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Methanethiol, dimethyl sulfide and acetone over biologically productive waters in the southwest Pacific Ocean

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Abstract. Atmospheric methanethiol (MeSHₐ), dimethyl sulfide (DMSₐ) and acetone (acetoneₐ) were measured over biologically productive frontal waters in the remote southwest Pacific Ocean in summertime 2012 during the Surface Ocean Aerosol Production (SOAP) voyage. MeSHₐ mixing ratios varied from below the detection limit (<10 ppt) up to 65 ppt and were 3 %–36 % of parallel DMSₐ mixing ratios. MeSHₐ and DMSₐ were correlated over the voyage ($R^2 = 0.3$, slope = 0.07) with a stronger correlation over a coccolithophore-dominated phytoplankton bloom ($R^2 = 0.5$, slope 0.13). The diurnal cycle for MeSHₐ shows similar behaviour to DMSₐ with mixing ratios varying by a factor of ∼2 according to time of day with the minimum levels of both MeSHₐ and DMSₐ occurring at around 16:00 LT (local time, all times in this paper are in local time). A positive flux of MeSH out of the ocean was calculated for three different nights and ranged from 3.5 to 5.8 µmol m⁻² d⁻¹, corresponding to 14 %–24 % of the DMS flux (MeSH / (MeSH + DMS)). Spearman rank correlations with ocean biogeochemical parameters showed a moderate-to-strong positive, highly significant relationship between both MeSHₐ and DMSₐ with seawater DMS (DMSₐ) and a moderate correlation with total dimethylsulfoniopropionate (total DMSP). A positive correlation of acetoneₐ with water temperature and negative correlation with nutrient concentrations are consistent with reports of acetone production in warmer subtropical waters. Positive correlations of acetoneₐ with cryptophyte and eukaryotic phytoplankton numbers, and high-molecular-weight sugars and chromophoric dissolved organic matter (CDOM), suggest an organic source. This work points to a significant ocean source of MeSH, highlighting the need for further studies into the distribution and fate of MeSH, and it suggests links between atmospheric acetone levels and biogeochemistry over the mid-latitude ocean.

In addition, an intercalibration of DMSₐ at ambient levels using three independently calibrated instruments showed ∼15 %–25 % higher mixing ratios from an atmospheric pressure ionisation chemical ionisation mass spectrometer (mesoCIMS) compared to a gas chromatograph with a sulfur chemiluminescence detector (GC-SCD) and proton transfer reaction mass spectrometer (PTR-MS). Some differences were attributed to the DMSₐ gradient above the sea surface and differing approaches of integrated versus discrete measurements. Remaining discrepancies were likely due to different calibration scales, suggesting that further investigation of the stability and/or absolute calibration of DMS standards used at sea is warranted.
1 Introduction

Volatile organic compounds (VOCs) are ubiquitous in the atmosphere, and they have a central role in processes affecting air quality and climate, via their role in formation of secondary organic aerosol and tropospheric ozone. The role of the ocean in the global cycle of several VOCs is becoming increasingly recognised, with recent studies showing that the ocean serves as a major source, sink, or both for many pervasive and climate-active VOCs (Law et al., 2013; Liss and Johnson, 2014; Carpenter and Nightingale, 2015).

The ocean is a major source of reduced volatile sulfur gases and the most well-studied of these is dimethyl sulfide (DMS) (CH$_3$SCH$_3$), with a global ocean source of $\sim$28 Tg S a$^{-1}$ (Lee and Brimblecombe, 2016). Since the publication of the CLAW hypothesis (Carlson et al., 1987), which proposed a climate feedback loop between ocean DMS concentrations and cloud droplet concentrations and albedo, extensive investigations have been undertaken into DMS formation and destruction pathways, ocean–atmosphere transfer, atmospheric transformation, and impacts on chemistry and climate (Law et al., 2013; Liss and Johnson, 2014; Carpenter et al., 2012; Quinn and Bates, 2011). Methanethiol or methyl mercaptan (MeSH) (CH$_3$SH) is another reduced volatile organic sulfur gas which originates in the ocean, with a global ocean source estimated to be $\sim$17 % of the DMS source (Lee and Brimblecombe, 2016).

The MeSH ocean source is twice as large as the total of all anthropogenic sources (Lee and Brimblecombe, 2016). However, the importance of ocean-derived MeSH as a source of sulfur to the atmosphere, and the impact of MeSH and its oxidation products on atmospheric chemistry and climate, is not well understood.

DMS and MeSH in seawater (DMS$_{sw}$ and MeSH$_{sw}$) are both produced from precursor dimethylsulfoniopropionate (DMSP), which is biosynthesised by different taxa of phytoplankton and released into seawater as a result of ageing, graz ing or viral attack (Yoch, 2002). DMSP is then degraded by bacterial catabolism (enzyme-catalysed reaction) via competing pathways that produce either DMS or MeSH (Yoch, 2002). Recent research showed that the bacterium Pelagibacter can simultaneously catabolise both DMS$_{sw}$ and MeSH$_{sw}$ (Sun et al., 2016), although it is not known how widespread this phenomenon is. DMS may also be produced by phytoplankton that directly cleave DMSP into DMS (Alcolombri et al., 2015). Once released, MeSH$_{sw}$ and DMS$_{sw}$ undergo further reaction in seawater. These compounds may be assimilated by bacteria, converted to dissolved non-volatile sulfur, be photochemically destroyed, or, in the case of MeSH$_{sw}$, react with dissolved organic matter (DOM) (Kiene and Linn, 2000; Kiene et al., 2000; Flock and Andreae, 1996). MeSH$_{sw}$ has a much higher loss rate constant than DMS$_{sw}$, with a lifetime of the order of minutes to an hour compared to approximately days for DMS$_{sw}$ (Kiene, 1996; Kiene and Linn, 2000). A fraction ($\sim$10 %) of DMS$_{sw}$ ventilates to the atmosphere where it can influence particle numbers and properties through its oxidation products (Simó and Pedrós-Alió, 1999; Malin, 1997). The fraction of MeSH$_{sw}$ ventilating to the atmosphere is poorly constrained.

While DMS$_{sw}$ measurements are relatively widespread, only a few studies have measured MeSH$_{sw}$. During an Atlantic Meridional Transect cruise in 1998 (Kettle et al., 2001) MeSH$_{sw}$ was higher in coastal and upwelling regions with the ratio of DMS$_{sw}$ to MeSH$_{sw}$ varying from unity to 30. Leck and Rodhe (1991) also reported ratios of DMS$_{sw}$/MeSH$_{sw}$ of 16, 20 and 6 in the Baltic Sea, Kattegat–Skagerrak, and North Sea, respectively. The drivers of this variability are unknown, but they are likely due to variation in the dominant bacterial pathway and/or spatial differences in degradation processes. More recent MeSH$_{sw}$ measurements in the northeast subarctic Pacific Ocean showed that the ratio of DMS$_{sw}$/MeSH$_{sw}$ varied from 2 to 5, indicating that MeSH$_{sw}$ was a significant contributor to the volatile sulfur pool in this region (Kiene et al., 2017). MeSH$_{sw}$ measurements from these three studies (Kettle et al., 2001; Leck and Rodhe, 1991; Kiene et al., 2017) were also used to calculate the ocean–atmosphere flux of MeSH, assuming control from the water side. The flux of MeSH/ (MeSH + DMS) ranged from 4 % to 5 % in the Baltic Sea and Kattegat, and it was 11 % in the North Sea (Leck and Rodhe, 1991), 16 % over the North Atlantic–South Atlantic transect (Kettle et al., 2001) and $\sim$15 % over the northeast subarctic Pacific (Kiene et al., 2017). In a review of global organo-sulfide fluxes, Lee and Brimblecombe (2016) estimated that ocean sources provide over half of the total global flux of MeSH to the atmosphere, with a total of 4.7 Tg S a$^{-1}$; however, this estimate is based on a voyage-average value from a single study in the North Atlantic–South Atlantic (Kettle et al., 2001) in which flux measurements varied by several orders of magnitude.

There are very few published atmospheric measurements of MeSH$_a$ over the ocean. To the best of our knowledge, the only prior MeSH$_a$ measurements over the ocean were made in 1986 over the Drake Passage and the coastal and inshore waters west of the Antarctic Peninsula (Berresheim, 1987). MeSH$_a$ was detected occasionally at up to 3.6 ppt, which was roughly 3 % of the measured atmospheric DMS$_a$ levels (Berresheim, 1987).

Once MeSH$_{sw}$ is transferred from ocean to atmosphere (MeSH$_a$), the main loss pathway for MeSH$_a$ is via reaction with OH and NO$_3$ radicals. MeSH$_a$ reacts with OH at a rate 2–3 times faster than DMS, and as such MeSH$_a$ has an atmospheric lifetime of only a few hours (Lee and Brimblecombe, 2016). The oxidation pathways and products that result from MeSH$_a$ degradation are still highly uncertain (Lee and Brimblecombe, 2016; Tyndall and Ravishankara, 1991), though they may be somewhat similar to DMS (Lee and Brimblecombe, 2016). This leads to uncertainty around the final atmospheric fate of the sulfur emitted via MeSH and also the overall impact of MeSH$_a$ oxidation on atmospheric chemi-
istry, particularly in regions when MeSH is a significant proportion of total sulfur emitted.

In the case of acetone, positive fluxes from the ocean have been observed in biologically productive areas (Taddei et al., 2009) and over some subtropical ocean regions (Beale et al., 2013; Yang et al., 2014a; Tanimoto et al., 2014; Schlundt et al., 2017); however, in other subropical regions, and generally in oligotrophic waters and at higher latitudes, net fluxes are zero (e.g. ocean and atmosphere in equilibrium) or negative (transfer of acetone into ocean) (Yang et al., 2014a, b; Marandino et al., 2005; Beale et al., 2015; Schlundt et al., 2017). Atmospheric acetone (acetone$_w$) also has significant terrestrial sources including direct biogenic emissions from vegetation, oxidation of anthropogenic and biogenic hydrocarbons (predominantly alkanes), and biomass burning (Fischer et al., 2012). In the ocean, acetone$_w$ is produced photochemically from chromophoric dissolved organic matter (CDOM), either directly by direct photolysis or via photosensitiser reactions (Zhou and Mopper, 1997; Dixon et al., 2013; de Bruyn et al., 2012; Kieber et al., 1990). There is also evidence of direct biological production by marine bacteria (Nemecek-Marshall et al., 1995) and phytoplankton (Schlundt et al., 2017; Sinha et al., 2007; Halsey et al., 2017). Furthermore, acetone$_w$ has been found to decrease with depth (Beale et al., 2015; Yang et al., 2014a; Beale et al., 2013; Williams et al., 2004), pointing to the importance of photochemistry and/or biological activity as the source. Studies have shown that acetone$_w$ production linked to photosynthetically active radiation (PAR) and net shortwave radiation (Sinha et al., 2007; Beale et al., 2015; Zhou and Mopper, 1997), and Beale et al. (2015) found higher acetone$_w$ concentrations in spring and summer compared to autumn and winter. Removal processes include uptake of acetone by bacteria as a carbon source (Beale et al., 2013, 2015; Halsey et al., 2017; Dixon et al., 2013), gas transfer into the atmosphere, vertical mixing into the deep ocean and photochemical destruction (Carpenter and Nightingale, 2015).

There are relatively few observations of acetone$_w$ and acetone$_a$ over the remote ocean, particularly in mid- and high-latitude regions. An understanding of the spatial distribution of acetone is particularly important due to the high degree of regional variation in the direction and magnitude of the acetone flux.

The Surface Ocean Aerosol Production (SOAP) voyage investigated the relationship between ocean biogeochemistry and aerosol and cloud processes in a biologically productive but under sampled region in the remote southwest Pacific Ocean (Law et al., 2017). In this work, we present measurements of DMS$_a$, MeSH$_a$ and acetone$_a$, including the largest observed mixing ratios of MeSH$_a$ in the marine boundary layer to date. We explore the relationship between DMS$_a$, MeSH$_a$ and acetone$_a$ as well as the relationship with ocean biogeochemical parameters. In particular, we investigate links between MeSH$_a$ and its precursor DMS$_w$ for the first time. We explore whether variability in acetone$_a$ is linked to biogeochemistry, including warmer subtropical waters and organic precursors such as CDOM as has been reported elsewhere.

Given the large uncertainty in the oceanic budget of MeSH, we estimate the importance of MeSH as a source of atmospheric sulfur in this region and compare results with other studies. Finally, we present results from a DMS$_a$ method comparison which was undertaken at sea between three independently calibrated measurement techniques.

2 Method
2.1 Voyage

The Surface Ocean Aerosol Production (SOAP) voyage took place on the NIWA RV Tangaroa over the biologically productive frontal waters of Chatham Rise (44$^\circ$ S, 174–181$^\circ$ E), east of New Zealand in the southwest Pacific Ocean. The 23-day voyage took place during the austral summer in February–March 2012. The scientific aim was to investigate interactions between the ocean and atmosphere, and as such the measurement programme included comprehensive characterisation of ocean biogeochemistry, measurement of ocean–atmosphere gas and particle fluxes, and measurement of distribution and composition of trace gases and aerosols in the marine boundary layer (MBL) (Law et al., 2017). During the voyage, NASA MODIS ocean colour images and underway sensors were used to identify and map phytoplankton blooms. Three blooms were intensively targeted for measurement: (1) a dinoflagellate bloom with elevated Chl $a$, DMS$_w$ and $p$CO$_2$ drawdown, and high irradiance (bloom 1 – B1); (2) a coccolithophore bloom (bloom 2 – B2); and (3) a mixed community bloom of coccolithophores, flagellates, and dinoflagellates sampled before (bloom 3a – B3a) and after (bloom 3b – B3b) a storm. For further voyage and measurement details, see Law et al. (2017).

2.2 PTR-MS

A high-sensitivity proton transfer reaction mass spectrometer (PTR-MS) (Ionicon Analytik) was used to measure DMS, acetone and methanethiol. The PTR-MS sampled from a 25 m 3/8 in. i.d. PFA (perfluoroalkoxy) inlet line, which drew air from the crow’s nest of the vessel 28 m above sea level (a.s.l.) at 10 L min$^{-1}$. A baseline switch based on relative wind speed and direction was employed to minimise flow of ship exhaust down the inlet (see Lawson et al., 2015).

The PTR-MS instrument parameters were as follows: inlet and drift tube temperature of 60$^\circ$C, a 600 V drift tube and 2.2 mbar drift tube pressure (E / N = 133 Td, townsend). The O$_2$ signal was <1 % of the primary ion H$_3$O$^+$ signal. DMS, acetone and MeSH were measured at m/z 63, m/z 59 and m/z 49, respectively, with a dwell time of 10 s. From day-of-year (DOY) 43–49, 19 selected ions including m/z 59 and m/z 63 were measured, resulting in 17 mass scans per
hour; however, from DOY 49 the PTR-MS measured in scan mode from m/z 21 to 155, allowing three full mass scans per hour. As such, MeSH measurements (m/z 49) were made only from DOY 49 onward.

VOC-free air was generated using a platinum-coated glass wool catalyst heated to 350°C; four times per day this air was used to measure the background signal resulting from interference ions and outgassing of materials. An interpolated background signal was used for background correction. Calibrations of DMS and acetone were carried out daily by diluting calibration gas into VOC-free ambient air (Galbally et al., 2007). Calibration gases used were a custom ∼ 1 ppm VOC mixture in nitrogen containing DMS and acetone (Scott Specialty Gases) and a custom ∼ 1 ppm VOC calibration mixture in nitrogen containing acetone (Apel Riemer). The calibration gas accuracy was ± 5 %. A calibration gas for MeSH was not available during this voyage. The PTR-MS response to a given compound is dependent on the chemical ionisation reaction rate, defined by the collision rate constant and to a given compound is dependent on the chemical ionisation in nitrogen containing acetone (Apel Riemer). The calibration gas accuracy was ± 5 %. MeSH has a similar structure and physical properties to DMS at pH < 10 (Sect. 3.2), and so inlet losses are likely to be similar. These small (< 5 %) losses could lead to a small underestimation in reported mixing ratios of DMSu, acetone, and MeSHu.

2.3 DMS intercomparison

During the SOAP voyage, DMSu measurements were made using three independently calibrated instruments: atmospheric pressure ionisation chemical ionisation mass spectrometer (mesoCIMS) from the University of California Irvine (UCI) (Bell et al., 2013, 2015), an Ionicon PTR-MS operated by the Commonwealth Scientific and Industrial Research Organisation (CSIRO) (Lawson et al., 2015) and a HP gas chromatograph with sulfur chemiluminescence detector (GC-SCD) operated by the National Institute of Water and Atmospheric Research (NIWA) (Walker et al., 2016).

Details of the mesoCIMS and GC-SCD measurement systems are provided by Bell et al. (2015) and Walker et al. (2016) with a brief description provided here. The mesoCIMS instrument (Bell et al., 2013) ionises DMS to DMS-H+: m/z = 63) by atmospheric pressure proton transfer from H3O+ by passing a heated air stream over a radioactive nickel foil (Ni-63). The mesoCIMS drew air from the eddy covariance set-up on the bow mast at approximately 12 m a.s.l. The inlet was a 1/2 in. i.d. PFA tube with a total inlet length of 19 m and a turbulent flow at 90 slpm (standard litres per minute). The mesoCIMS subsampled from the inlet at 1 L m⁻¹. A gaseous tri-deuterated DMS standard (D3DMS) was added to the air sample stream at the entrance to the inlet. The internal standard was ionised and monitored continuously in the mass spectrometer at m/z = 66, and the atmospheric DMS mixing ratio was computed from the measured 63/66 ratio. The internal standard was delivered from a high-pressure aluminium cylinder and calibrated against a DMS permeation tube prior to and after the cruise (Bell et al., 2015).

The GC-SCD system included a semiautomated purge and trap system, a HP 6850 gas chromatograph with cryogenic pre-concentrator and thermal desorber, and sulfur chemiluminescence detection (Walker et al., 2016). The system was employed during the voyage for discrete DMS seawater measurements and gradient flux measurement bag samples (Smith et al., 2018). The system was calibrated using an internal methylethylsulfide (MES) permeation tube and external DMS permeation tube located in a Dynacalibrator® with a twice daily five-point calibration and a running standard every 12 samples (Walker et al., 2016).

A DMS measurement intercomparison between the mesoCIMS, GC-SCD and PTR-MS was performed during the voyage on DOY 64 and DOY 65. Tedlar bags (70 L) with blackout polythene covers were filled with air containing DMS at sub-parts-per-billion levels, and they were sequentially distributed between all instruments for analysis within a few hours. On DOY 64, two bags were prepared including ambient air filled from the foredeck and a DMS standard prepared using a permeation device (Dynacalibrator®) and dried compressed air (DMS range 384–420 ppt from permeation uncertainty). On DOY 65, two additional bags were prepared including one ambient air sample from the
foredeck with tri-deuterated DMS added and a DMS standard prepared using the Dynacalibrator and dried compressed air (DMS range 331–363 ppt). MesoCIMS values are not available for DOY 64 due to pressure differences between bag and instrument calibration measurements; this was resolved by using an internal standard on DOY 65. For those analyses, the mesoCIMS and PTR-MS measured DMS at m/z 63 and tri-deuterated DMS at m/z 66, while the GC-SCD measured both DMS and deuterated DMS as a single peak.

2.4 Biogeochemical measurements in surface waters

Continuous seawater measurements were obtained from surface water sampled by an intake in the vessel’s bow at a depth of ~7 m during the SOAP voyage and included underway temperature and salinity (Sea-Bird thermosalinograph SBE-21), underway chlorophyll a (Chl a) and backscatter (Wetlabs (Sea-Bird) ECOrtripllet), and dissolved DMS (DMS-sw) (miniCIMS) (Bell et al., 2015). Quenching obscured the Chl a signal during daylight when irradiance was >50 W m$^{-2}$.

The following parameters were measured in surface waters (depths 2–10 m) in discrete samples from Niskin bottles on a conductivity–temperature–depth (CTD) rosette: nutrients according to methods described in Law et al. (2011), particulate nitrogen concentration (Nodder et al., 2016), phytoplankton speciation, groups and numbers (optical microscopy of samples preserved in Lugol’s solution) (Safi et al., 2007), and flow cytometry (Hall and Safi, 2001). In addition, the organic parameters measured included high-molecular-weight (HMW) reducing sugars (Somogyi, 1926, 1952; for details see Burrell, 2015), DMSP (Walker et al., 2016) and CDOM measured using a liquid waveguide capillary cell (Gall et al., 2013). See Table S1 for measurement specifications and Law et al. (2017) for further details and results for these parameters.

3 Results and discussion

3.1 DMS atmospheric intercalibration

This section describes a comparison of DMS$_a$ measurements from bag samples of ambient air and DMS standard mixtures (analysed by GC-SCD, PTR-MS and mesoCIMS; see Sect. 2), as well as comparison of ambient DMS$_a$ measurements (PTR-MS and mesoCIMS).

3.1.1 Comparison of bag samples

Table 1 summarises the comparison between the GC-SCD, PTR-MS and mesoCIMS instruments for ambient and DMS standard bags prepared and analysed on DOY 64 and 65 (see Sect. 2.2). The highest DMS levels were measured by the mesoCIMS, with GC-SCD and PTR-MS being ~20%–25% and ~20%–30% lower, respectively. The GC-SCD and PTR-MS agreed reasonably well, with a mean difference of 5% (range 0%–10%) between instruments for different diluted standard and ambient air bags. There was no clear influence of dry versus humid (ambient) bag samples on the differences between instruments.

3.1.2 Comparison of in situ ambient measurements

Measurements from the PTR-MS and mesoCIMS were interpolated to a common time stamp for comparison, and differences were examined only where data were available for both instruments. PTR-MS results for DMS were reported for 10 s every 4 min until DOY 49 and then 10 s every 20 min until the end of the voyage (Sect. 2.2). The mesoCIMS measured DMS continuously and reported 10 min averages. As such the PTR-MS measured only a “snapshot” of the DMS$_a$ levels in each measurement cycle of 4 or 20 min. This was a potential source of difference between the two instruments when DMS levels changed rapidly (Bell et al., 2015).

The mesoCIMS was deployed primarily for DMS eddy covariance measurements, while the PTR-MS was deployed to measure atmospheric mixing ratios of a range of VOCs. As such, the mesoCIMS was situated on the foredeck and sampled from the eddy covariance set-up on the bow mast (12 m a.s.l.), while the PTR-MS was sited further back in the vessel and sampled from the crow’s nest (28 m a.s.l.). Therefore, due to different intake heights, a further source of the difference between the PTR-MS and mesoCIMS measurements is likely due to vertical gradients in DMS caused by turbulent mixing of the local surface DMS flux into the atmospheric surface layer. On days with a strong DMS source and/or more stable stratification in the boundary layer, a significant decrease with height is expected (Smith et al., 2018).

If all the DMS observed was due to local emissions, the vertical gradient would be described by Eq. (2) from Smith et al. (2018):

$$F \equiv -u^*C^* = -u^* \frac{k}{\varphi_c(z/L)} \left( \frac{\partial C}{\partial \ln z} \right),$$

(1)

where $u^*$ is friction velocity, $C^*$ is scaling parameter for gas concentration, $k$ is the von Kármán constant, $\varphi_c$ is the stability function for mass, $z$ is the height above mean water level and $L$ is the Monin–Obukhov scaling length representing atmospheric stability. Atmospheric stability is a measure of the degree of vertical motion in the atmosphere, where $z/L = 0$ indicates neutral stability, $z/L > 0$ indicates a stable atmosphere and $z/L < 0$ indicates an unstable atmosphere.

Figure 1 shows wind speed, absolute wind direction and atmospheric stability, DMS$_a$ levels from the voyage measured by PTR-MS and mesoCIMS, relative percent difference between the two measurements (normalised to the mesoCIMS), and observed absolute difference in DMS$_a$ between the two measurements, as well as the expected calculated difference (Eq. 1) between the two measurements due to the DMS$_a$ concentration gradient.
Table 1. Results of the DMS bag sample intercomparison study undertaken during the SOAP voyage. Note that a 1 s PTR-MS dwell time for m/z 63 and m/z 66 was used during the intercomparison compared to the 10 s during ambient measurements; as such, the PTR-MS standard deviation reported here is expected to be ∼3 times higher than during ambient measurements. “Total” refers to the ambient DMS + spiked tri-deuterated DMS bag sample on DOY 65.

| DOY | Comparison          | GC-SCD (ppt) | PTR-MS (ppt) | mesoCIMS (ppt) | GC-SCD / PTR-MS | PTR-MS / mesoCIMS | GC-SCD / mesoCIMS |
|-----|---------------------|--------------|--------------|----------------|-----------------|------------------|------------------|
| 64  | Standard (dry)      | 354 ± 6      | 339 ± 64     | n/a            | 1.04 ± 0.2      | n/a              | n/a              |
| 65  | Standard (dry)      | 289 ± 2      | 262 ± 43     | 383 ± 30       | 1.1 ± 0.18      | 0.68 ± 0.12      | 0.75 ± 0.06      |
| 64  | Ambient             | 168 ± 5      | 158 ± 49     | n/a            | 1.06 ± 0.33     | n/a              | n/a              |
| 65  | Ambient             | n/a          | 127 ± 43     | 141 ± 5        | 0.90 ± 0.30     | n/a              | n/a              |
|     | +Tri-deuterated DMS | n/a          | 197 ± 49     | 260 ± 2        | n/a             | 0.76 ± 0.19      | n/a              |
|     | Total               | 323 ± 9      | 324 ± 66     | 401 ± 6        | 1.0 ± 0.2       | 0.81 ± 0.16      | 0.81 ± 0.03      |

Figure 1. From top to bottom, wind speed and stability (a), DMS
measurements from mesoCIMS and PTR-MS (b), relative difference
(normalised to mesoCIMS) according to absolute wind direction (c), and absolute observed and calculated difference between mesoCIMS and PTR-MS (d), taking into account the expected DMS concentration gradient (Eq. 1).

The mesoCIMS and PTR-MS DMS₃ data showed similar temporal behaviour over the voyage (Fig. 1). From DOY 44 to 46 there was an average of 50% (±10%) relative difference between measurements, yet on DOY 47 this difference decreased suddenly to an average of ∼20% (±20%).

Overall, agreement between instruments improved with time during the voyage, with differences of several hundred parts per trillion (ppt) of DMS observed in the first few days decreasing to differences of only 10–20 ppt by the end of the voyage. The agreement between instruments improves with increasing wind speeds (Fig. 1). The expected calculated difference between DMS₃ at the two inlet heights due to the DMS concentration gradient also decreases throughout the voyage. This indicates that the increasing agreement between instruments during the voyage was likely influenced by a progressively well-mixed atmosphere leading to weaker DMS vertical gradients.

The reason for the improved agreement between mesoCIMS and PTR-MS at DOY 47 is unlikely due to a decrease in the DMS concentration gradient (Fig. 1d, bottom...
Figure 2. (a) DMS<sub>a</sub> measured by mesoCIMS (x axis) and PTR-MS (y axis) and PTR-MS (y axis) DMS data corrected for the expected concentration gradient (observed PTR-MS DMS + calculated delta DMS). Dashed lines represent the reduced major axis regression and solid lines represent a 1:1 relationship.

Table 2. MeSH<sub>a</sub>, DMS<sub>a</sub> and acetone<sub>a</sub> measured with PTR-MS during the SOAP voyage, with reaction rate constant for OH and calculated lifetime with respect to OH.

| Substance | Mean range (ppt) | k<sub>OH</sub> (cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup>) | Lifetime (days) |
|-----------|-----------------|---------------------|----------------|
| MeSH      | 18 (BDL–65)     | 3.40 × 10<sup>-11</sup> | 0.4            |
| DMS       | 208 (BDL–957)   | 1.29 × 10<sup>-11</sup> | 1              |
| Acetone   | 237 (54–1508)   | 2.20 × 10<sup>-13</sup> | 60             |

BDL represents below detection limit. *Reaction rate constants from Atkinson (1997) (MeSH), Berresheim et al. (1987) (DMS) and Atkinson (1996) (acetone).

panel), but it is more likely due to changes in instrument calibration or other differences. However, careful inspection of the instrument parameters, configurations and calibration responses prior to DOY 47 did not identify the cause of the disagreement.

Figure 2a shows paired DMS<sub>a</sub> data from the mesoCIMS versus PTR-MS over the whole voyage, and Fig. 2b shows paired mesoCIMS data versus PTR-MS data converted to the same height as the mesoCIMS with the expected DMS difference calculated from the eddy covariance estimate of DMS flux (from mesoCIMS) and eddy diffusivity (PTR-MS DMS<sub>a</sub> + calculated difference between the two intake heights). The reduced major axis regression relationship between the two measurement systems for uncorrected data gives a slope of 0.74 ± 0.02, while for the corrected data the relationship gives 0.81 ± 0.02 (R<sup>2</sup> = 0.69). The gradient-corrected slope agrees with the ambient bag sample ratio from the method comparison (PTR-MS / mesoCIMS = 0.81 ± 0.16) (Table 1). Correcting for the DMS gradient improved the comparison between PTR-MS and mesoCIMS. The remaining ∼20% difference is likely due to instrument calibration differences and differing approaches of integrated versus discrete measurements.

There was no obvious impact of absolute wind direction on the differences observed between measurement systems. Note that due to the baseline switch which was employed to avoid sampling ship exhaust down the PTR-MS inlet (Lawson et al., 2015), the PTR-MS did not sample during certain relative wind directions. However, this does not affect the comparison which was undertaken only when data were available for both instruments.

3.2 Ambient atmospheric data

Atmospheric mixing ratios of MeSH<sub>a</sub>, DMS<sub>a</sub> and acetone<sub>a</sub> are shown along the voyage track in Fig. 3 with bloom locations highlighted. Figure 4 shows a time series of MeSH<sub>a</sub>, DMS<sub>a</sub>, acetone<sub>a</sub> and MeSH<sub>a</sub> / DMS<sub>a</sub> (all measured with PTR-MS), as well as DMS<sub>s</sub> (miniCIMS) from Bell et al. (2015), Chl<sub>a</sub>, irradiance, wind speed, wind direction, and sea and air temperature. Note that MeSH<sub>a</sub> measurements started on DOY 49 (the last day of bloom B1). The fraction of backward trajectories arriving at the ship that had been in contact with land masses in the previous 10 days is also shown with a value of 0 indicating no contact with land masses in the preceding 10 days. This was calculated using the Lagrangian Numerical Atmospheric-dispersion Modelling Environment (NAME) for the lower atmosphere (0–100 m) as time-integrated particle density (g s m<sup>-3</sup>), every 3 h from the ship location (Jones et al., 2007) as shown in Law et al. (2017). Where air contacted land masses, this was the New Zealand land mass in almost all cases.

MeSH<sub>a</sub> ranged from below detection limit (<10 ppt) to 65 ppt, DMS<sub>a</sub> ranged from below detection limit (∼22 ppt) up to 957 ppt, and acetone<sub>a</sub> ranged from 50 to 1500 ppt (Table 2). The ratio of MeSH<sub>a</sub> to DMS<sub>a</sub> ranged from 0.03 to 0.36 (mean 0.14) for measurements when both were above the minimum detectable limit. Periods of elevated DMS<sub>a</sub> gen-
Figure 3. Atmospheric mixing ratios of (a) MeSH$_a$, (b) DMS$_a$ and (c) acetone$_a$ as a function of the voyage track. Location of the blooms are shown.

erally correspond to periods of elevated DMS$_{sw}$. Both DMS$_a$ and DMS$_{sw}$ were very high during B1, during the transect to B2, and the first half of B2 occupation. MeSH$_a$ variability broadly correlates with DMS$_a$ and DMS$_{sw}$, with highest levels during B2 (no data available for B1). The highest acetone$_a$ levels observed occur during B2, and a broad acetone peak during B1 of 700 ppt (≈ DOY 49) overlaps with but is slightly offset from the largest DMS$_a$ peak during the voyage (≈ 957 ppt). DMS$_a$, acetone$_a$ and MeSH$_a$ were somewhat lower during B3a and lowest during B3b (the post-storm part of that bloom B3) (see Law et al., 2017). In general, DMS$_a$ levels during B1 were at the upper range of those found in prior studies elsewhere (Lana et al., 2011; Law et al., 2017). MeSH$_a$ levels during B2 ranged from below detection limit (≈ 10 ppt) up to 65 ppt (mean 25 ppt), which is substantially higher than the only comparable measurements from the Drake Passage and the coastal and inshore waters west of the Antarctic Peninsula (3.6 ppt) (Berresheim, 1987). The average acetone$_a$ levels during this study were broadly comparable to those from similar latitudes reported in the South Atlantic and Southern Ocean (Williams et al., 2010) and at Cape Grim (Galbally et al., 2007). Acetone$_a$ during SOAP was generally lower than at similar latitudes at Mace Head (Lewis et al., 2005), the southern Indian Ocean (Colomb et al., 2009) and also the marine subtopics (Read et al., 2012; Schlundt et al., 2017; Warneke and de Gouw, 2001; Williams et al., 2004).

There were two occasions when elevated acetone$_a$ corresponded closely to an increased land influence – during B1 on DOY 48–49 (maximum land influence 12 %) and DOY 60 (maximum land influence 20 %) (Fig. 4). Both these periods corresponded to winds from the north, and back trajectories show that the land mass contacted was the southern tip of New Zealand’s North Island (including the city of Wellington and the northern section of the South Island in both cases). The acetone measured during these periods may have been emitted from anthropogenic and biogenic sources as well as from photochemical oxidation of hydrocarbon precursors (Fischer et al., 2012). The acetone enhancement relative to the degree of land influence was higher on DOY 48–49 than DOY 60 possibly due to different degrees of dilution of the terrestrial plume or different terrestrial source strengths.

The period with the highest acetone levels during B2 (1508 ppt) corresponds with a period of negligible land in-
Figure 4. Time series of measurements during the SOAP voyage according to DOY. Atmospheric DMS and MeSH measurements below detection limit have had half detection limit substituted. WS represents wind speed, wind dir represents wind direction, Irrad. represents irradiance and Chl \( a \) represents chlorophyll \( a \).
Differences behaving a much faster loss rate in seawater than DMS \(_{sw}\) (Lizotte et al., 2017). The relative loss rates of lysis and senescence) or direct processes (algal DMSP-late afternoon minimum has been observed in many previous studies and is attributed to photochemical destruction by OH. This includes Cape Grim baseline station, which samples air from the Southern Ocean (average minimum and maximum \(~\sim 40–70\) ppt) (Ayers and Gillett, 2000), over the tropical Indian ocean (average minimum and maximum \(~\sim 25–60\) ppt (Warneke and de Gouw, 2001) and at Kiritimati in the tropical Pacific (average minimum and maximum \(~\sim 120–200\) ppt) Bandey et al., 1996). The higher atmospheric levels in this study are due to high DMS \(_{sw}\) concentrations (> 15 nM). The amplitude of the DMS diurnal cycle is likely to have been influenced by stationing the vessel over blooms with high DMS \(_{sw}\) from 08:00 each day and regional mapping of areas with lower DMS \(_{sw}\) overnight (Law et al., 2017).

The diurnal cycle for MeSH \(_{a}\) and acetone \(_{a}\) is entrainment of air from the free troposphere into the MBL. For short-lived DMS and MeSH (Table 2), free tropospheric air is most likely to be depleted in these gases compared to air sampled close to the ocean surface. Acetone is relatively long lived (Table 2) and has significant terrestrial sources (Fischer et al., 2012), and so, depending on the origin of the free tropospheric air, it could be enhanced or depleted relative to MBL air.
Figure 5. Correlation between (a) DMS$_a$ and MeSH$_a$ all data (DOY 49 onwards) and (b) DMS$_a$ and MeSH$_a$ bloom (B2) only.

Figure 6. Diurnal cycles of (a) DMS$_a$, (b) MeSH$_a$, (c) acetone$_a$ with land-influenced data removed. Average values from 00:00 to 03:00 are excluded because of lower data collection during this period, due to calibrations and zero air measurements.
of ~2 with the minimum mixing ratio occurring at around 16:00 (the same time as minimum DMS$_a$). The most important sink of MeSH$_a$ is thought to be oxidation by OH (Lee and Brimblecombe, 2016), and the minima in late afternoon may be due to destruction by OH. The decoupling of the DMS and MeSH diurnal cycles between 04:00 and 08:00, with DMS increasing and MeSH decreasing, is likely due to the differing production pathways as well as the possibility of additional sinks for MeSH in the ocean during this time. This period may also have been influenced by mapping areas with lower DMS$_{sw}$ overnight and stationing the vessel over blooms with high DMS$_{sw}$ from 08:00 each day, as described above.

The acetone$_a$ diurnal cycle (Fig. 6c) with land-influenced data removed shows reasonably consistent mixing ratios from the early morning until midday, with an overall increase in acetone levels during the afternoon hours from 14:00 onwards, then decreasing again at night, which is the opposite to the behaviour of DMS$_a$ and MeSH$_a$. Acetone is long lived (~60 days – Table 2) with respect to oxidation by OH. The increase of acetone$_a$ mixing ratios in the afternoon may indicate photochemical production from atmosphere or sea surface precursors but there was no correlation between irradiance and acetone$_a$ during the voyage.

3.3 Flux calculation from nocturnal accumulation of MeSH

MeSH and DMS fluxes ($F$) were calculated according to the nocturnal accumulation method (Marandino et al., 2007). This approach assumes that nighttime photochemical losses are negligible and that sea surface emissions accumulate overnight within the well-mixed marine boundary layer (MBL). Horizontal homogeneity and zero flux at the top of the boundary layer are also assumed. The air–sea flux is calculated from the increase in MeSH and DMS. For example,

$$F = \frac{\partial [\text{MeSH}]}{\partial t} \times h,$$

where [MeSH] is the concentration of MeSH (mol m$^{-2}$) and $h$ is the average nocturnal MBL for the voyage of 1135 m ± 657 m, estimated from nightly radiosonde flights.

DMS and MeSH fluxes were calculated for three nights (DOY 52, 54 and 60) (Table 4) when linear increases in mixing ratios occurred over several hours (Fig. 4). The MeSH flux was lowest on DOY 52 prior to B2 (3.5 ± 2 µmol m$^{-2}$ d$^{-1}$), higher on DOY 60 during B3a (4.8 ± 2.8 µmol m$^{-2}$ d$^{-1}$) and highest on DOY 42 during B2 (5.8 ± 3.4 µmol m$^{-2}$ d$^{-1}$). There are no MeSH measurements during B1. The percentage of MeSH/(DMS + MeSH) emitted varied from 14% for DOY 60 (B3a) up to 23% and 24% for DOY 54 (B2) and DOY 52 (prior to B2), respectively.

For comparison, the DMS fluxes measured using eddy covariance (EC) at the same time are given in Table 4 (Bell et al., 2015). DMS fluxes calculated using the nocturnal accumulation method are within the variability of the EC fluxes (Bell et al., 2015).

The average MeSH flux calculated from this study (4.7 µmol m$^{-2}$ d$^{-1}$) was more than 4 times higher than average MeSH fluxes from previous studies in the North Atlantic–South Atlantic transect (Kettle et al., 2001) and in the Baltic Sea, Kattegat, and North Sea (Leck and Rodhe, 1991) (Table 5). The MeSH fluxes calculated from this work are comparable to maximum values reported by Kettle et al. (2001), which were observed in localised coastal and upwelling regions. The average emission of MeSH compared to DMS (MeSH/(DMS + MeSH)) was higher in this study (20%) compared to previous studies (Table 5) including the Baltic Sea, Kattegat, and North Sea (5%, 4% and 11%); North Atlantic–South Atlantic (16%); and a recent study from the northeast subarctic Pacific (~15%) (Kiene et al., 2017). Note that other sulfur species such as dimethyl disulfide (DMDS), carbon disulfide (CS$_2$) and hydrogen sulfide (H$_2$S) typically make a very small contribution to the total sulfur compared to DMS and MeSH (Leck and Rodhe, 1991; Kettle et al., 2001; Yvon et al., 1993), and so they are neglected from this calculation.

3.4 Correlation with ocean biogeochemistry

To investigate the influence of biogeochemical parameters on atmospheric mixing ratios of MeSH$_a$, DMS$_a$ and acetone$_a$, Spearman rank correlations were undertaken to identify relationships significant at the 95% confidence interval (CI). Table 6 summarises the correlation coefficients and $p$ values for significant correlations. MeSH$_a$, DMS$_a$ and acetone$_a$ data
were averaged for 1 h either side of the CTD water entry time for the analysis.

Sulfur gases MeSH$_{a}$ and DMS$_{a}$ are short lived and so the air–sea flux is controlled by the seawater concentration. By contrast, acetone$_{a}$ is much longer lived in the atmosphere (≈60 days), so the air–sea gradient can be influenced by both oceanic emissions and atmospheric transport from other sources. As such, the variability in acetone$_{a}$ mixing ratios may be driven by ocean–air exchange and/or input of acetone$_{a}$ to the boundary layer from terrestrial sources, the upper atmosphere or in situ production. This means that correlation analyses to explore ocean biogeochemical sources of acetone$_{a}$ may be confounded by atmospheric sources. Removal of land-influenced data reduces the likelihood of this, but observed increases in atmospheric acetone could still be from in situ processes such as oxidation of organic aerosol or mixing from above the boundary layer.

Both MeSH$_{a}$ and DMS$_{a}$ have a strong positive and highly significant relationship with DMS$_{sw}$, and a moderate correlation with discrete measurements of DMSP$_{p}$ (total) and DMSP$_{p}$ (particulate). The correlation of DMS$_{a}$ with DMS$_{sw}$ can be attributed to the positive flux of DMS out of the ocean; however, the correlation of MeSH$_{a}$ with DMS$_{sw}$ is likely due to a common ocean precursor of both gases (DMSP$_{p}$), albeit via different production pathways. DMS$_{a}$ and MeSH$_{a}$ correlate with DMSP$_{p}$ (particulate) but not with DMSP$_{d}$ (dissolved). For DMS$_{a}$, the correlation may reflect that a proportion of the DMS observed was derived directly from phytoplankton rather than being bacterially mediated, which is in agreement with findings by Lizotte et al. (2017); however, as demethylation of DMSP$_{d}$ represents the primary source of MeSH, the lack of correlation is surprising. The latter may reflect MeSH sinks in surface water associated with organics and particles (Kiene, 1996), and this could be confirmed via incubation experiments. DMS$_{a}$ also correlated with particulate nitrogen and showed a moderate negative correlation with silicate that may reflect lower DMS production in diatom-dominated waters.

Acetone$_{a}$ shows a positive correlation with temperature and negative correlation with nutrients. This is consistent with reported sources of acetone$_{sw}$ in warmer subtropical waters (Beale et al., 2013; Yang et al., 2014a; Tanimoto et al., 2014; Schlundt et al., 2017). The positive relationship with organic material including HMW sugars and CDOM may reflect a photochemical ocean source (Zhou and Mopper, 1997; Dixon et al., 2013; de Bruyn et al., 2012; Kieber et al., 1990) or possibly a biological source (Nemecek-Marshall et al., 1995, 1999; Schlundt et al., 2017; Sinha et al., 2007; Halsey et al., 2017) as indicated by the correlations with cryptophyte and picoeukaryote abundance. Correlation with particle backscatter suggests potential links between acetone$_{a}$ and coccolithophores (Sinha et al., 2007). Alternatively, the positive correlations of acetone$_{a}$ with these organic components of sea water may reflect acetone production in the atmosphere from photochemical oxidation of ocean-derived organic aerosols (Pan et al., 2009; Kwan et al., 2006; Jacob et al., 2002). Seawater acetone measurements would allow for further elucidation of the relationships between acetone$_{a}$ and biogeochemical parameters identified in this study. More generally, mesocosm or laboratory studies could be employed to identify the explicit sources and production mechanisms of these gases in Chatham Rise waters.

### Table 5. MeSH flux from this and previous studies (voyage averages).

| Location                        | MeSH flux (µmol m$^{-2}$ d$^{-1}$) | Flux MeSH / MeSH + DMS (%) | Reference                        |
|---------------------------------|----------------------------------|---------------------------|----------------------------------|
| Baltic Sea                      | 0.2                              | 5%                        | Leck and Rodhe (1991)            |
| Kattegat sea                    | 0.8                              | 4%                        |                                  |
| North Sea                       | 1.6                              | 11%                       |                                  |
| North Atlantic–South Atlantic   | 1.2                              | 16%                       | Kettle et al. (2001)             |
| Northeast subarctic Pacific     | Not reported                     | ~15%                      | Kiene et al. (2017)              |
| Southwest Pacific               | 4.7                              | 20%                       | This study                       |

### 4 Implications and conclusions

Mixing ratios of short-lived MeSH$_{a}$ over the remote ocean of up to 65 ppt in this study are the highest observed to date, and they provide evidence that MeSH transfers from the ocean into the atmosphere and may be present at non-negligible levels in the atmosphere over other regions of high biological productivity. The average MeSH flux calculated from this study (4.7 µmol m$^{-2}$ d$^{-1}$) was at least 4 times higher than average MeSH fluxes from previous studies, and it is comparable to maximum MeSH flux values reported in localised coastal and upwelling regions of the North Atlantic–South Atlantic (Kettle et al., 2001) (Table 5). The average emission of MeSH compared to DMS (MeSH / (DMS + MeSH)) was higher in this study (20%) compared to previous studies (4%–16%), indicating that MeSH provides a significant transfer of sulfur to the atmosphere in this region. Taken together with other studies, the magnitude of the ocean MeSH...
Table 6. Spearman rank correlations between acetone, DMS, and MeSH, as well as biogeochemical parameters, using data from the 14 February–4 March 2012 (acetone, DMS) and 20 February–4 March 2012 (MeSH). Correlations shown are significant at the 95 % confidence interval (CI). Correlation coefficient (and \( p \) value) are shown. No entry indicates there was no correlation at 95 % CI. Land-influenced acetone data excluded (see text for details).

|                          | Acetone  | DMS     | MeSH    |
|--------------------------|----------|---------|---------|
| **Positive correlations** |          |         |         |
| Salinity (psu)           | 0.55 (0.005) |   n = 25 |         |
| Sea temperature (°C)     | 0.77 (< 0.0001) |   n = 25 |         |
| Beta-660 backscatter (m⁻² sr⁻¹) | 0.67 (0.004) |   n = 25 |         |
| DMS\(_{sw}\) (nM)        | 0.49 (0.025) |   n = 21 | 0.73 (0.0002) |   n = 22 | 0.59 (0.011) |   n = 18 |
| Chl \( \alpha \)/mixed layer depth | 0.50 (0.014) |   n = 25 |         |
| Particulate nitrogen (mg m⁻³) |          |         | 0.79 (0.048) |   n = 7      |
| Cryptophyte algae (cells mL⁻¹) | 0.47 (0.019) |   n = 25 |         |
| Eukaryotic picoplankton (cells mL⁻¹) | 0.48 (0.016) |   n = 25 |         |
| DMSP\(_{t}\) (nmol L⁻¹)   |          | 0.54 (0.011) |         | 0.59 (0.014) |   n = 17 |
| DMSP\(_{p}\) (nmol L⁻¹)   |          | 0.56 (0.007) |         | 0.53 (0.032) |   n = 17 |
| CDOM (ppb)               | 0.48 (0.041) |   n = 20 |         |
| HMW reducing sugars (µg L⁻¹) | 0.67 (0.011) |   n = 14 |         |
| **Negative correlations** |          |         |         |
| Chl \( \alpha \)/backscatter 660 | −0.47 (0.019) |   n = 25 |         |
| Mixed layer depth (m)    | −0.66 (0.0005) |   n = 25 |         |
| Dissolved oxygen (µmol kg⁻¹) | −0.45 (0.030) |   n = 24 |         |
| Phosphate (µmol L⁻¹)      | −0.54 (0.006) |   n = 25 |         |
| Nitrate (µmol L⁻¹)        | −0.60 (0.002) |   n = 25 |         |
| Silicate (µmol L⁻¹)       | −0.50 (0.012) |   n = 25 | −0.43 (0.031) |   n = 26 |
| Monounsaturated fatty acids (µg L⁻¹) | −0.82 (0.007) |   n = 10 |         |

flux to the atmosphere appears to be highly variable as is the proportion of S emitted as MeSH compared to DMS. For example, MeSH fluxes in the Kettle et al. (2001) study varied by several orders of magnitude, and in some cases the MeSH flux equaled the DMS flux. Similarly, DMS\(_{sw}\)/MeSH\(_{sw}\) concentration ratios have varied substantially (Kettle et al., 2001; Leck and Rodhe, 1991; Kiene et al., 2017). As such, further studies are needed to investigate the spatial distribution of MeSH both in seawater and the atmosphere as well as the importance of MeSH as a source of atmospheric sulfur. The fate of atmospheric MeSH sulfur in the atmosphere is also highly uncertain, in terms of its degradation pathways, reactions, and intermediate and final degradation products. For example, the impact that oxidation of MeSH has on the...
oxidative capacity of the MBL, and on other processes such as particle formation or growth, to the best of our knowledge remains largely unknown, and further work is needed on its atmospheric processes and fate.

A correlation analysis of MeSH$_a$ and biogeochemical parameters was undertaken for the first time, and it showed that MeSH$_a$, as well as DMS$_a$, correlated with their ocean precursor, DMSP, and also correlated with seawater DMS (DMS$_{sw}$). The correlation of MeSH$_a$ with DMS$_{sw}$ is likely due to a common ocean precursor of both gases (DMSP), which is produced via different pathways.

Correlation of acetone$_a$ with biogeochemical parameters suggests a source of acetone from warmer subtropical ocean waters, in line with other studies, with positive correlations between acetone$_a$ and ocean temperature, high-molecular-weight sugars, cryptophyte, eukaryote phytoplankton, chromophoric dissolved organic matter (CDOM), and particle backscatter, and a negative correlation with nutrients. While data with a terrestrial source influence were removed from this analysis, it is still possible that the acetone peaks observed may not have been due to a positive flux of acetone from the ocean but rather from in situ processes, leading to acetone production such as oxidation of marine-derived organic aerosol.

Finally, the SOAP voyage provided the opportunity to compare three independently calibrated DMS$_a$ measurement techniques at sea (PTR-MS, mesoCIMS and GC-SCD). Agreement between the three techniques was generally good; however, some systematic differences between the datasets were observed. Some of these differences were attributed to the near-surface DMS gradient and the use of different inlet heights (28 and 12 m a.s.l. for the PTR-MS and mesoCIMS, respectively), as well as differing approaches of integrated versus discrete measurements. The remaining discrepancies are likely due to differences in calibration scales, suggesting that further investigation of the stability and/or absolute calibration of DMS standards used at sea is warranted.

**Data availability.** DMS, acetone and MeSH data are available via the CSIRO data access portal (DAP) at https://doi.org/10.25919/5d914b00e5759 (Lawson, 2019). Further data are available by emailing the corresponding author or the voyage leader: cliff.law@niwa.co.nz.

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