The sensitivity of Southern Ocean atmospheric dimethyl sulfide (DMS) to modeled oceanic DMS concentrations and emissions

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Abstract. The biogeochemical formation of dimethyl sulfide (DMS) from the Southern Ocean is complex, dynamic, and driven by physical, chemical, and biological processes. Such processes, produced by marine biogenic activity, are the dominant source of sulfate aerosol over the Southern Ocean. Using an atmosphere-only configuration of the United Kingdom Earth System Model (UKESM1-AMIP), we performed eight 10-year simulations for the recent past (2009–2018) during austral summer. We tested the sensitivity of atmospheric DMS to four oceanic DMS datasets and three DMS transfer velocity parameterizations. One oceanic DMS dataset was developed here from satellite chlorophyll $a$. We find that the choice of oceanic DMS dataset has a larger influence on atmospheric DMS than the choice of DMS transfer velocity. Simulations with linear transfer velocity parameterizations show a more accurate representation of atmospheric DMS concentration than those using quadratic relationships. This work highlights that the oceanic DMS and DMS transfer velocity parameterizations currently used in climate models are poorly constrained for the Southern Ocean region. Simulations using oceanic DMS derived from satellite chlorophyll $a$ data, and when combined with a recently developed linear transfer velocity parameterization for DMS, show better spatial variability than the UKESM1 configuration. We also demonstrate that capturing large-scale spatial variability can be more important than large-scale interannual variability. We recommend that models use a DMS transfer velocity parameterization that was developed specifically for DMS and improvements to oceanic DMS spatial variability. Such improvements may provide a more accurate process-based representation of oceanic and atmospheric DMS, and therefore sulfate aerosol, in the Southern Ocean region.


1 Introduction

The representation of aerosols over the Southern Ocean (40 to 60°S) is a large source of uncertainty in climate models due to the lack of observational data and large seasonal variability (Revell et al., 2019). Poor representation of aerosols contributes to the large biases in future climate projections over the Southern Ocean (Myhre et al., 2014). Sea spray and dimethyl sulfide (DMS; CH$_3$SCH$_3$) are fundamental sources for aerosol formation over this region (Revell et al., 2021; Bhatti et al., 2022). The dominant source of sulfate over the marine atmosphere is the biogenic marine aerosol precursor DMS (Keller et al., 1989; Bates et al., 1987; Kiene and Bates, 1990; Curson et al., 2011). Revell et al. (2019) found that sulfate aerosol production from DMS was responsible for around 60% of the austral summer aerosol optical depth over the Southern Ocean. Atmospheric DMS therefore has the potential to greatly influence cloud condensation nuclei during austral summer (Kloster et al., 2006; Revell et al., 2019; Korhonen et al., 2008; Pandis et al., 1994).

The Southern Ocean contains extremely high phytoplankton and marine biota productivity during austral summer (DJF or December–February) (Deppeler and Davidson, 2017). Marine biogenic activity, controlled by marine biota, plays a key role in chlorophyll (chl) a production and is considered to be a key driver of oceanic DMS production (e.g., Uhlig et al., 2019; Townsend and Keller, 1996; Anderson et al., 2001; Deppeler and Davidson, 2017). Earth system models (ESMs) represent the process of oceanic DMS formation through multiple approaches that are dependent on chl a, nutrients, light, mixed-layer depth, zooplankton, and dimethylsulfoniopropionate concentration (Bock et al., 2021). The UKESM1 and MIROC-ES2L models use a diagnostic approach to represent chl a (Sellar et al., 2019; Anderson et al., 2001; Hajima et al., 2020). The CNRM-ESM2-1 and NorESM2-LM models use a prognostic approach, closely related to zooplankton and dimethylsulfoniopropionate abundance, which is a precursor of oceanic DMS (Seland et al., 2020; Séférian et al., 2019). Bock et al. (2021) evaluated oceanic DMS in the Coupled Model Intercomparison Project Phase 6 (CMIP6) and found that all models are biased in comparison with observational climatologies of DMS in the Southern Ocean region.

Atmosphere-only global climate models use climatologies to prescribe the global concentration of oceanic DMS in the surface seawater layer. Lana et al. (2011), Kettle et al. (1999), and Hulswar et al. (2022) constructed observational climatologies of oceanic DMS, which are used by such models. However, there is a limited amount of data available within the Southern Ocean, which can lead to errors in the representation of oceanic DMS (e.g., Bock et al., 2021; Mulcahy et al., 2020). A limitation of representing oceanic DMS as a static climatology is that it does not account for the large temporal variations in DMS concentrations observed. For instance, El Niño–Southern Oscillation (ENSO) events, wildfires, and volcanic eruptions all significantly influence oceanic DMS within the Southern Ocean (e.g., Yoder and Kennelly, 2003; Tang et al., 2021; Wang et al., 2022; Browning et al., 2015; Longman et al., 2022). Calculating oceanic DMS online using a biological proxy would resolve these perturbing events to some degree (Galli et al., 2018).

The flux of DMS from the ocean to the atmosphere depends on the gas transfer velocity ($K$), which in turn depends on the surface wind speed (e.g., Fairall et al., 2011). The flux of DMS is calculated as

$$\text{DMS}_{\text{flux}} = K \times \Delta C = K (\text{DMS}_{\text{w}} - \text{DMS}_{\text{a}}).$$

(1)

$\Delta C$ represents the concentration gradient across the air–sea interface, where DMS$_{\text{w}}$ is the concentration of DMS in water, and DMS$_{\text{a}}$ is the concentration in the air but is negligible, as this concentration is substantially smaller than that of the oceanic concentration.

Many DMS transfer velocity parameterizations have been developed, but most use transfer velocities measured for gases other than DMS (Wanninkhof, 1992, 2014; Nightingale et al., 2000; Liss and Merlivat, 1986). Some studies, including Blomquist et al. (2017) and Yang et al. (2011), used DMS measurements to derive a relationship between wind speed and DMS. Depending on the solubility of the gas measured, gas transfer velocities typically have a linear or quadratic dependence on wind speed. Linear relationships best represent gases with intermediate solubilities, such as DMS (e.g., Blomquist et al., 2017; Goddijn-Murphy et al., 2016; Bell et al., 2015; Yang et al., 2011; Huebert et al., 2010), while quadratic equations are better suited for highly soluble gases like CO$_2$ (Wanninkhof, 2014; Nightingale et al., 2000; Wanninkhof, 1992).

Uncertainty in DMS emissions remains high, particularly in the Southern Ocean region, where wind speeds are high and observational data sparse (e.g., Elliott, 2009; Smith et al., 2018; Zhang et al., 2020). ESMs use a variety of transfer velocities to represent DMS emissions (Bock et al., 2021). UKESM1 uses the Liss and Merlivat (1986) parameterization, even though it was constructed for gases other than DMS.

Here we examine whether incorporating realistic oceanic DMS variability based on remotely sensed chl a observations improves the simulation of atmospheric DMS. Using a nudged configuration of the atmosphere-only United Kingdom Earth System Model (UKESM1-AMIP), we use three established oceanic DMS datasets and three transfer velocity parameterizations. We also test a 10-year monthly time series in which oceanic DMS is calculated offline from MODIS Aqua satellite chl a data, using the Anderson et al. (2001) oceanic DMS parameterization, which is used by UKESM1 (Sellar et al., 2019). We evaluate sea-to-air fluxes of DMS and oceanic and atmospheric DMS concentrations relative to station and ship-based observations. The observational datasets are described in Sect. 2.4, the model configuration is described in Sect. 2.1, and details of the oceanic DMS
datasets and transfer velocity parameterizations tested are in Sect. 2.2 and 2.3, respectively. Results follow in Sect. 3.

2 Methods

2.1 Model configuration and evaluation

Simulations were performed using the atmosphere-only configuration of the coupled UK Earth System Model (UKESM1; Yool et al., 2020; Sellar et al., 2019; Mulcahy et al., 2020). By default, atmospheric DMS is produced via the Lana et al. (2011) oceanic DMS dataset and Liss and Merlivat (1986) transfer velocity parameterization. Atmospheric DMS then oxidizes to form sulfate aerosols. In UKESM1, aerosol growth, chemistry and removal are handled by the GLOMAP-mode scheme (Mulcahy et al., 2020).

Wind and temperatures are nudged to 6 h ERA5 reanalysis data (Hersbach et al., 2020). The full description of the nudging configuration is outlined in Telford et al. (2008). Nudging ensures that wind speeds, which are pivotal to the formation of atmospheric DMS, are accurately represented (Pithan et al., 2012; Kuma et al., 2020) and allows like-for-like comparisons against observations. Sea surface temperature and sea ice data from the Hadley Centre Global Sea Ice and Sea Surface Temperature were used (HadISST; Titchener and Rayner, 2014). Simulations are 10 years long, spanning from 2009 to 2018. This period was chosen to coincide with the availability of recent DMS observations (Sect. 2.4).

Atmospheric DMS concentrations are analyzed at the lowest model level, at 20 m during DJF, which is the most productive season for DMS (Deppeler and Davidson, 2017; Jarníková and Tortell, 2016). Hourly output was saved to compare with observations where applicable (for example, voyages provide observations at hourly temporal frequency). To evaluate the variability, we use the coefficient of variation (CoV). A higher CoV suggests that the variability or dispersion of the data is relatively large compared to its mean. Where uncertainty is reported, 1 standard deviation calculated over the relevant domain and time period is stated.

2.2 Oceanic DMS

We input four oceanic DMS datasets into the model, namely three climatologies and one 10-year time series. Observational-based climatologies are from Lana et al. (2011) (hereafter “Lana”) and Hulswar et al. (2022) (“Hulswar”). The “MEDUSA” climatology (1979–2014) originates from the UKESM1 CMIP6 (Yool et al., 2021; Sellar et al., 2019; Tang et al., 2019). Table 1 outlines the oceanic DMS datasets used. Ocean biogeochemistry is simulated in the UKESM1 via MEDUSA2.0 Yool et al. (the Model of Ecosystem Dynamics, nutrient Utilization, Sequestration, and Acidification; 2020, 2013). The time series was calculated offline, using a combination of satellite data and the UKESM1 approach to calculating oceanic DMS, as described below.

In UKESM1, oceanic DMS concentrations are calculated using a diagnostic method from Anderson et al. (2001), using surface daily shortwave radiation (J), dissolved inorganic nitrogen (Q), and chl a (C) as follows:

\[ \text{Oceanic DMS} = a, \text{ for } \log(CJQ) \leq s \]  

(2)

\[ \text{Oceanic DMS} = b(\log(CJQ) - s) + 1, \text{ for } \log(CJQ) > s. \]  

(3)

The parameter values are \( a = 1, b = 8, \) and \( s = 1.56 \), as described by Sellar et al. (2019). \( Q, \) chl \( a, \) and \( J \) are averaged from CMIP6 for the MEDUSA climatology. The Anderson et al. (2001) parameterization produces positive biases in DMS over the Southern Ocean within MEDUSA (Bock et al., 2021), due to the set minimum oceanic concentration of 1, which leads to large average DMS concentrations (Yool et al., 2021; Bock et al., 2021). Recent research suggests that chl \( a \) may not be an appropriate proxy for oceanic DMS (Uhlig et al., 2019; Bell et al., 2021), and future work will explore alternative methods for calculating oceanic DMS within UKESM1. Nonetheless, chl \( a \) is widely used by CMIP6-era models to calculate oceanic DMS, and we explore here whether using an observationally derived chl \( a \) concentration field leads to changes in the spatial and temporal variability in the atmospheric DMS. Monthly mean chl \( a \) concentrations from the Moderate Resolution Imaging Spectroradiometer (MODIS) Aqua satellite instrument were used to construct a time series of oceanic DMS between 2009 and 2018 (Table 1; Hu et al., 2019; O’Reilly and Werdell, 2019). This time series, which we term the “MODIS-DMS” dataset, is calculated offline, using the same diagnostic parameterization as Eqs. (2) and (3). The \( J \) and \( Q \) values used to calculate MODIS-DMS remain the same as MEDUSA. Through this, we capture spatial and interannual chl \( a \) variability, indicating biological productivity. Bilinear interpolation is used to fill in small gaps (around 1% for monthly averages) of spatial chl \( a \) data. Oceanic DMS concentrations are masked where they coincide within the sea ice zone from HadISST.

In general, the MODIS Aqua Ocean Color chl \( a \) retrieval underestimates Southern Ocean chlorophyll concentrations by up to 25% (Zeng et al., 2016; Haëntjens et al., 2017; Jena, 2017; Gregg and Casey, 2007; Johnson et al., 2013). Simulated oceanic DMS may therefore be systematically underestimated. Nonetheless, the high spatial and temporal availability of chl \( a \) observations during summertime makes it useful to explore spatiotemporal variability in atmospheric DMS.

2.3 DMS sea-to-air flux

Three DMS transfer velocities are tested (Fig. 1, Table 2). Two are linear equations from Liss and Merlivat (1986)
(hereafter “LM86”) and Blomquist et al. (2017) (hereafter “B17”). LM86 is a piece-wise linear equation and the default parameterization within UKESM1 (Sellar et al., 2019) and was evaluated in combination with all oceanic DMS datasets. The quadratic formula from Wanninkhof (2014) (hereafter “W14”) is also tested. Using these different parameterizations provides an appropriate estimate for the spread of DMS emissions due to the upper and lower limits of DMS transfer velocity tested from in situ DMS measurements (e.g., Goddijn-Murphy et al., 2016; Blomquist et al., 2017). Table 2 summarizes the sensitivity simulation names of simulations that were performed. Simulations are named with the oceanic DMS concentration used and subscripted with the transfer velocity used. For example, Lana\textsubscript{LM86} means that the simulation used the Lana et al. (2011) climatology as its oceanic DMS dataset and the DMS transfer velocity parameterization of Liss and Merlivat (1986).

The Schmidt number for DMS is used to calculate the DMS emission. The Schmidt number represents the viscosity or diffusion properties of a gas and varies with respect to sea surface temperature ($T$ in °C). We update the Schmidt number of DMS ($\text{Sc}_{\text{DMS}}$) used in the UKESM1 from the formulation used in Saltzman et al. (1993) to Wanninkhof (2014), as shown in Eq. (3):

$$\text{Sc}_{\text{DMS}} = 2855.7 + (-177.63 + 6.0438 + (-0.11645 + 0.00094743 \cdot T) \cdot T) \cdot T,$$

where $T$ is derived from HadISST (Titchner and Rayner, 2014). $U_{10}$ (m s\(^{-1}\)) represents near-surface (10 m) wind speed, and $K_w$ (cm h\(^{-1}\)) represents the transfer velocity of DMS. Equation (5) represents the LM86 transfer velocity of DMS as follows:

For $u_{10} \leq 3.6$:

$$K_w = 0.17 \left( \frac{600}{\text{Sc}_{\text{DMS}}} \right)^{\frac{3}{2}} u_{10}, \text{for } 3.6 \leq u_{10} < 13 :$$

$$K_w = (2.85 u_{10} - 9.65) \left( \frac{600}{\text{Sc}_{\text{DMS}}} \right)^{\frac{1}{2}}, \text{for } u_{10} > 13 :$$

$$K_w = (5.8 u_{10} - 49.3) \left( \frac{600}{\text{Sc}_{\text{DMS}}} \right)^{\frac{1}{2}}.$$

W14 uses a quadratic formula (Eq. 6) for sea-to-air transfer. W14 is also used to calculate DMS emissions amongst

$$K_w = 0.251 \cdot u_{10}^2 \left( \frac{660}{\text{Sc}_{\text{DMS}}} \right)^{\frac{1}{2}} \quad (6)$$

B17 is the only parameterization tested in this study for which the transfer velocity is based on real-world observation of DMS (Eq. 7). B17 is a super-linear parameterization; however, for simplicity and the wind speeds used in this study, we label B17 as a linear parameterization.

$$K_w = 0.7432 \cdot u_{10}^{1.33} \left( \frac{660}{\text{Sc}_{\text{DMS}}} \right)^{\frac{1}{2}} \quad (7)$$

To assess the interannual variability in the DMS emissions and atmospheric DMS concentrations, we performed an additional 10-year simulation, MODIS\textsubscript{B17}CLIM. While MODIS\textsubscript{B17} used a 10-year time series of oceanic DMS derived from MODIS chlorophyll $a$ data, MODIS\textsubscript{B17}CLIM used a climatology calculated from monthly mean data for the 10-year MODIS\textsubscript{B17} time series.
Table 2. Simulations used in this study, named with the oceanic DMS concentration used and subscripted with the transfer velocity used. N/A stands for not available.

<table>
<thead>
<tr>
<th>Simulation name</th>
<th>Oceanic DMS</th>
<th>DMS transfer velocity parameterization</th>
</tr>
</thead>
<tbody>
<tr>
<td>LanaLM86</td>
<td>Lana et al. (2011)</td>
<td>Liss and Merlivat (1986)</td>
</tr>
<tr>
<td>Lanag17</td>
<td>Lana et al. (2011)</td>
<td>Blomquist et al. (2017)</td>
</tr>
<tr>
<td>HulswarLM86</td>
<td>Hulswar et al. (2022)</td>
<td>Liss and Merlivat (1986)</td>
</tr>
<tr>
<td>MEDUSA_LM86</td>
<td>Anderson et al. (2001), Sellar et al. (2019)</td>
<td>Liss and Merlivat (1986)</td>
</tr>
<tr>
<td>MODISB17LM86</td>
<td>N/A (produced for this study)</td>
<td>Liss and Merlivat (1986)</td>
</tr>
<tr>
<td>MODISB17W14</td>
<td>N/A (produced for this study)</td>
<td>Blomquist et al. (2017)</td>
</tr>
<tr>
<td>MODISB17CLIM</td>
<td>N/A (climatology produced for this study)</td>
<td>Wanninkhof (2014)</td>
</tr>
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2.4 Observational datasets

2.4.1 Oceanic DMS and atmospheric DMS datasets

Two Southern Ocean voyages are used to evaluate our simulations, namely the SOAP campaign (Surface Ocean Aerosol Production; Bell et al., 2015; Law et al., 2017) and research vessel (R/V) Tangaroa voyage (TAN1802; Kremser et al., 2021). The SOAP voyage measured oceanic and atmospheric DMS from February–March 2012 near the Chatham Rise (within 42–47°S, 172–180°E) off the east coast of Aotearoa / New Zealand (Bell et al., 2015; Smith et al., 2018). The TAN1802 voyage measured oceanic DMS along the Southern Ocean during February–March 2018 between 40 to 70°S, 180°E (Kremser et al., 2021). We also extend the simulations to cover the ANDREXII voyage between February–April 2019 for atmospheric DMS concentrations, as this voyage mostly conducted measurements during autumn (Wohl et al., 2020). ANDREXII traveled longitudinally around 60°S. Although outside our simulation range, we also consider the Southern Ocean Iron ReRelease Experiment (SOIREE) for atmospheric DMS analysis from February 1999 (Boyd and Law, 2001) between 42–63°S, 139–172°E. We used oceanic DMS measurements for TAN1802 Kremser et al. (2021), SOAP (Bell et al., 2015), and ERA5 surface wind speeds (Hersbach et al., 2020) to calculate hourly DMS emissions. The Wanninkhof (2014) DMS Schmidt number is calculated using the same parameters used within the simulations to ensure consistency with the comparisons to simulated fluxes. The HadISST and ERA5 wind speed data were obtained for the same time and location for both of the two voyages (within the nearest-neighbor grid cell). We applied three different transfer velocity parameterizations (LM86, B17, and W14) to both SOAP and TAN1802 voyage paths (see Sect. 3.2).

We compare our simulations to the voyage dataset using the hourly model output and identify the nearest-neighbor grid cell to the ship location. Analysis of oceanic DMS data used in the models is also synchronized to TAN1802 and SOAP voyages, using the same timescales for comparing the voyages with model data.

We also validate the model using atmospheric DMS concentrations measured at two stations, namely Kennaook / Cape Grim (1989 to 1996; 41°S and 145°E) and King Sejong Station (2018 to 2020; 62°S, 58°W). King Sejong is located on the Antarctic Peninsula, where sea ice melt occurs during our study period, which can profoundly increase DMS emissions, as previously found by Berresheim et al. (1998) and Read et al. (2008).

2.4.2 Cloud and aerosol observations

MODIS Aqua aerosol optical depth (AOD) measurements at 550 nm (Platnick et al., 2017) are compared with each daily mean model output. Daily averaged observations from Grosvenor et al. (2018) and Bennartz and Rausch (2017) were used to compare the cloud droplet number concentration (CDNC) with our daily averaged simulations. Finally, to evaluate cloud condensation nuclei (CCN), we used observations from Choudhury and Tesche (2023) at 818 m in comparison with simulated CCN at 800 m. The description and evaluation of using MODIS-observed AOD compared with a related configuration of UKESM1-AMIP is discussed in more detail in Revell et al. (2019) and Mulcahy et al. (2020). We calculate an austral summertime climatology for these observational datasets, which we use over the Southern Ocean.

3 Results and discussion

3.1 Oceanic DMS

Figure 2a–d shows the spatial distribution of each oceanic DMS dataset. Each distribution has key defining characteristics, although Hulswar (Fig. 2d) is an update to Lana (Fig. 2c). The distinction between MODIS-DMS and MEDUSA oceanic DMS calculations is chl a, which results in distinctly different distributions, as shown in Fig. 2e. Observationally based climatologies, like Lana or Hulswar, do
Figure 2. Summertime (DJF) Oceanic DMS in the Southern Ocean (40–60 S). The spatial distribution (a–d) shows the (a) UKESM1 climatology from MEDUSA, (b) the climatology from MODIS-DMS, and observationally based climatologies of (c) Lana and (d) Hulswar. (e) The box plot shows the distribution of each oceanic DMS dataset used, where MODIS-DMS contains all 10 years of data, while the climatologies contain 12 months.

not align with the chl $a$ distribution in the Southern Ocean, particularly along the Antarctic Circumpolar Current, concentrating oceanic DMS in specific regions based only on observations of oceanic DMS (Lana et al., 2011; Hulswar et al., 2022). The mean difference between the lowest (MODIS-DMS) and highest (MEDUSA) mean of all the oceanic DMS datasets used is 107%.

MEDUSA produces the most homogeneous oceanic DMS distribution in the summertime Southern Ocean, with the highest mean and smallest standard deviation (4.88 ± 0.87 nM). It also has the lowest CoV of ± 17%, indicating a small spread of variance. Chl $a$ in MEDUSA shows a positive bias against summer observations in the Southern Ocean (Yool et al., 2013, 2021). In contrast, MODIS-DMS has low oceanic DMS concentrations in open-ocean regions and high concentrations in biologically productive regions (near the subtropical front), such as the Chatham Rise and coastal South America (Behrens and Bostock, 2023). MODIS-DMS has the largest spatial variability in oceanic DMS overall (CoV 67%). The mean oceanic DMS in MODIS-DMS is 2.36 ± 1.57 nM, which is outside the range of MEDUSA, highlighting the sensitivity of the Anderson et al. (2001) parameterization to chl $a$ concentrations.

In the MODIS-DMS simulation, oceanic DMS concentrations vary each summer across the Southern Ocean over a 10-year climatology (see Fig. S1a in the Supplement). The most significant interannual variability occurs around Aotearoa / New Zealand and South America’s east coast, likely from phytoplankton blooms influenced by ENSO (Santoso et al., 2017; Thompson et al., 2015; Yoder and Kennelly, 2003; Fig. 2a). The Lana and Hulswar simulations have similar means (3.87 and 3.51 nM, respectively) but differ in their distribution (Fig. 2e). Oceanic DMS maximizes at 30 nM in Lana and at 14 nM in Hulswar. The MEDUSA simulation using the Anderson et al. (2001) parameterization shows oceanic DMS maximizing at 11 nM, while when a variable chl $a$ concentration field is used in the MODIS-DMS simulation, oceanic DMS maximizes at 18 nM (64% higher than in the MEDUSA simulation).

To examine how the simulations compare with observations, we compare the oceanic DMS distribution against TAN1802 and SOAP voyages for the regions and times at which those voyages took place (Figs. 3, 4). For the TAN1802 voyage (40–70° S, 180° E), the distribution of measured oceanic DMS aligns closely with the Lana simulation. MODIS-DMS and MEDUSA have lower means of 1.19 and 1.52 nM, respectively, but MODIS-DMS has a high CoV of 79%, due to higher concentrations at lower latitudes (45° S) of the Southern Ocean. Oceanic DMS in the Hulswar simulation overestimates DMS concentrations by a factor of 2 between 45–65° S.

For the SOAP voyage, which targeted phytoplankton bloom events (42–47° S, 172–180° E), the measured DMS distribution is skewed toward higher concentrations com-
Figure 3. Violin plots of TAN1802 data (gray). Overlaid are the oceanic DMS datasets used in the model simulations (February to March 2018; 40 to 70°S, 180°E) from MEDUSA (purple), MODIS-DMS (blue), Lana (green), and Hulswar (yellow). Violin plots depict data distribution and density. The width of each “violin” corresponds to the frequency of data points within that value range, while the length indicates the range of values. The frequency axis, represented by the width, allows for an immediate visual comparison of how often particular ranges of values occur in each category. This offers a comprehensive view of both the distribution and frequency of data across different categories.

pared with the TAN1802 voyage (Fig. 4). In contrast, TAN1802 transected the Southern Ocean without a specific focus on bloom activity, yielding a range of DMS concentrations. We consider that SOAP is still useful, as it offers insights into extreme conditions not reflected in other datasets. All simulations fail to capture the higher concentrations measured by SOAP. Oceanic DMS in the MODIS-DMS exhibits the highest variability (CoV of 36%), mean, and maximum concentration. MODIS-DMS also aligns best with SOAP in that it captures some of the high DMS concentrations resulting from phytoplankton blooms. The MODIS-DMS simulation captures around half of the variability in the SOAP measurements, whereas the other simulations only match between 7% to 18%. MODIS-DMS is within 11% of the SOAP mean, whereas the other simulations are 22% to 218% lower. See Figs. S2 and S3 for simulated comparisons of DMS emission to SOAP and TAN1802.

The Anderson et al. (2001) parameterization assumes chl $a$ is central to oceanic DMS formation. Previous correlations between chl $a$ and oceanic DMS, given by the coefficient of determination ($R^2$), range globally from 0.11 to 0.93, with higher latitudes having increased $R^2$ values due to factors like nutrient availability and prolonged summer daylight, coupled with heightened wind speeds (Uhlig et al., 2019; Townsend and Keller, 1996; Tison et al., 2010; Matrai et al., 1993). Gros et al. (2023) estimated an $R^2$ of 0.93 towards sea ice latitudes, while Bell et al. (2021) found chl $a$ explains just 15% of oceanic DMS variability. Using the Anderson et al. (2001) parameterization in MODIS-DMS, we determined a large $R^2$ of 0.75 in the Southern Ocean. While associating chl $a$ with oceanic DMS has discrepancies (Gros et al., 2023; Bell et al., 2021), we show that using Anderson et al. (2001) with satellite chl $a$ data better represents Southern Ocean summertime DMS compared with the MEDUSA configuration.

Chl $a$ is used to calculate oceanic DMS in two of the four ESMs with interactive biogeochemistry in CMIP6 (Bock et al., 2021). These models reveal discrepancies between each other and observed oceanic DMS datasets, indicating ongoing uncertainties in CMIP6 ESMs concerning oceanic DMS and its flux to the atmosphere (Bock et al., 2021). Bock et al. (2021) emphasize the need for enhanced understanding and observations to accurately capture DMS–climate feedbacks. CNRM-ESM2-1 adopts an approach considering zooplankton and dimethylsulfoniopropionate (DMSP) rather than chl $a$, but its validation is challenging due to limited observational data (Belviso et al., 2012). NorESM2 uses an alternative mechanism for DMS production, by using detritus export production and sea surface temperature (Tjiputra et al., 2020). An oceanic DMS algorithm developed by Gali et al. (2018) includes sea surface temperature, chl $a$, photosynthetically active radiation, and the mixed-layer depth, but oceanic DMS has a general overestimation along coastal regions (Gali et al., 2019; Hayashida et al., 2020). Gali et al. (2018) also produced a time series of oceanic DMS over parts of the Northern Hemisphere, finding high interannual variability by using chl $a$ satellite data. Adopting temporally variable oceanic DMS inputs within the model may
better reflect interannual Southern Ocean variability due to ENSO events and biologically productive years. One such way to achieve this for future projections would be through a stochastic approach of capturing all chl α years from the satellite (e.g., SeaWiFS and MODIS Aqua) archive.

3.2 DMS flux

Having established that oceanic DMS from the MODIS-DMS simulation compares reasonably with summertime observational voyages, as seen in Figs. 3 and 4, we now assess the sensitivity of atmospheric DMS to various sea-to-air transfer functions (Figs. 5, S4). Figure 5 shows the DMS flux during the austral summer in the Southern Ocean, averaging between 2.9 and 7.3 TgS yr⁻¹. This is consistent with the Jarníková and Tortell (2016) estimation of 3.4 TgS, aligning most with the MODIS-DMS linear parameterizations (LM86 and B17). The spread in average Southern Ocean summertime DMS fluxes across the eight simulations is 153 %, which is greater than the spread between all the simulations testing different oceanic DMS datasets, which is at 107 %. The lowest CoVs within both oceanic DMS and DMS emissions are found in the MODIS-DMS simulations and, specifically, the Blomquist et al. (2017) parameterization (MODIS-B17) with a mean of 2.9 ± 0.84 TgS yr⁻¹. The upper range of simulated DMS flux, 7.3 ± 1.8 TgS yr⁻¹, comes from the W14 quadratic formula used with the Lana DMS climatology (Lana_W14).

The largest DMS emissions are seen in the MEDUSA_LM86 simulations, due to the relatively large underlying oceanic DMS values spread throughout the Southern Ocean (Fig. 2a). The Lana_W14 simulation also shows large DMS emissions due to the quadratic dependence of the gas transfer velocity on wind speed (Fig. 1). Overall, the W14 quadratic formula yields about 33 % more emissions than the LM86 and B17 linear formulas. For the transfer velocity parameterizations using a linear relationship to wind (LM86 and B17), LM86 exhibits a higher transfer velocity than B17 for wind speeds above 7.5 m s⁻¹ (Fig. 1). Given the Southern Ocean’s predominantly high wind speeds (Bracegirdle et al., 2020), simulations indicate that LM86 yields 14 % more emitted DMS than B17 (Fig. 5).

The LM86 transfer velocity parameterization was tested with all oceanic DMS datasets, as it is currently the parameterization used by default in UKESM1-AMIP. Simulations using LM86 have a spread in the average summertime Southern Ocean DMS emissions of 112 % (3.3 to 6.9 TgS yr⁻¹). In contrast, simulations using the same oceanic DMS dataset (MODIS-DMS and Lana) but including transfer velocity parameterizations (LM86, B17, and W14) have a spread in the average summertime Southern Ocean DMS emissions from 51 % (MODIS-DMS simulations) to 62 % (Lana simulations). The choice of the oceanic DMS dataset therefore impacts DMS emissions more than the transfer velocity parameterization within these simulations.

Many CMIP6 models use the quadratic transfer velocity parameterization detailed in Wanninkhof (2014) for DMS emissions (e.g., Salzmann et al., 2022; Seland et al., 2019; Neubauer et al., 2019; Tatebe and Watanabe, 2018; Wu et al., 2019). Yet, recent studies indicate a linear relationship between DMS and wind speed (e.g., Blomquist et al., 2017; Yang et al., 2011; Bell et al., 2013, 2015). We demonstrate that linear DMS transfer velocities represent the DMS flux ranges better than the quadratic W14 flux when compared to Southern Ocean observations.
3.3 Atmospheric DMS

We next evaluate atmospheric DMS in our sensitivity simulations. Figure 6 compares all simulated atmospheric DMS with the observational datasets. Data in Fig. 6 are from three Southern Ocean voyages (SOAP, SOIREE, and ANDREXII; Fig. 6a–c) and two stations (Kennaook / Cape Grim and King Sejong Station; Fig. 6d–e). Figure 6f shows the aggregate-averaged DMS concentrations from all five observational sources and has an average summertime concentration of 129 ± 74 ppt (parts per trillion) (Fig. 6f; Smith et al., 2018; Wohl et al., 2020; Boyd and Law, 2001). The simulations using the MODIS-DMS oceanic dataset and linear DMS transfer models (LM86 and B17) show the closest agreement with the observational mean of 106 ± 66 ppt and 100 ± 60 ppt for MODISL86 and MODISB17, respectively. The mean total spread in the summertime Southern Ocean atmospheric DMS across all simulations is 171 %, compared with the spread of 153 % in DMS emissions.

Our simulations, compared to coastal Antarctic measurements, offer insights into the performance of sea-ice-influenced regions (Galf et al., 2021). In summer, Berresheim et al. (1998) recorded mean atmospheric DMS of 119 ppt at 64.8° S, 64° W, closely matching MODISB17 at 121 ppt. All other oceanic DMS datasets show concentrations which are more than twice as large as this measurement. Read et al. (2008) measured atmospheric DMS concentrations of 45 ± 50 ppt at Halley VI Research Station, Antarctica (75.4° S, 26.2° W), best aligning with LanaB17 at 42 ppt. It should be noted that all simulations fall within 1 standard deviation of the measurements reported at Halley Station. Preunkert et al. (2007) measured high interannual variation in the atmospheric DMS at Dumont d’Urville Station (66.4° S, 140° E) during January, from 244 ppt in 2002 to only 60 ppt in 2003. The average January concentration over 13 years was 170 ± 180 ppt. Here, the Lana and Hulswar simulations are in closest agreement and simulate average DMS concentrations between 92 and 141 ppt. Last, Lee et al. (2010)
measured a 61 ppt average over the Pacific–Southern Ocean in February, closest to MODIS\textsubscript{B17} and MODIS\textsubscript{LM86} (64 and 53 ppt, respectively).

Multiannual studies emphasize high yearly variability (Read et al., 2008; Preunkert et al., 2007). Measurements during austral summer over the Southern Ocean show significant variability, especially in higher latitudes. The climatologies produce higher concentrations along the coastal regions of Antarctica, as illustrated in Fig. 2a–d, but MODIS-DMS still captures much of the spatial variability (Fig. S5). MEDUSA performs the worst over these higher-latitude regions, where sea ice can have a large role in producing atmospheric DMS (Gali et al., 2021). MODIS\textsubscript{B17} represents atmospheric DMS more accurately than models like MEDUSA, Lana, and Hulswar, based on observations over the Southern Ocean during summertime.

### 3.4 Effects from interannual and spatial variability

To assess the impact of interannual variability in oceanic DMS on simulated atmospheric DMS, we compare the MODIS\textsubscript{B17} simulation with MODIS\textsubscript{B17}\textsubscript{CLIM}, which used a climatology of oceanic DMS calculated from the MODIS-DMS dataset (Fig. 7). Both simulations are similar ($R^2 = 0.92$) in terms of interannual variability across the Southern Ocean as a whole (Fig. 7c). Rolling means are presented in Fig. S1b and c. While there are small differences in Southern Ocean atmospheric DMS between the simulations, the overwhelming similarities between Fig. 7a and b suggest that an oceanic DMS climatology results in similar interannual variability in the atmospheric DMS probability density function (PDF), suggesting that oceanic DMS is not a strong driver of interannual variability in atmospheric DMS. This result is in contrast to that of Gali et al. (2018), who used a different algorithm for producing oceanic DMS. This difference may be due to our use of the Anderson et al. (2001) algorithm, which is known to produce limited variability (Belviso et al., 2004; Bock et al., 2021).

To assess the impact of spatial variability in oceanic DMS on simulated atmospheric DMS, we compare simulations performed using the MEDUSA and MODIS-DMS datasets (with low and high spatial variability in oceanic DMS, respectively) in Fig. 8. The larger variability in the MODIS-DMS oceanic DMS dataset leads to larger variability in simulated atmospheric DMS when compared with the MEDUSA simulations. The spatial CoV from MEDUSA\textsubscript{LM86} is 45 % lower than MODIS\textsubscript{LM86}, showing greater spatial variability from MODIS-derived chl $\alpha$. The oceanic DMS signal in the atmosphere is strong but includes large fluctuations driven by the wind variations.

### 3.5 Aerosol and cloud response

Figures 9 and S6 show the effect on cloud and aerosol properties of changing the atmospheric DMS distribution. Changing the atmospheric DMS concentration yields a spread across all our simulations for AOD, CDNC, and CCN by 6 %, 15 %, and 11 %, respectively, over the austral summer Southern Ocean. As DMS predominately oxidizes into sulfate within the smaller aerosol modes, it has a smaller influence on the AOD than the larger modes from sea salt aerosol (Mulcahy et al., 2020). However, these smaller aerosols influence cloud seeding, as our simulations show. Changes to the oceanic DMS dataset increase the spread in simulated CCN and CDNC over the Southern Ocean rather than changing the DMS emissions, which is consistent with our findings for atmospheric DMS concentrations. Altering the oceanic DMS dataset produces a 73 % greater change in AOD than altering the DMS emissions over the Southern Ocean, emphasizing the role of the ocean in producing atmospheric DMS. Box plots of AOD, CCN, and CDNC (Fig. 9e, a, c) show that the simulations do not capture the maxima in CDNC, CCN, or AOD over the Southern Ocean.

### 4 Conclusions

We examined the sensitivity of atmospheric DMS to different oceanic DMS datasets and transfer velocity parameterizations using the UKESM1-AMIP model. Modeled atmospheric DMS over the Southern Ocean is sensitive to both oceanic DMS concentrations and sea-to-air emissions. The current approach to calculating oceanic DMS within UKESM1 (MEDUSA; Anderson et al., 2001) shows little spatial variability and high average biases in the Southern Ocean region, emphasizing the need for further refinement (e.g., Bock et al., 2021; Mulcahy et al., 2020; Yool et al., 2021). Incorporating satellite chlorophyll $\alpha$ observations within the Anderson et al. (2001) oceanic DMS parameterization produces larger spatial variability than MEDUSA.
Figure 8. Atmospheric DMS concentrations comparing (a–d) MEDUSA_LM86 with (e–h) MODIS_LM86 across 4 of the same summertime days (15 December) in (a, e) 2009, (b, f) 2010, (c, g) 2013, and (d, h) 2015. The area-weighted Southern Ocean mean is shown below each plot.

Figure 9. Summertime climatology between 60° to 40° S showing the (a, b) cloud droplet number concentrations, (c, d) cloud conversation nuclei (800 m in altitude), and (e, f) aerosol optical depth at 550 nm. The violin plots (a, c, e) represent all spatial and temporal data points across the 10 years over the Southern Ocean in December–February (DJF). The lowest 1% of values are excluded from the violin plots. (b, d, f) The gray lines represent observational datasets, where (b) Grosvenor et al. (2018) (dashed) and Bennartz and Rausch (2017) (solid) are shown for CDNC, (d) Choudhury and Tesche (2023) is shown at 818 m, (f) AOD climatology by the MODIS satellite retrieval is shown (Platnick et al., 2017). The error bars represent 1 standard deviation on either side of the observational mean.

MODIS-DMS simulations indicate that large open-water areas in the Southern Ocean have lower oceanic DMS concentrations than MODIS-DMS, including along coastal regions and at higher latitudes.

Current oceanic DMS climatologies in climate models lack similar spatial distribution to ocean chlorophyll a during Southern Ocean summer and perform poorly relative to observations from voyages and atmospheric DMS comparisons. We show how using chlorophyll a data from the MODIS Aqua satellites offers an alternative spatial representation of oceanic DMS based on the chlorophyll a distribution. Approaches like this, and that of Galí et al. (2018), offer promising avenues for realistically capturing spatial variability in oceanic DMS associated with marine biogenic activity.

During the austral summer over the Southern Ocean, the Wanninkhof (2014) quadratic DMS parameterization leads to 33% more DMS emissions than the Liss and Merlivat (1986) and Blomquist et al. (2017) parameterizations. Linear transfer velocity parameterizations also align better with observations for DMS emissions, particularly for the MODIS-DMS simulations.

Atmospheric DMS in the 10-year MODIS-DMS time series simulation shows similar interannual variability to the climatology simulation. We show that capturing large-scale spatial variability is more important for oceanic DMS concentrations than capturing large-scale interannual variability.

In simulations with different oceanic DMS datasets but the same transfer velocity parameterization, the Southern Ocean summertime DMS emissions vary by 112% (3.3 to 6.9 TgS yr⁻¹). This is approximately twice as much as the simulations using the same oceanic DMS dataset, but different transfer velocity parameterizations, in which DMS emissions vary by 50%–60% (2.9 to 4.7 TgS yr⁻¹). The choice of oceanic DMS dataset therefore has a larger influence on...
atmospheric DMS than the choice of DMS transfer velocity. The total spread in average Southern Ocean DMS emissions across all simulations is 153%. Both oceanic DMS and DMS transfer velocity parameterization changes significantly influence atmospheric DMS, emphasizing the need for careful consideration in future research. Changing the oceanic DMS concentrations and transfer velocity results in a mean spread between the simulations of 6% for AOD and 15% for CDNC.

Future work will adopt more recent parameterizations of oceanic DMS concentrations and test various sulfate chemistry schemes. In future, we recommend that models use up-to-date transfer velocity parameterizations specific to DMS, such as Blomquist et al. (2017).

Data availability. CDNC observations are available from Grosvenor et al. (2018) and Gryspeerdt et al. (2022) (https://doi.org/10.5194/amt-15-3875-2022). MODIS AOD and chlorophyll a observations were accessed via the Giovanni online data system, which is developed and maintained by the NASA Goddard Earth Sciences Data and Information Services Center (GES DISC: https://giovanni.gsfc.nasa.gov/). Platnick et al., 2015). DMS measurements from Amsterdam Island were obtained from the World Data Centre for Greenhouse Gases at https://ebas-data.nilu.no/ (EBAS, 2023). TAN1802 measurements are available from Kremser et al. (2021). DMS measurements from the SOIREE campaign are available from Boyd (2009) (http://lod.bco-dmo.org/id/dataset/3212). For the ANDREXII DMS datasets, see Wohl et al. (2020).

Model simulation data are archived at the Aotearoa / New Zealand eScience Infrastructure (https://www.nesi.org.nz/, NeSI, 2023) and are available by contacting the corresponding author.

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Author contributions. YAB implemented model developments, performed model simulations, and wrote the article, with assistance from all co-authors. LER, AJM, and AJS assisted with the experimental design and the model evaluation by comparing with the observational datasets and sensitivity analysis. ATA advised on DMS chemistry and aerosols over the Southern Ocean. CH provided assistance for lodging DMS emissions into the UKESM1 trunk. JW and EB provided technical expertise in running the model simulations.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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