawater (Kiene and Slezak, 2006) and analysed within 2 months of collection, again as described in Archer et al. (2013). DMS calibrations were performed using alkaline cold hydrolysis (10 MNaOH) of DMSP (> 98 % purity, Centrum voor Analyse, Spectroscopie and Synthese, Rijksuniversiteit Groningen) diluted 3 times in Milli-Q water to give working standards in the range 0.03–3.3 ng S mL$^{-1}$, and multi-point calibrations were performed every 2–4 days throughout the cruise.

Table 1. Overview of initial conditions for experimental bioassays. MLD stands for mixed layer depth, DMSPt is total DMSP and DMSPp is particulate DMSP. Data shown are means (± standard error) for triplicate measurements made at 0 h for each experiment. Standard errors are not shown for experimental $pCO_2$ and pH as values are based on single measurements.

<table>
<thead>
<tr>
<th></th>
<th>E01</th>
<th>E02</th>
<th>E03</th>
<th>E04</th>
<th>E05</th>
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<td>Stratified</td>
<td>Mixed</td>
<td>Stratified</td>
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<td>Sample depth (m)</td>
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<td>5</td>
<td>10</td>
<td>5</td>
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<td>MLD (total depth) (m)</td>
<td>13 (190)</td>
<td>76 (76)</td>
<td>44 (4760)</td>
<td>33 (33)</td>
<td>14 (72)</td>
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<td>$T$ (°C)</td>
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<td>11.8</td>
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<td>Salinity</td>
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<td>35.0</td>
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<td>$pCO_2$ (µatm)</td>
<td>In situ 342.6 ± 2.4</td>
<td>333.7 ± 3.1</td>
<td>339.8 ± 3.5</td>
<td>400.6 ± 3.3</td>
<td>368.1 ± 2.5</td>
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<td>345.4</td>
<td>395.4</td>
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<td>~ 550</td>
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<td>674.0</td>
<td>691.4</td>
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<td>~ 1000</td>
<td>969.6</td>
<td>862.7</td>
<td>877.8</td>
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<td>pH</td>
<td>In situ</td>
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<td>~ 1000</td>
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<td>Nutrients and Chl a</td>
<td>NO$_3^-$ (µM)</td>
<td>1.1 ± 0.1</td>
<td>0.3 ± 0.01</td>
<td>0.6 ± 0.0</td>
<td>0.9 ± 0.14</td>
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<td>Si(OH)$_4$ (µM)</td>
<td>2.1 ± 0.04</td>
<td>0.5 ± 0.01</td>
<td>0.6 ± 0.01</td>
<td>0.8 ± 0.01</td>
</tr>
<tr>
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<td>PO$_4^{3-}$ (µM)</td>
<td>0.1 ± 0.2</td>
<td>0.1 ± 0.01</td>
<td>0.1 ± 0.001</td>
<td>0.1 ± 0.01</td>
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<td>Total Chl a (µg L$^{-1}$)</td>
<td>3.3 ± 0.04</td>
<td>3.5 ± 0.07</td>
<td>0.8 ± 0.04</td>
<td>1.3 ± 0.03</td>
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<td></td>
<td>&gt; 10µm Chl a (µg L$^{-1}$)</td>
<td>no data</td>
<td>2.8 ± 0.12</td>
<td>0.3 ± 0.02</td>
<td>0.3 ± 0.03</td>
</tr>
<tr>
<td>DMS parameters</td>
<td>DMSPt (nmol L$^{-1}$)</td>
<td>25.9 ± 1.9</td>
<td>59.6 ± 0.7</td>
<td>44.6 ± 1.3</td>
<td>8.0 ± 0.4</td>
</tr>
<tr>
<td></td>
<td>DMS (nmol L$^{-1}$)</td>
<td>1.1 ± 0.02</td>
<td>0.7 ± 0.02</td>
<td>2.1 ± 0.2</td>
<td>1.1 ± 0.02</td>
</tr>
<tr>
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<td>DMSPt : Chl a (nmol µg$^{-1}$)</td>
<td>18.3 ± 0.2</td>
<td>9.2 ± 0.2</td>
<td>58.1 ± 3.8</td>
<td>6.2 ± 0.3</td>
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<td></td>
<td>DMS : DMSPt</td>
<td>0.02 ± 0.001</td>
<td>0.03 ± 0.002</td>
<td>0.05 ± 0.004</td>
<td>0.14 ± 0.004</td>
</tr>
</tbody>
</table>

2.3 DMSP synthesis and production rates

De novo DMSP synthesis and gross production rates were determined for all bioassay experiments at each experimental time point, except for E01. The methods used are based on the approach of Stefels et al. (2009) and are described in detail in Archer et al. (2013). For each CO$_2$ level, triplicate rate measurements were determined. For each rate measurement $3 \times 500$ mL polycarbonate bottles were filled by gently siphoning water directly from each replicate bioassay bottle. Trace amounts of NaH$^{13}$CO$_3$, equivalent to ~6% of in situ dissolved inorganic carbon (DIC), were added to each 500 mL bottle. The bottles were incubated in the bioassay incubation container with temperature and light levels as described above. Samples were taken at 0 h and then at two further time points over a 6–9 h period. At each time point, 250 mL was gravity-filtered in the dark through a 47 mm GF/F filter, the filter was gently folded and placed in a 20 mL serum vial with 10 mL of Milli-Q and one NaOH pellet, and the vial was crimp-sealed. Samples were stored at $-20^\circ$C until analysis by proton transfer reaction mass spectrometer (PTR–MS) (Stefels et al., 2009).
in Milli-Q water. DMSpp concentrations were subsequently measured as DMS following alkaline hydrolysis (see above).

2.4 Biological DMS consumption (BC) and estimation of gross DMS production (GP)

Gross DMS production (GP) by the whole microbial community and bacterial DMS consumption (BC), were determined by dark incubations of whole seawater with $^{13}$C-DMS additions. Incubations were performed at each experimental time point of the five bioassay experiments. Six incubations were performed per experiment (i.e. three time points for both ambient and 750 µam CO$_2$ treatments).

$^{13}$C-DMS solutions were prepared by reducing $^{13}$C-dimethyl sulfoxide (DMSO) (100 µL 99 % DMSO, Sigma Aldrich Co., diluted in 900 µL Milli-Q) using sodium borohydride (NaBH$_4$, 1 g left for 5 min). The reactants were purged (20 mL min$^{-1}$ for 20 min) and $^{13}$C-DMS collected in a 1 / 16” PTFE loop submerged in liquid nitrogen. The $^{13}$C-DMS was rinsed into a 20 mL glass serum vial by syringing 15 mL of Milli-Q through the sample loop. The serum vial was crimp-sealed and stored in the dark at 4°C. The primary solution underwent two serial dilutions in Milli-Q to produce a working solution.

For each incubation, 500 mL of seawater were siphoned from the bioassay bottle into an acid-washed (1 % HCl) and thoroughly rinsed 1 L Tedlar bag. Once filling was complete, any bubbles and headspace were gently expelled from the bag. Each bag was spiked with small volumes (100–150 µL) of a working solution of $^{13}$C-DMS (in Milli-Q water) to give concentrations of 0.1–0.3 nM. After spiking, the bags were left for 1 h to allow complete homogenisation of the tracer. Bags were incubated in the dark, and concentrations of $^{13}$C-DMS and $^{12}$C-DMS were monitored to determine $^{13}$C-DMS loss rates and net and gross DMS. Subsamples (20 mL) were withdrawn using a glass syringe at 0 h and at three further time points over a 10–12 h period. The samples were gently filtered through a Millipore filtration unit containing a 25 mm GF/F filter, directly into a 10 mL glass syringe, and 8 mL of filtered seawater were injected into a glass purge tower. The addition of air and bubbles was avoided or minimised at all times. The sample was purged for 8 min at 90 mL min$^{-1}$, dried with a PTFE counterflow Nafion drier and trapped in a 1 / 16” PTFE loop held in liquid nitrogen. Once purging was complete, the sample loop was rapidly submerged in boiling water, injecting the sample into an Agilent 5973 N gas chromatograph with a 60 m DB-VRX capillary column and mass spectral detector (MSD). The oven temperature was held at 60 °C for 8 min and was increased sharply to 220 °C for the remainder of the 10 min runtime. DMS and $^{13}$C-DMS eluted from the column at $\sim 5.3$ min. The MSD was operated in single ion mode (SIM) and was programmed to detect the following ions: m/z 62, 61 and 47 for $^{13}$C-DMS; m/z 64 for $^{13}$C-DMS; and m/z 68 for deuterated DMS (CD$_3$SCD). of this, 100 µL of a 5 ppmv gas standard was injected into the

The specific growth rate of DMSP ($\mu$DMSP) was calculated assuming exponential growth from

$$
\mu_t = \alpha_k \times AVG \left[ \ln \left( \frac{64\text{MP}_{eq} - 64\text{MP}_{t-1}}{64\text{MP}_{eq} - 64\text{MP}_{t}} \right) \right.
$$

$$
\left. \left( \ln \left( \frac{64\text{MP}_{eq} - 64\text{MP}_{t-1}}{64\text{MP}_{eq} - 64\text{MP}_{t+1}} \right) \right) \right]
$$

(Stefels et al., 2009), where $64\text{MP}_{t-1}$ and $64\text{MP}_{t+1}$ are the proportion of $1 \times 13$C labelled DMSP relative to total DMSP at time $t$, at the preceding time point ($t - 1$) and at the subsequent time point ($t + 1$), respectively. Values of $64\text{MP}$ were calculated from the protonated masses of DMS as follows: mass $64/$(mass$63+$mass$64+$mass$65$), determined by PTR–MS. $64\text{MP}_{eq}$ is the theoretical equilibrium proportion of $1 \times 13$C based on a binomial distribution and the proportion of tracer addition. An example of the change in proportion of mass ratio $64(64\text{MP})$ during incubations of water from experiment E05 at T48 for the different $p$CO$_2$ treatments is shown in Fig. S1 in the Supplement. An isotope fractionation factor $\alpha_k$ of 1.06 is included, based on laboratory culture experiments using Emiliania huxleyi (Stefels et al., 2009).

Gross DMS production rates during the incubations (nmol L$^{-1}$ h$^{-1}$) were calculated from $\mu$DMSP and the initial particulate DMSP (DMSpp) concentration of the incubations. Concentrations of DMSpp were determined at each time point by gravity filtering 7 mL of sample onto a 25 mm GF/F filter and preserving the filter in 7 mL of 35 mM H$_2$SO$_4$.
purge gas stream for every sample in order to monitor and correct for system sensitivity and drift. By taking the ratio of \( m/\nu \) 62 or 64 to \( m/\nu \) 68, a greatly improved precision of analysis was attained.

Calibrations were performed using pure DMS (\( >99\%\), Sigma Aldrich Co.) diluted in Milli-Q water. A primary standard was gravimetrically prepared, and then serial dilutions were performed to obtain a working standard of 0.3 ng S mL\(^{-1}\). Multi-point calibrations, from 0 to 3.5 ng S mL\(^{-1}\), were performed every 3 days for the duration of the cruise. Detection limits for \(^{12}\text{C}-\text{DMS}\) and \(^{13}\text{C}-\text{DMS}\) were 0.1 ng S and 0.03 ng S, respectively. The lower detection limit for \(^{13}\text{C}-\text{DMS}\) is the result of lower background concentrations of this isotope in blank measurements. Average analytical precision based on triplicate samples was \( \sim 10\%\).

Rates of BC (nmol L\(^{-1}\) d\(^{-1}\)) were estimated from the slope of the linear decrease in \(^{13}\text{C}-\text{DMS}\) concentrations over the 10–12 h incubation period (see Fig. S2 in the Supplement). DMS loss via photochemistry and gas exchange was excluded, so the \(^{13}\text{C}-\text{DMS}\) loss is considered to be equivalent to biological consumption. To scale to in situ DMS concentrations, BC rates were first divided by the concentration of \(^{13}\text{C}-\text{DMS}\) tracer (giving first-order rate constants, d\(^{-1}\)) and multiplied by in situ DMS concentrations. The \(^{12}\text{C}-\text{DMS}\) change during the incubations represents the net DMS production rate. The GP was estimated as the sum of net DMS production and BC.

Additions of \(^{13}\text{C}-\text{DMS}\) ranged from 10 to 100% of ambient DMS concentrations. Large relative additions of \(^{13}\text{C}-\text{DMS}\) were unavoidable when ambient DMS concentrations were low (<1 nmol L\(^{-1}\)) due to the limits of detection of the method. The following equation was used to correct BC rates for any enhancement caused by tracer addition (rdMS\(_{t}\)), (Rees et al., 1999), which was originally described for the correction of \(^{15}\text{N}\) ammonium uptake rates:

\[
\text{rdMS}_{t} = \frac{\text{rdMS}_{a}}{\text{DMS}_{\text{tot}}/\left(K_s + \text{DMS}_{\text{tot}}\right) - \left(K_s + \text{DMS}_{a}\right)/\text{DMS}_{a}},
\]

where rdMS is the loss rate adjusted for stimulation by tracer (nmol L\(^{-1}\) d\(^{-1}\)), rdMS\(_{a}\) is the original loss rate (nmol L\(^{-1}\) d\(^{-1}\)), DMS\(_{\text{tot}}\) is [DMS] ambient + [\(^{13}\text{C}-\text{DMS}\)] (nmol L\(^{-1}\)), DMS\(_{a}\) is [DMS] ambient (nmol L\(^{-1}\)), and \( K_s \) is the half-saturation parameter (nmol L\(^{-1}\)). The value of \( K_s \) was taken as the mean of the \( K_s \) determined through three kinetic experiments described in the section below. The applied corrections resulted in decreases in the biological DMS consumption rates of 8–51%; the larger uncertainties are associated with relatively low in situ DMS concentrations, thus resulting in relatively high tracer additions. Applying the maximum (25.0 nmol L\(^{-1}\)) and minimum (4.5 nmol L\(^{-1}\)) \( K_s \) values to the correction gives the following uncertainties on the loss rates: E01 13.8–24.9 %, E02 23.1–39.7 %, E03 7.0–9.1 %, E04 9.7–12.7 % and E05 1.1–7.7 %. Using the standard error of mean \( K_s \) to the correction (shown in Table 2) results in uncertainty on loss rates of E01 3.8–6.9 %, E02 6.3–9.5 %, E03 1.6–2.0 %, E04 1.9–2.3 % and E05 0.3–1.6 %.

### 2.5 Biological DMS consumption: kinetic parameters

Three additional experiments, hereafter referred to as KE1, KE2 and KE3, were conducted to determine BC kinetic parameters. As it was not logistically feasible to conduct the kinetic experiments in parallel to the bioassay incubations, the three sites were chosen to encompass contrasting NW European shelf waters. The locations of sampling for kinetic experiments are shown in Fig. 1. Unfiltered surface seawater (3–5 m) was siphoned into five 1 L Tedlar bags that had been acid-washed with 1 % HCl and thoroughly rinsed three times with ultrapure water. Once filling was complete, any bubbles and headspace were gently expelled from the bag. For each experiment, increasing concentrations of \(^{13}\text{C}-\text{DMS}\) were added to each bag, ranging from at or below in situ concentrations up to 74 nmol L\(^{-1}\). After the addition, the bags were left for 1 h to allow complete homogenisation of the tracer. The bags were incubated for up to 12 h in the dark at in situ seawater temperature, and samples were processed and analysed as described above.

Initial DMS concentrations showed a wide range, from 1.0 nmol L\(^{-1}\) in KE1 to 16.8 nmol L\(^{-1}\) in KE3 (see Table 2 and water column profiles in Fig. S3 in the Supplement). A large range of DMS : DMSP\(_{t}\) ratios were also encountered, with a value of 0.2 for KE3, which was an order of magnitude higher than the values of 0.02 and 0.03 for KE1 and KE2.

A summary of the Michaelis–Menten kinetic parameters for the three experiments, obtained through non-linear regression using Minitab 16.0 statistical software, are given in Table 2. Kinetic curves (see example Fig. S4 in the Supplement) were based on five to six data points; replicate samples were unfeasible due to the time taken to analyse all samples at each of four time points over a \( \sim 10\) h period. However, the non-linear regressions for each experiment were significant (\( P < 0.05\)). The kinetic parameters generated in this study represent the activity of natural assemblages rather than the activity of single enzymes or species and showed a similarly broad range to in situ DMS(P) characteristics. Half-saturation constants (\( K_s \)) ranged from 4.5 nmol L\(^{-1}\) in KE3 to 25.0 L\(^{-1}\) in KE1, with KE2 displaying an intermediate \( K_s \) value of 11.4 nmol L\(^{-1}\). Thus, an inverse relationship between in situ DMS concentrations and \( K_s \) was apparent. Maximum DMS consumption rates (\( V_{\text{max}} \)) were also variable. KE3 displayed the lowest \( V_{\text{max}} \) of only 1.3 nmol L\(^{-1}\) d\(^{-1}\), and this time KE2 gave the highest values with 25.9 nmol L\(^{-1}\) d\(^{-1}\). Despite having the highest \( K_s \) and lowest in situ DMS concentrations, KE1 yielded an intermediate \( V_{\text{max}} \) of 10.9 nmol L\(^{-1}\) d\(^{-1}\).
2.6 Total bacteria abundance

Samples for the determination of total bacteria abundance were taken in triplicate at the start of all incubations for BC and GP. Twenty µL of glutaraldehyde solution (Grade 1, 50% in H2O, Sigma Aldrich Co.) was added to 2 mL samples, and the vials were fixed for 30 min at 4°C, followed by snap freezing in liquid nitrogen and storage at −80°C until analysis. Bacteria were counted by flow cytometry according to Marie et al. (1999). Briefly, thawed samples were diluted with Tris-EDTA buffer (10 mM Tris–HCl and 1 mM EDTA, pH 8) and stained with the green fluorescent nucleic acid–specific dye SYBR-Green I (Molecular Probes, Invitrogen Inc.) at a final concentration of 1 × 10⁻⁴ of the commercial stock in the dark at room temperature for 15 min. Bacteria were discriminated in bivariate scatter plots of green fluorescence versus side scatter.

2.7 Ancillary parameters

A description of methodologies for all ancillary parameters described in Table 1 (carbonate chemistry, nutrients, total and size-fractionated chlorophyll a) and for the enumeration of small phytoplankton by flow cytometry (Fig. 2) is given in Richier et al. (2014a).

3 Results

3.1 Experimental bioassays: starting conditions

The design of this study allowed an assessment of the response of surface ocean communities to high CO2 during the most biologically productive time of year by sampling a variety of seasonally stratified (E01, E03, E05) and perennially mixed (E02, E04) sites, both on- and off-shelf (Fig. 1). Initial conditions generally displayed typical NW European shelf summertime characteristics, with low concentrations of nutrients (< 1.1 µM total organic nitrogen (TON), ~ 0.1 µM P) and a range of Chl a concentrations (0.8–3.3 µg L⁻¹) reflecting the heterogeneous spatial distribution of marine productivity in shelf sea waters. In situ pCO2 concentrations were similarly variable, ranging from 334 µatm for E02 to 401 µatm for E04 (See Table 1). Further detail is given elsewhere (Richier et al., 2014a).

3.2 DMSPt and DMS: concentrations and ratios

Triplicate bottles for each treatment displayed similar values and trends of DMSPt over the course of each experiment. Initial concentrations in the five bioassays spanned a wide range, from 8.0 nmol L⁻¹ for E04 to 59.6 nmol L⁻¹ for E01 (Table 1). There was a net increase in DMSPt for all ambient control incubations after 96 h, with the greatest increase seen over the first 48 h. For all experiments, a clear response to increasing levels of CO2 was observed (Fig. 2a–e), with a large number of significant reductions in DMSPt with increasing CO2 concentrations after 48 h (analysis of variance (ANOVA), Holm–Sidak, P < 0.05; Table S2 in the Supplement). After 96 h, concentrations showed some recovery in high-CO2 treatments for E02 and E04, but the response persisted for E01, E03 and E05. In addition, when all data were included in a single analysis, a significant negative relationship between DMSPt and [H⁺] data was seen (ANOVA, significance of F-ratio P < 0.001; Table S3 in the Supplement).

Again, concentrations and trends of DMS from triplicate bottles within each treatment were similar for all experiments. Initial mean concentrations of DMS (Fig. 2f–j) varied by a factor of 3, from 0.7 nmol L⁻¹ (E02) to 2.1 nmol L⁻¹ (E03), and higher DMS concentrations tended to be associated with lower Chl a (Table 1). Net DMS accumulation was observed in all ambient control incubations over 96 h. There was a clear response to high CO2 in all experiments, with significantly elevated DMS concentrations, predominantly after 48 h (ANOVA, Holm–Sidak, P < 0.05; Table S2 in the Supplement). The response became more variable by 96 h, with rapid rises in DMS in ambient controls for E01, E03 and E04 between 48 and 96 h, resulting in concentrations similar to those under high CO2. For all experiments at each time point (with the exception of E04 at 96 h) significant positive relationships between DMS concentrations and [H⁺] were observed (Table S3 in the Supplement). The relationship between these variables was strongest at 48 h (r² = 0.58 – 0.92, Table 2).

Table 2. Kinetic parameters (Ks (half-saturation constant), Vmax (maximum ¹³C-DMS consumption rate)) and turnover time of DMS due to BC (τBC) for ¹³C-DMS loss rates at three contrasting sites in NW European shelf waters. Kinetic parameters were calculated by fitting data to the Michaelis–Menten equation through non-linear regression of loss rate vs. ¹³C-DMS concentration data from three kinetic experiments (See Fig. S4 in the Supplement). Initial in situ DMS concentrations measured at 0 h of each kinetic experiment are also shown.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Date</th>
<th>Initial [DMS] (nmol L⁻¹)</th>
<th>DMS : DMSPt (nmol L⁻¹)</th>
<th>Ks ± SE (nmol L⁻¹ d⁻¹)</th>
<th>Vmax ± SE (nmol L⁻¹ d⁻¹)</th>
<th>r² (d)</th>
<th>τBC (d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KE1</td>
<td>19 June 2011</td>
<td>1.0</td>
<td>0.02</td>
<td>25.0 ± 13.8</td>
<td>10.9 ± 2.5</td>
<td>0.91</td>
<td>0.09</td>
</tr>
<tr>
<td>KE2</td>
<td>24 June 2011</td>
<td>3.8</td>
<td>0.03</td>
<td>11.4 ± 6.7</td>
<td>25.9 ± 7.3</td>
<td>0.96</td>
<td>0.15</td>
</tr>
<tr>
<td>KE3</td>
<td>5 July 2011</td>
<td>16.8</td>
<td>0.2</td>
<td>4.5 ± 1.1</td>
<td>1.3 ± 0.1</td>
<td>0.89</td>
<td>12.9</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td>13.6 ± 7.2</td>
<td>12.7 ± 3.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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$P < 0.01$ and generally weakened at 96 h ($r^2 = 0.35 - 0.79$, $P < 0.05$).

In summary, large increases in mean DMS concentrations relative to ambient controls were observed in all bioassays: 110% (28–223%) at 550 µatm, 153% (56–295%) at 750 µatm and 225% (79–413%) at 1000 µatm. By contrast, mean DMSPt concentrations showed a consistent decrease, but to a lesser extent than DMS: 28% (18–40%) at 550 µatm, 44% (18–64%) at 750 µatm and 52% (24–72%) at 1000 µatm.

Initial DMS: DMSPt ratios ranged from 0.02 for E01 to 0.14 for E04 (Fig. S5 in the Supplement). This ratio gradually increased over 96 h (except in E04) and for all experiments was significantly correlated with increasing [H+] after both 48 h and 96 h (Fig. 3 and Table S3 in the Supplement; ANOVA of regression, $F$ ratio $P < 0.05$). DMS: Chl $a$ ratios generally followed similar trends to DMS: DMSPt (Fig. S5 in the Supplement). Significant positive relationships between DMS: Chl $a$ and [H+] were identified for all experiments at 48 h (ANOVA of regression, $F$ ratio $P < 0.001$, except E04 $P < 0.05$; Table S3 in the Supplement). By 96 h, this relationship remained significant for all experiments but was negative for E01 and E04.

DMSPt: Chl $a$ also showed large variation between experiments, from $< 30$ nmol µg$^{-1}$ in E02 up to $∼ 150$ nmol µg$^{-1}$ in E03 (Fig. S5 in the Supplement). In general, increasing [H+] was associated with a decrease in DMSPt: Chl $a$, but this relationship was found to be significant only for E01 and E04 at 48 h. By 96 h, the majority of experiments displayed a significant negative relationship between these variables (ANOVA of regression, $F$ ratio $P < 0.01$; Table S3 in the Supplement).

### 3.3 Plankton community and OA response

In general, an increase in the abundance of small phytoplankton ($< 10 \mu$m, pico- and nanophytoplankton) was seen over the experimental period, indicating growth within the bioassay bottles (Fig. 2k–o). For E02–E05, abundances were significantly lower under high CO$_2$ at both 48 and 96 h (ANOVA, Holm–Sidak, $P < 0.05$; Table S4 in the Supplement). By contrast, there were significant increases in abundance under high CO$_2$ (750 and 1000 µatm) at 96 h for E01 (ANOVA, Holm–Sidak, $P < 0.001$; Table S4 in the Supplement). The reader is directed to Richier et al. (2014a) for further description of the response of small phytoplankton.

### 3.4 DMSP synthesis and production

Initial DMSP synthesis rate constants ($\mu$DMSP) ranged from 0.33 d$^{-1}$ in E05 to 0.96 d$^{-1}$ in E03. Rate constants tended to decrease over the course of 96 h to $\sim 0.10$–0.50 d$^{-1}$ (Fig. 4a–d). The effect of CO$_2$ treatment on $\mu$DMSP was variable. Marked and significant decreases in $\mu$DMSP at all high-CO$_2$ treatments relative to ambient controls were seen for E02 at 48 h and 96 h (analysis of covariance (ANCOVA), $P < 0.01$; Table S5 in the Supplement) and for E05 at 48 h (ANCOVA, $P < 0.01$; Table S5 in the Supplement). For E03 and E04, the response of $\mu$DMSP to high-CO$_2$ treatments was more variable, with a number of significant differences from ambient CO$_2$ identified (see Table S5 in the Supplement for results of ANCOVA), but of an inconsistent direction relative to ambient CO$_2$.

Temporal trends in DMSP production rates corresponded closely with those for $\mu$DMSP (Fig. 4e–h). Initial DMSP production rates ranged from 7.1 nmol L$^{-1}$ d$^{-1}$ in E05 to 37.3 nmol L$^{-1}$ d$^{-1}$ in E03. For E02 and E05, clear decreases in DMSP production rates were observed in the high-CO$_2$ treatments. In E02, this difference was maintained for the duration of the experiment with a mean of 12.0 nmol L$^{-1}$ d$^{-1}$ at ambient CO$_2$, compared to 1.8–3.1 nmol L$^{-1}$ d$^{-1}$ at elevated CO$_2$. Similarly large decreases in mean DMSP production were seen in E05 (2.9–11.1 nmol L$^{-1}$ d$^{-1}$ high CO$_2$, 13.4 nmol L$^{-1}$ d$^{-1}$ ambient CO$_2$). Similarly to $\mu$DMSP, the response to high CO$_2$ was more variable for E03 and E04.

### 3.5 Bacterially mediated DMS processes

#### 3.5.1 Total bacteria abundance

Initial abundances of bacteria (Fig. 5a–e) in the sub-incubations for BC and GP rates were statistically similar (Kruskal–Wallis $H = 3.273$, $df = 3$, $P = 0.415$), ranging from 0.83 ($± 0.01$) to 1.00 ($± 0.16$) $\times 10^6$ cells mL$^{-1}$ (no data available for E01 0 h). For E01, E02, E04 and E05 bacterial abundance increased with increasing CO$_2$. Small differences in abundance at 48 h were followed by large increases in bacteria at 750 µatm by 96 h. By contrast, a decrease in bacterial abundance in response to increased CO$_2$ was observed for E03.

#### 3.5.2 Biological DMS consumption (BC)

Rates of biological DMS consumption (BC) across the five bioassay experiments ranged from 0.2 ± 0.1 to 8.6 ± 1.6 nmol L$^{-1}$ d$^{-1}$, with the highest overall mean values observed during E01 at Mingulay Reef (3.6 ± 0.7 nmol L$^{-1}$ d$^{-1}$) and the lowest during E02 in the Irish Sea (0.5 nmol L$^{-1}$ d$^{-1}$) (see Fig. 5f–j and Table S6 in the Supplement). All rates fell well within the range of a variety of previous studies (0.02–8.8 nmol L$^{-1}$ d$^{-1}$; Bailey et al., 2008, Kiene et al., 2007, Toole and Siegel 2004, Toole et al., 2006, del Valle et al., 2009, Vila-Costa et al., 2008, Zubkov et al., 2002, 2004). Overall, no consistent response of BC to CO$_2$ treatment was observed. For E01 and E02, BC was greater at high CO$_2$, with significant differences after 48 h (ANOVA $F$ ratio $P < 0.05$). For E03, BC was lower under high CO$_2$, although the differences were not significant. For E04 and E05, significant decreases in BC after 96 h were observed (ANOVA $F$ ratio $P < 0.05$).
3.5.3 Biological DMS turnover

DMS turnover time due to biological consumption ($\tau_{BC}$), calculated as the inverse of the loss rate constant ($k_{BC}$, d$^{-1}$) (Fig. 5k–o and Table S6 in the Supplement), ranged from < 1 d (E01, E05) to a maximum of 12.3 d (E02). Similarly to BC, there was a lack of consistent response to high CO$_2$. For experiments E03–E05, turnover over times were consistently higher at high CO$_2$, but the differences were only significant for E03 (48 h) and E04 (96 h).

3.5.4 Gross DMS production (GP)

Mean gross production rates of DMS (GP) in the five bioassay experiments ranged from undetectable levels during E02 up to a maximum of $8.7 \pm 1.4$ nmol L$^{-1}$ d$^{-1}$ during E05 (Fig. 5p–t, Table S6 in the Supplement). The negative value observed at 48 h in the ambient treatment for E02 was not significantly different from 0, so GP was considered to be undetectable in this case. Overall, a response of GP to high CO$_2$ was variable or undetectable. GP was significantly elevated at high CO$_2$ for E02 at 96 h and E03 after 48 h, whilst a significant decrease in GP was observed for E04 at 96 h (ANOVA $F$ ratio $P<0.05$). No significant differences were seen for E01, E04 and E05.

4 Discussion

Despite variability in physical, biological and biogeochemical characteristics of sampling stations, experimental bioassays gave highly consistent DMS(P) responses to OA, affirming the strength of the experimental approach adopted here. Large increases in mean DMS concentrations relative to ambient CO$_2$ controls were observed in all bioassays: 110% (28–223%) at 550 µatm, 153% (56–295%) at 750 µatm and 225% (79–413%) at 1000 µatm. By contrast, mean DMSPt concentrations showed a consistent decrease: 28% (18–40%) at 550 µatm, 44% (18–64%) at 750 µatm and 52% (24–72%) at 1000 µatm. Our results are in opposition to the majority of results from mesocosm studies. We examine the possible drivers of the observed responses in the following section.

4.1 Influence of plankton community response on DMS(P)

Primarily as a result of the characteristic community dynamics of mesocosm experiments, our current knowledge of the algal physiological effects of OA on the production of DMSP and DMS is minimal. During a mesocosm experiment in Arctic waters, increasing CO$_2$ resulted in elevated gross primary productivity, partly accounted for by an increase in net production of the autotrophic dinoflagellate *Heterocapsa rotundata* (Archer et al., 2013). The net increase in *H. rotundata*
was evident as an increase in DMSP production rates and concentrations at high CO$_2$. Despite observations of a stimulating effect of CO$_2$ on photosynthesis and carbon fixation in a range of other phytoplankton taxa and natural assemblages (Riebesell and Tortell, 2011), there was no evidence for enhanced productivity during the bioassay experiments. Indeed, primary production appeared to decrease with increasing [H$^+$]. $^{14}$C-based measurements of primary production for E02 and E03 showed a tendency to decrease under high CO$_2$ (A. Poulton, personal communication, 2013), whilst in E03, E04 and E05 significantly lower [Chl a] with increasing CO$_2$ was also observed (see Richier et al., 2014a).

Over the shorter timescale of the present study, changes in phytoplankton composition were not expected to influence the DMS(P) response to the same extent as seen in mesocosm experiments. However, it is apparent in the abundance of small phytoplankton (Fig. 2k–o), and is elaborated on by Richier et al. (2014a), that phytoplankton composition altered rapidly between treatments in the majority of the experiments. In all experiments, except E01, the abundance of small phytoplankton (<10 µm) was significantly lower under high CO$_2$ (Fig. 2k–o, Table S4). Furthermore, specific rates of DMSP synthesis (µDMSP) and DMSP production rates were either insensitive to high CO$_2$ (E03, E04) or showed marked declines (E02, E05) (Fig. 4a–d). As small phytoplankton can contribute between 40% and 57% to the total DMSP pool (Archer et al., 2011), it is likely that decreases in DMSP$_t$ were primarily driven by the observed decreases in abundance of this group, accompanied in some cases (E02, E05) by the influence of a decrease in DMS synthesis.

This general decrease in growth of small phytoplankton also coincided with significant increases in DMS : DMSP$_t$, suggesting a relationship between these two responses. OA results in a reduction in buffer capacity that may affect proton concentration ([H$^+$]) and/or regulation at the cell membrane surface of phytoplankton. Larger cells (>25 µm) possess thicker boundary layers and under present-day conditions may experience such changes over the course of a diel cycle (Flynn et al., 2012). As such, they may be better adapted than smaller cells (<25 µm) to the higher [H$^+$] that may be encountered under future OA. During the experimental bioassays, rapid changes to seawater chemistry may have induced changes to external cell surface [H$^+$] beyond those experienced by small cells in the present-day oceans, resulting in deleterious effects on the growth of this fraction of the population. This may explain the reduced growth of small phytoplankton (<10 µm) under high CO$_2$ in most experiments, summarised here in Fig. 2 (k–o and Table S4 in the Supplement) and discussed further by Richier et al. (2014a).

Figure 3. Relationship between DMS : DMSP$_t$ and H$^+$ concentration, ([H$^+$] x 10$^{-8}$ equivalents L$^{-1}$) at 48 h (a) and 96 h (b) for all data from all five bioassay experiments.

This response may be compared to those observed when phytoplankton communities are subjected to other stressors. Gali et al. (2013) proposed that exposure to sublethal or lethal levels of ultraviolet radiation (UVR) could induce an increase in cell membrane permeability, eventually triggering apoptosis. This would lead to the release of intracellular DMSP, increasing its availability for catabolism by bacteria and/or extracellular DLA (DMSP lyase activity). In the bioassays, DMSP concentrations fell and DMS production was stimulated, suggesting that the induced changes to carbonate chemistry may have resulted in an increase in cell permeability and lysis (Flynn et al., 2012; Richier et al., 2014a), leading to increased DMS release from cells. As discussed above, small phytoplankton can contribute a large proportion to the total DMSP pool, and so possess a large potential for DMS production (Archer et al., 2011). Thus, sublethal/lethal cellular damage to this fraction of the community induced by rising [H$^+$] may have resulted in the strong increase in DMS concentrations.

The same response was seen in E01 despite an increase in abundance of small phytoplankton cells with increasing CO$_2$ (Fig. 2k). Uniquely, E01 at Mingulay Reef was dominated by cryptophytes (∼10 µm), whereas all others comprised a mainly <10 µm-sized community. Despite a similar overall response in DMS standing stocks, differences in initial community structure could result in a different physiological response to increased CO$_2$. Rather than cells releasing DMSP/DMS as described above, a physiologically mediated overflow mechanism may have played a role. The cellular release of DMS is thought to facilitate the removal of surplus
sulfur, carbon and energy, allowing the cell to continue functioning during periods of cellular stress or unbalanced growth (Stefels, 2000). This process depends on a direct up-regulation of the conversion of DMSP to DMS via DLA. Previous studies have demonstrated an increase of up to an order of magnitude in DMS production in phytoplankton exposed to UV stress (Sunda et al., 2002; Archer et al., 2010; Gali et al., 2013). The observed increases of DMS : DMSPt with increasing CO$_2$ suggest that elevated [H$^+$] may drive an analogous response, stimulating increases in turnover of DMSP to DMS. This could be driven by the apparent susceptibility of smaller phytoplankton cells to changes in [H$^+$] (Flynn et al., 2012; Richier et al., 2014a). Such a process could, to some extent, be relevant to the results of all bioassay experiments.

**Figure 4.** De novo synthesis of DMSP ($\mu$DMSP, d$^{-1}$) (a–d) and DMSP production rates (nmol L$^{-1}$ d$^{-1}$) (e–h) for E02–E05 (data not available for E01). Asterisks (*) denote significant differences from ambient CO$_2$ bioassays (significance of $F$ ratio from ANCOVA, $p < 0.05$). Smoothed lines on plots do not represent extrapolation of data between time point measurements but are used to highlight trends.

**Figure 5.** DMS consumption rates (nmol L$^{-1}$ d$^{-1}$) (a–e), biological DMS turnover (d) (f–i), DMS gross production rates (k–o) and total bacteria counts (cells mL$^{-1}$) (p–t) during five bioassay experiments at ambient pCO$_2$ and $\sim$750 µatm. DMS turnover times (d) = 1 / loss rate constant ($k_{BC}$, d$^{-1}$). Data are summarised in Table S6 in the Supplement. Error bars indicate standard error on triplicate rate measurements. Asterisks (*) denote significant difference from ambient CO$_2$ bioassays (significance of $F$ ratio from ANOVA, $p < 0.05$). No measurements were made at 0 h for E01. Smoothed lines on plots do not represent extrapolation of data between time point measurements but are used to highlight trends.
4.2 Influence of bacterial community response on DMS(P)

Previous OA studies suggest that increased primary production and photosynthesis at high CO₂ may stimulate bacterial production by increasing the availability of organic substrates for bacterial utilisation (Weinbauer et al., 2011; Engel et al., 2013; Piontek et al., 2013). Such conditions could generate a greater bacterial demand for DMSP sulfur, leading to an increase in bacterial DMSP catabolism via the de/de pathway and to reductions in gross DMS production (Kiene et al., 2000). This mechanism could explain some of the reduction in DMS concentrations under high CO₂ seen during mesocosm studies (Hopkins et al., 2010; Avgoustidi et al., 2012; Archer et al., 2013). These previous findings suggest that increased CO₂ may stimulate bacterial production and (i) increase bacterial DMSP demethylation/demethiolation (de/de) and/or (ii) increase bacterial DMS consumption resulting in reduced gross DMS production.

However, there were few instances of significantly elevated dark DMS gross production rates in response to high CO₂ (Fig. 5 and Table S6 in the Supplement). For some experiments (E02, E03 and E04), trends in GP rates were comparable to temporal trends in net concentrations from the bioassays suggesting microbial DMS production is an important contributor to total DMS production, particularly in the latter stages of the bioassays (>48 h). However, it seems unlikely that the rapid increases in DMS yield seen in the first 48 h can be fully explained by an increase in bacterial DMS production or a reduction in consumption rates. The observed increase in DMS in the bioassays could imply a CO₂ effect resulting in a decrease in the activity of the de/de pathway, making more DMSP available for bacterial cleavage to DMS.

We also examined the response of biological consumption (BC) of DMS to elevated CO₂, a process thought to account for 50–80 % of total DMS loss in the surface oceans (Gabric et al., 1999; Simó, 2004). Increased bacterial production at high CO₂ may lead to increased bacterial consumption of DMS, resulting in a decrease in net DMS production. Rates of BC during this study ranged from 0.2 to 8.6 nmol L⁻¹ d⁻¹ falling well within the range of previous studies (0.02–8.8 nmol L⁻¹ d⁻¹; Bailey et al., 2008; Kiene et al., 2007; Toole et al., 2004, 2006; del Valle et al., 2009; Vila-Costa et al., 2008; Zubkov et al., 2002, 2004). Similarly, turnover times ranged from 0.2 to 12.3 d, again similar to earlier studies (0.7–18.7 d). The wide range in rates seen here reflects the highly heterogeneous spatial variability of both biomass and surface seawater DMS in NW European shelf waters during summertime.

In addition, the results from three kinetic experiments revealed a large range in values of Kₛ and Vₘₐₓ in the study waters implying contrasting levels of control of BC on surface ocean DMS concentrations in the study region (Table S6 in the Supplement). A broad range in these parameters is unsurprising given that the measured rates represent the activity of natural assemblages that will vary greatly in space and time in the dynamic shelf sea environment, rather than the activity of specific single enzymes or species. The three sites (KE1, KE2 and KE3) also encompassed a wide range of surface DMS concentrations of 1.0, 3.8 and 16.8 nmol L⁻¹, respectively, and this was likely a reflection of the contrasting BC characteristics of the sites (Fig. S3 in the Supplement). Vₘₐₓ ranged from 1.3 to 25.9 nmol L⁻¹ d⁻¹. Kₛ was inversely proportional to in situ DMS, demonstrating the saturation of BC at higher concentrations. For KE3, BC appeared to be close to saturation, suggesting a slow response of DMS consumers to rapidly increasing DMS concentrations at this site, reflected in low Kₛ of 4.5 nmol L⁻¹ and a long turnover time of ∼13 d (Table S6 in the Supplement). Such large accumulations of DMS concentrations occur during the breakdown of phytoplankton blooms dominated by DMSP producers and have previously been attributed to low saturation kinetics of DMS consumers (Wolfe et al., 1999; Simó et al., 2000). Where DMS concentrations were lower for KE1 and KE2, Kₛ attained greater values of 11–25 nmol L⁻¹, and rapid turnover times of ∼0.1 d were observed. Clearly, at these sites DMS consumers were able to exert a strong control over the accumulation of seawater DMS.

It is important to reiterate that it was not feasible to perform the kinetic experiments in parallel to the bioassay incubations for rates of BC. Therefore, the three sites chosen for kinetic experiments are assumed to give a good representation of the consumption kinetics likely to be encountered around NW European seas and, of course, within the bioassay experiments, with the recognised caveat that they do not precisely represent the in situ kinetics for each bioassay experiment. The uncertainties associated with the use of mean Kₛ determined from the three kinetic experiments and used to correct BC rates for tracer additions are given in the methodology section. However, given the broad range of consumption kinetics observed and the large range in situ DMS concentrations for each kinetic experiment (Table 2), it seems feasible that our kinetic experiments give a reasonable representation of the consumption kinetics within the bioassay incubations.

In the context of the bioassays, DMS concentrations generally did not exceed 9 nmolL⁻¹, apart from in two cases where concentrations of >10 nmol L⁻¹ were measured at 1000 µatm in E01 and E03 (Fig. 2f and h). With Kₛ in the range of 4.5–25.0 nmol L⁻¹, it was unlikely that this process was saturated during the bioassays, allowing consumption to continue unabated despite rapid rises in DMS concentrations. In the two cases where DMS did exceed 10 nmol L⁻¹, it is possible that consumption kinetics reached saturation. However this was likely a result of, rather than cause of, the DMS CO₂ response. With this in mind, were the observed increases in DMS concentrations driven by CO₂-driven reductions in BC? Although variations in BC between ambient and 750 µatm were observed (Fig. 5f–j, Table S6 in the Supplement), the observed differences were not consistent.
enough to explain the changes in bulk DMS concentrations. Thus bacterial consumption of DMS in the bioassays did not appear to be sensitive to increasing CO$_2$ concentrations.

4.3 Exploring the regional variability

Despite this study targeting sites with a range of physical, biological and biogeochemical characteristics, the overall response to high CO$_2$ was remarkably consistent, suggesting limited regional variability. In Fig. 6, DMSP$_t$ and DMS concentrations under high CO$_2$ are normalised to those in ambient controls to allow comparison of the relative changes seen for each experiment. The general response is clear; however analysis of variance reveals some significant differences in the extent of the response at each station, suggesting varying levels of resilience in the communities.

The relative decreases in DMSP$_t$ concentrations were broadly consistent, ranging from 10 to 30 % relative to controls – except for E05 with a mean decrease of $\sim$50% (Fig. 6a). However, the relative increases in DMS were more variable between stations, from only $\sim$ 10 % for E04 up to $\sim$ 190 % for E01 (though accompanied by the greatest range) (Fig. 6b). Such variability is not surprising given the complex nature of DMSP and DMS seawater dynamics. Furthermore, the extent of the response to high CO$_2$ may be a reflection of the potential community DMS yield. Is it possible to attribute this variability to differences in the characteristics of each station given the available information?

Differences in phytoplankton abundance, speciation and size structure are likely to exert an influence on DMS yields. Initial Chl $a$ concentrations were up to 12 times higher at E01 and E02 compared to E03–E05 (see Table 1), accompanied by clear differences in community size structure. E03–E05 were strongly dominated by the < 10 $\mu$m size class (29 724–155 764 cells mL$^{-1}$, compared to 5 795 cells mL$^{-1}$ for E02 – no data for E01). By contrast, E01 and E02 saw a greater dominance by the > 10 $\mu$m size fraction (90–268.4 cells mL$^{-1}$, compared to 37.4–83.2 cells mL$^{-1}$ for E03–E05) (Richier et al., 2014a). The results of the ANOVA shown in Fig. 6 show that the sites also fall within these groupings based on the relative DMS response to high CO$_2$. As discussed in the previous section, size-related differences in community structure (< 10 $\mu$m E03–E05 vs. > 10 $\mu$m E01–E02) and differing responses/resiliencies of phytoplankton species are likely to contribute to differences in sensitivity to increasing CO$_2$ between the stations.

Further insight may be gained by considering the influence of microbial community processes on the observed differences between the stations. We observed differences in gross DMS production rates (dark) between stratified (E01, E03, E05) and mixed (E02, E04) stations. Measurements made at 0 h are most representative of in situ rates, and showed low levels of GP of 0.1–0.7 nmol L$^{-1}$ d$^{-1}$ at the mixed sites (E02 and E04) but relatively high rates of 4.9 and 8.7 nmol L$^{-1}$ d$^{-1}$ at E03 and E05 (no data for E01) (see Table 5 and Table S6 in the Supplement). This may indicate that the communities of stratified sites have a greater potential DMS yield, perhaps due to an increased importance of bacterial cleavage of DMSP to DMS in stratified waters, compared to a dominance of bacterial catabolism via the de/de pathway at the mixed sites. This may, for example, explain the large difference in relative DMS response between E01 and E04, despite a similar relative decrease in DMSP$_t$ (Fig. 6).

However, the above explanations are not sufficient to explain all of the variation in relative DMS response to high CO$_2$ that was observed across the five stations, as a range of interacting factors are likely to be involved. Further investigation is now needed to verify the extent of the influence of phytoplankton community structure and bacterial processes.
on the sensitivity of surface ocean DMS systems to elevated CO$_2$ and to ascertain how the physical and biogeochemical characteristics of different regions may determine this response.

5 Summary and conclusions

We observed that rapid and short-term experimental manipulations of CO$_2$ and [H$^+$] induced consistent, marked increases in DMS and decreases in DMSP, in contrast to results from mesocosm experiments. Mesocosms focus on longer timescales (up to 5 weeks) within the framework of a nutrient-induced phytoplankton bloom. This allows a “winners and losers” dynamic, encouraging shifts to species with greater resilience for change. The resultant DMS(P) response reflects the ensuing taxonomic changes (Vogt et al., 2008; Hopkins et al., 2010; Avgoustidi et al., 2012; Archer et al., 2013). The bioassay results we present represent an “acclimatory” response characterised by the lack of ability of small phytoplankton (< 10 μm) to adapt to the altered carbonate chemistry. Our data are suggestive of an increase in stress-induced algal processes (increased cell permeability resulting in increased DMSP release and cleavage to DMS and/or direct up-regulation of intracellular DLA and DMS release) induced by the rapid changes to carbonate chemistry. However, this cannot be validated without direct measurements, so future studies could include better determination of algal-related processes, including measurements of dissolved DMSP concentrations to give an indication of loss of DMS from phytoplankton cells and direct measurements of DLA (Steinke et al., 2000). In addition, more extensive tracer approaches may be useful to disentangle the fate of DMSP, particularly the balance between DMSP synthesis and breakdown or release by the algae. Finally, the influence of OA effects on grazing and subsequent DMS cycling was not determined during this study and may warrant further investigation.

Although it would be inappropriate to extrapolate our small-scale, short-term results to future DMS sea-to-air fluxes, our study provides further evidence that the DMS system has a capacity to change in the face of future environmental change. Only one modelling study has attempted to quantify the climate response to OA-induced changes to sea surface DMS concentrations (Six et al., 2013). Though an important step forward, the study used model parameterisations based on empirical relationships between DMS and [H$^+$] from mesocosm studies and is thus limited by the low level of understanding of the processes behind the observed responses to OA. The variables and rates we report improve our understanding of the sensitivity of the reduced sulfur cycle to future OA and may contribute to the development of a mechanistic approach to modelling future DMS concentrations (Polimene et al., 2012).

Acknowledgements. This work is a contribution to the UK Ocean Acidification Research Programme (UKOA) which was jointly funded by the Department for Environment, Food and Rural Affairs (Defra), the Natural Environment Research Council (NERC) and the Department for Energy and Climate Change (DECC) under grant agreement no. NE/H017259/1. We are grateful to all members of the scientific party for aiding the success of cruise D366. In particular, we thank Toby Tyrrell as lead PI of the UKOA Sea Surface Consortium, Eric Achterberg for leadership as chief scientist, and Sophie Richier and Mark Moore for the design and management of the experimental bioassays. Thanks also to Ross Holland for flow cytometry data, Alex Poulton for primary production data, Michelle Barnett for microscopy counts, Mark Stinchcombe for nutrient data and Cynthia Dumousseaud for carbonate chemistry measurements. Finally, we thank the captain and the crew of the RRS Discovery and the technical staff of the National Marine Facilities for support and assistance during the cruise. We are grateful to Tom Bell and two anonymous referees for comments that helped to significantly improve the manuscript.

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F. E. Hopkins and S. D. Archer: Consistent increase in dimethyl sulfide (DMS) 4939


