



# Decadal increase in Arctic dimethylsulfide emission

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**Dimethylsulfide (DMS), a gas produced by marine microbial food webs, promotes aerosol formation in pristine atmospheres, altering cloud radiative forcing and precipitation. Recent studies suggest that DMS controls aerosol formation in the summertime Arctic atmosphere and call for an assessment of pan-Arctic DMS emission (EDMS) in a context of dramatic ecosystem changes. Using a remote sensing algorithm, we show that summertime EDMS from ice-free waters increased at a mean rate of  $13.3 \pm 6.7$  Gg S decade<sup>-1</sup> (~33% decade<sup>-1</sup>) north of 70°N between 1998 and 2016. This trend, mostly explained by the reduction in sea-ice extent, is consistent with independent atmospheric measurements showing an increasing trend of methane sulfonic acid, a DMS oxidation product. Extrapolation to an ice-free Arctic summer could imply a 2.4-fold ( $\pm 1.2$ ) increase in EDMS compared to present emission. However, unexpected regime shifts in Arctic geo- and ecosystems could result in future EDMS departure from the predicted range. Superimposed on the positive trend, EDMS shows substantial interannual changes and nonmonotonic multiyear trends, reflecting the interplay between physical forcing, ice retreat patterns, and phytoplankton productivity. Our results provide key constraints to determine whether increasing marine sulfur emissions, and resulting aerosol–cloud interactions, will moderate or accelerate Arctic warming in the context of sea-ice retreat and increasing low-level cloud cover.**

dimethylsulfide | Arctic Ocean | plankton | sea ice | aerosols

The Arctic region is warming more than 2 times faster than the global average, and ice-free summers could be a reality in the next few decades (1). Removal of the ice barrier boosts ocean–atmosphere exchanges of energy, gases, and particles, with profound effects on marine ecosystems and climate. Enhanced heat and moisture fluxes are increasing the abundance of low-level clouds (2) and, very likely, the prevalence of liquid-state clouds and precipitation (3). Ice retreat allows more solar radiation to penetrate into the ocean surface, driving a pan-Arctic increase in phytoplankton primary production (2, 4–6). Meanwhile, changes in stratification and nutrient supply to the sunlit ocean layer modulate phytoplankton productivity (4–7) and alter phytoplankton bloom phenology (7–10) and the occurrence of species with distinct biogeochemical traits (10–12) such as their capacity to produce the climate active gas dimethylsulfide (DMS) (13, 14). DMS is produced through microbial decomposition of dimethylsulfoniopropionate (DMSP), a compound synthesized in variable amounts by different phytoplankton groups (13, 14). Despite complex biogeochemical cycling (14, 15), high-latitude DMS production scales to first order with phytoplankton biomass and productivity over the seasonal cycle at large scales (16–19). In a scenario of changing ice cover and phytoplankton dynamics, changes in the magnitude, timing, and spatial distribution of Arctic DMS emission (EDMS) are expected (14, 20, 21).

Previous estimates of Arctic Ocean EDMS have been made using either sea-surface DMS climatologies (16), produced through interpolation of sparse in situ data, or prognostic models (20, 21). Both types of estimates depict the Arctic as a region with relatively low sea–air DMS flux per unit area (FDMS) on an annual basis (mean FDMS lower than  $2 \mu\text{mol m}^{-2}\text{d}^{-1}$ ) compared to temperate and tropical oceans (mean FDMS of about  $4\text{--}5 \mu\text{mol m}^{-2}\text{s}^{-1}$ ) (16, 20). However, Arctic FDMS is concentrated in the short productive

summer season, and relatively high daily fluxes have been reported associated with phytoplankton blooms that form in the wake of melting sea ice, often exceeding  $10 \mu\text{mol m}^{-2}\text{s}^{-1}$  (14, 15). Ice margin phytoplankton blooms are a major feature of the Arctic ecosystem. They typically last for 1–3 wk after ice breakup and are promptly detected using ocean color remote sensing (8, 9, 22). Given the patchy and ephemeral nature of Arctic EDMS, accurate estimates of its magnitude and spatial–temporal distribution based on climatological datasets are severely limited.

Once emitted to the atmosphere, the influence of DMS on atmospheric particles does not depend strictly on the magnitude of FDMS. Rather, it is the background concentration of aerosol particles that critically determines whether atmospheric DMS oxidation products can nucleate new particles or condense on preexisting ones (23–25). In summer, different processes isolate the Arctic marine boundary layer from southern aerosol sources (both natural and anthropogenic), namely: the northward migration of the atmospheric polar front, the efficient wet scavenging by drizzling stratocumulus clouds, and the formation of surface inversion layers (3, 25, 26). These processes result in extremely low aerosol concentrations, which favor new particle formation from local gaseous precursors (23, 25, 27, 28). Recent measurements and associated modeling have shown instances where DMS controls the formation of ultrafine particles (23, 28), which can grow large enough to act as cloud condensation nuclei (CCN) (23, 25). Thus, changes in Arctic Ocean EDMS could alter aerosol populations, light scattering, and cloud-seeding activity (26, 27, 29,

## Significance

**As Arctic sea-ice cover declines because of climate warming, the emission of reactive gases produced by marine microbes increases. One of such gases, dimethylsulfide, forms new atmospheric particles that contribute to cloud formation. This can either cool the Earth's surface by reflecting incoming sunlight, or warm it due to the blanket effect. Here we quantify Arctic Ocean dimethylsulfide emission between 1998 and 2016 using satellite observations of microalgal biomass and physical variables. We report an increasing trend, driven by sea-ice loss, and substantial year-to-year variability modulated by biological productivity. Our results can help understand the impacts and feedbacks of marine plankton on Arctic climate and foresee their future trajectories under the pressure of global change.**

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Data deposition: The DMSSAT algorithm code has been deposited in GitHub (<https://github.com/mgalii>), and curated in situ datasets and DMSSAT datasets have been deposited at Zenodo (<https://doi.org/10.5281/zenodo.3243967>).

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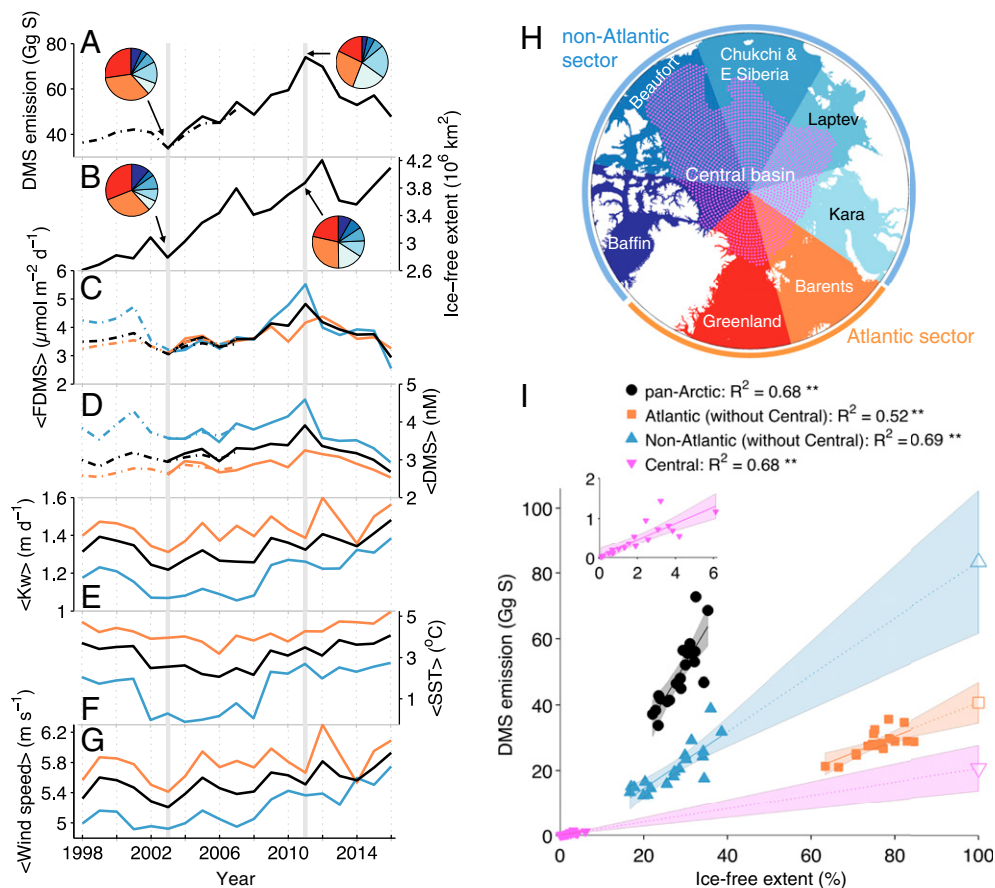
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**Fig. 4.** Interannual variations in EDMS and its driving factors north of 70°N. All panels show May–August integrals (EDMS) or means (other variables). (A) EDMS, (B) mean ice-free ocean extent, (C) mean FDMS, (D) mean DMS concentration, (E) sea–air DMS transfer coefficient ( $K_{wa}$ ), (F) SST, (G) wind speed. Values in C–G correspond to ice-free pixels only. Different lines correspond to the whole domain >70°N and its division into 2 sectors: Atlantic-influenced and non-Atlantic regions, according to the map in H. The pie charts in A and B illustrate the relative contribution of different source regions (colored following H) to EDMS, and their corresponding share of ice-free extent, in the years with lowest (2003) and highest (2011) EDMS. (I) shows the domain-specific relationship between % ice-free extent and EDMS. In this plot, the Central Arctic basin (pink stippling in H) is treated separately, and its area subtracted from the other regions to calculate the regressions (the regression for the Central basin is also shown in the *Inset*). Empty symbols show extrapolation to 100% ice-free extent for the Atlantic, non-Atlantic, and Central domains, and shaded areas show the 95% CI of predictions.

The scenario described above is consistent with conceptual (7) and numerical (5) model predictions suggesting that salinity stratification and resulting nutrient limitation will prevent large increases in phytoplankton production in the Arctic as sea-ice loss proceeds. However, our EDMS estimates for an idealized ice-free scenario suffer from multiple sources of uncertainty. First, the observed linear relationship between ice-free extent and EDMS will not necessarily hold in the future, and its extrapolation is particularly speculative in domains with high ice cover at present (Fig. 4I). Second, estimation of future EDMS is confounded by additional layers of complexity that interact with each other (14): the response of plankton communities to multiple stressors, with a potentially prominent but poorly understood role for acidification (38); the strong taxonomic dependence of DMSP synthesis; and the complex biogeochemical cycling of DMS in seawater (13, 14).

The relationship between airborne MSA and phytoplankton production in historical records could help constrain EDMS projections (14, 18, 19). MSA trapped in Greenland ice cores shows a robust positive relationship with in situ sampled phytoplankton and satellite-observed net primary production (NPP) around southern Greenland over several decades (19). This observation supports extrapolation of the satellite-era EDMS trends. Yet, the relationship between NPP and MSA may be confounded by variable atmospheric MSA yields (14, 30) and by differences in phytoplankton taxonomy across source regions (18, 36). In the

light of these findings, the response of Arctic phytoplankton to environmental forcing appears particularly critical. A warmer, more stable, and irradiated water column (7, 15) might favor nanoplanktonic strong DMS producers, like coccolithophores (10) and *Phaeocystis pouchetii* (20). Such a taxonomic shift could enhance EDMS from the seasonal ice zone, offsetting other processes suspected to affect negatively EDMS, e.g., acidification (38).

Understanding and predicting how changes in marine EDMS will affect the Arctic climate requires progress in many fronts. Although atmospheric models still strive to represent aerosol (24, 26) and cloud (3, 33) dynamics, there is growing consensus that 1) DMS is an essential ingredient for Arctic marine boundary layer nucleation; and 2) nucleation rates will increase in the future (3, 24, 26) owing to concomitant increases in atmospheric humidity, aerosol wet removal, and marine aerosol precursor emissions. In line with these predictions, increasing frequency of aerosol nucleation events has been clearly linked to ice retreat at the Mt. Zeppelin observatory (78.9°N) (27).

Current knowledge suggests that CCN concentrations are unlikely to increase as much as new particle formation (nucleation) rates, due to a concomitant increase in aerosol removal (24, 26). Yet, DMS will still play a critical role in seeding and sustaining CCN populations, and could also affect precipitation (26, 39). Widening the focus, the impact of increasing EDMS on CCN populations will also depend on changing anthropogenic sulfur



emissions and their transport to the Arctic. Ongoing reductions in power plant emissions in the Northern Hemisphere (40) may magnify the role of DMS or extend its seasonal dominance, unless they are compensated by increasing shipping, industrialization, or oil and gas extraction in the Arctic.

The future response of cloud radiative forcing is also uncertain. Unlike in lower latitudes, low-level marine clouds in the Arctic act to retain heat in the ocean–atmosphere system during most of the year (31). Net cloud cooling is currently restricted to a short midsummer period when high solar elevation and low ice cover cooccur, but the seasonal radiation budget will change as ice recedes. Extremely low CCN concentrations that generally occur over the ice pack imply strong sensitivity to CCN changes (32), amplifying uncertainty in indirect aerosol forcing (33). An increase in CCN concentrations is generally associated with enhanced cloud albedo and a cooling effect (3, 29, 33), with recent estimates suggesting a shortwave forcing of  $-1$  to  $-2$   $W\ m^{-2}$  in response to a 2- to 5-fold increase in Arctic EDMS (3) (*SI Appendix*). However, a CCN increase might also enhance longwave cloud forcing over the CCN-depleted pack ice (32). This response is poorly quantified and could offset shortwave forcing, causing net warming and further accelerating ice melt (25, 27).

Our study highlights the key role of atmospheric forcing in driving Arctic EDMS through the control of ice retreat (41), plankton dynamics (7, 14), and gas exchange. Since large-scale weather systems also determine air-mass transport pathways (25), the fate of atmospheric DMS and its interaction with aerosols and clouds cannot be fully understood by analyzing climatological fields in the variable and heterogeneous Arctic environment. The space- and time-resolved FDMS estimates presented here provide a key constraint for atmospheric models and can help reduce uncertainty in projections of aerosol direct and indirect forcing (3, 24, 26, 33). This can in turn improve our understanding of contemporary plankton-climate feedbacks through the interaction of multiple processes, including ocean–atmosphere exchange of  $CO_2$ , other greenhouse gases, and aerosol precursors such as DMS (29, 38). Changing EDMS has wide implications for the vulnerable Arctic environment, its human populations, and the weather and climate of lower latitudes (1).

## Methods

**Remote Sensing Algorithms.** Daily level 3 composites of remote sensing reflectance spectra acquired by SeaWiFS and MODIS-Aqua were used to retrieve Chl and euphotic layer depth (Zeu) [along with absorption coefficients of colored detrital matter ( $a_{CDM(412)}$ )]. These data were used as input to the DMS<sub>SAT</sub> algorithm (17). First, sea-surface DMS<sub>Pt</sub> ( $nmol\ L^{-1}$ ) was estimated as a function of chlorophyll *a* (Chl) concentration using 2 different equations depending on the phytoplankton light exposure regime. Second, sea-surface DMS concentration ( $nmol\ L^{-1}$ ) was estimated from DMS<sub>Pt</sub> and photosynthetically available radiation, after binning these variables to 8-d 28-km resolution to achieve full coverage. Remotely sensed sea-ice concentration (SIC) was used to screen out ice-contaminated pixels ( $SIC > 10\%$ ). DMS<sub>SAT</sub> was calibrated and validated for the Arctic region using in situ DMS and DMS<sub>Pt</sub> data from a public database supplemented with recent datasets. DMS<sub>SAT</sub> match-ups with in situ DMS yielded similar validation statistics for SeaWiFS and MODIS-Aqua. Detailed information on algorithm tuning, implementation, and validation is provided in *SI Appendix*.

**DMS Sea–Air Flux.** FDMS was estimated as the product of the sea–air gas transfer coefficient ( $K_w$ ) and the DMS gradient across the sea–air interface using standard gas-exchange parameterizations based on wind speed. We took into account air- and water-side resistance and the effects of SST and salinity on DMS diffusivity and solubility (*SI Appendix, section 4*).

**Large-Scale EDMS.** We estimated EDMS by integrating FDMS over different periods and spatial domains (e.g., zonal bands and longitude sectors). The 200-m isobath was used to partition EDMS into open-ocean and continental shelf domains. Within continental shelves, a threshold of colored detrital matter  $a_{CDM(412)} > 0.25\ m^{-1}$  was used to screen for riverine influence. High Arctic summer EDMS, defined as the May–August emission north of  $70^\circ N$ , was further regionalized into 7 longitude sectors (Fig. 4H). The Greenland and Barents Seas and the remaining 5 sectors were grouped into the Atlantic and non-Atlantic domains, respectively.

**Relationship Between Ice Cover and EDMS.** We computed linear least-squares regressions between EDMS and the percentage of open-ocean water (pixels with  $\leq 10\%$  SIC) for the high Arctic ( $> 70^\circ N$ ) and for 3 domains within it: the Atlantic sector, the non-Atlantic sector, and the Central Arctic basin. Extrapolation to 100% ice-free water gave an estimate of future EDMS within a given domain, and the sum of extrapolated quantities gave a pan-Arctic estimate of future EDMS (Fig. 4). Extrapolation based on the regression over the entire domain (which yielded  $200 \pm 54\ Gg\ S$ , larger than the sum of domain-specific EDMS estimates) was discarded owing to uneven ice-free extent (%) and mean FDMS across domains. Uncertainty in future EDMS was propagated by adding in quadrature the 95% CI of the extrapolated predictions for each domain. Uncertainty in the fold change with respect to present-day EDMS also took into account the uncertainty in present-day EDMS estimates (2 SDs of 2011–2016 mean EDMS). Slopes obtained from alternative types of regression (type II major axis) were not significantly different from those obtained from regular linear least squares. Additional regressions between EDMS and ice-free extent (million  $km^2$ ) in smaller longitude sectors are shown in *SI Appendix*.

**Multiyear Trends and Means.** We computed linear least-squares regression slopes of satellite-diagnosed quantities (e.g., EDMS) over time (in decades). Regressions were calculated for the entire study period (1998–2016,  $n = 19$  y) and, north of  $70^\circ N$ , for 3 subperiods showing distinct trends. We also calculated mean EDMS during the initial (1998–2003) and final (2011–2016) 6 y, which represent the contiguous years with highest and lowest sea-ice extent, respectively, and without significant trends in sea-ice extent ( $P > 0.05$ ). Division of regression slopes by the mean EDMS during 1998–2003 yielded relative rates of change ( $\% decade^{-1}$ ). To compute 19-y trends we checked the coherence between SeaWiFS and MODIS-Aqua records between 2003 and 2007, and corrected for small offsets ( $< 3.5\%$ ; Fig. 4A) prior to regression analysis.

**Uncertainty Assessment.** We assessed the sensitivity of EDMS to variations in input satellite products and algorithm configuration (*SI Appendix, section 5 and Table S3*). This analysis was conducted only for the MODIS-Aqua record, which largely drives observed temporal trends. The sensitivity tests included random perturbation of DMS algorithm coefficients, use of alternative Chl products and gas-exchange schemes, and replacement of time-varying DMS fields by climatologies (DMS<sub>SAT</sub> and L11).

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