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Boundary layer and free-tropospheric dimethyl sulfide in the Arctic spring and summer

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Abstract. Vertical distributions of atmospheric dimethyl sulfide (DMS(g)) were sampled aboard the research aircraft *Polar* 6 near Lancaster Sound, Nunavut, Canada, in July 2014 and on pan-Arctic flights in April 2015 that started from Longyearbyen, Spitzbergen, and passed through Alert and Eureka, Nunavut, and Inuvik, Northwest Territories. Larger mean DMS(g) mixing ratios were present during April 2015 (campaign mean of 116 ± 8 pptv) compared to July 2014 (campaign mean of 20 ± 6 pptv). During July 2014, the largest mixing ratios were found near the surface over the ice edge and open water. DMS(g) mixing ratios decreased with altitude up to about 3 km. During April 2015, profiles of DMS(g) were more uniform with height and some profiles showed an increase with altitude. DMS reached as high as 100 pptv near 2500 m.

Relative to the observation averages, GEOS-Chem (www. geos-chem.org) chemical transport model simulations were higher during July and lower during April. Based on the simulations, more than 90 % of the July DMS(g) below 2 km and more than 90 % of the April DMS(g) originated from Arctic seawater (north of 66° N). During April, 60 % of the DMS(g), between 500 and 3000 m originated from Arctic

seawater. During July 2014, FLEXPART (FLEXible PAR-Ticle dispersion model) simulations locate the sampled air mass over Baffin Bay and the Canadian Arctic Archipelago 4 days back from the observations. During April 2015, the locations of the air masses 4 days back from sampling were varied: Baffin Bay/Canadian Archipelago, the Arctic Ocean, Greenland and the Pacific Ocean. Our results highlight the role of open water below the flight as the source of DMS(g) during July 2014 and the influence of long-range transport (LRT) of DMS(g) from further afield in the Arctic above 2500 m during April 2015.

1 Introduction

The Arctic has experienced rapid climate change in recent decades (IPCC, 2013). Its high climate sensitivity distinguishes the Arctic from the rest of the world. The Arctic Ocean moderates Arctic climate and has variable surface temperature and salinity as ice cover melts and freezes (Bourgain et al., 2013). This ocean is an important source of atmospheric gases and particles (e.g. dimethyl sulfide,

as well as sea salt, organic and biogenic particles) (e.g. Bates et al., 1987; Andreae, 1990; Yin et al., 1990; Leck and Bigg, 2005a, b; Barnes et al., 2006; Ayers and Cainey, 2007; Sharma et al., 2012). Aerosols affect the climate by scattering/reflecting sunlight (direct effects), changing the number/size of cloud droplets and altering precipitation efficiency (indirect effects) (Twomey, 1974; Albrecht, 1989). Shupe et al. (2013) provided the evidence for the formation of clouds and transport of moisture and aerosol particles, likely accompanied by warm air masses, from lower latitudes into the central Arctic during summer. The study of these particles has been of interest for numerous researchers because of their importance in Arctic climate change. Najafi et al. (2015) estimated that the net effect of aerosol is cooling the Arctic. However, there are many uncertainties related to the estimation of effects and sources of aerosol particles. In this study, we focus on one of those sources: DMS(g) (atmospheric dimethyl sulfide).

Atmospheric oxidation of DMS(g) is the main source of biogenic sulfate aerosols in the Arctic (Norman et al., 1999). DMS(aq) (DMS in aqueous phase) is produced by the breakdown of dimethylsulfonopropionate (DMSP) by oceanic phytoplankton and bacteria DMSP lyases (Levasseur, 2013) and transported to the atmosphere via turbulence, diffusion and advection (Lunden et al., 2010). Sulfur compounds from atmospheric DMS(g) oxidation are able to form new particles and condense on pre-existing aerosols in the atmosphere (Chang et al., 2011). If sufficient condensable vapours are available, the particles may grow large enough to act as cloud condensation nuclei (CCN), and Charlson et al. (1987) hypothesised that DMS could provide a negative feedback to stabilise the global warming (CLAW hypothesis). Although no evidence in support of the hypothesis has been found (Quinn and Bates, 2011), DMS(g) emissions may play an important role in the climate of remote areas with low aerosol concentrations, such as in the Arctic (Carslaw et al., 2013; Leaitch et al., 2013; Levasseur, 2013; Croft et al., 2016a).

Dimethyl sulfide production and emission to the atmosphere vary seasonally. Production and emission are particularly strong during the Arctic summertime due to high temperature, biological activity, and the amount of ice-free surface area. Melting ice in the marginal ice zone, ice edge and under-ice are favourable for the production of DMSP and aqueous DMS(aq) by oceanic phytoplankton (Leck and Persson, 1996; Matrai and Vernet, 1997; Levasseur, 2013). After summer, aqueous phase DMS(aq) concentrations decrease by about 3 orders of magnitude between August and October in the central Arctic Ocean (Leck and Persson, 1996).

Dimethyl sulfide oxidation in the atmosphere occurs by the radical addition pathway (by hydroxyl radicals OH and halogen oxides) and by the H abstraction pathway (by the nitrate radical NO₃, OH and halogens) (Barnes et al., 2006; von Glasow and Crutzen, 2004). In general, the DMS(g) oxidation rate and pathway depends on the available oxidants and temperature. The final products of DMS(g) oxidation by the addition pathway are dimethyl sulfoxide (DMSO) and methanesulfonic acid (MSA). DMSO oxidises in cloud droplets to methanesulfonic acid, due to its high solubility, and MSA likely condenses onto pre-existing aerosols (von Glasow and Crutzen, 2004). On the other hand, DMS(g) oxidation by the abstraction pathway leads to the formation of SO₂. Some of SO₂ removes from the atmosphere via dry and wet deposition, and the remaining SO₂ may form sulfuric acid (H₂SO₄) in the gas and aqueous phases (Pierce et al., 2013). Sulfuric acid formed in the gas phase is a key atmospheric nucleation component which is able to form new particles that may grow to the size of CCN and affect climate (Kulmala et al., 2004).

Previous measurements of DMS(g) in the Arctic atmosphere are limited to a few studies and field campaigns at different locations (e.g. Sharma et al., 1999; Rempillo et al., 2011; Mungall et al., 2016). The study of the vertical distribution of DMS(g) in the Arctic atmosphere is also limited to a few observations. Ferek et al. (1995) reported the first measurements of DMS(g) vertical profiles over the Arctic Ocean near Barrow in early summer 1990 and spring 1992. They reported low DMS(g) mixing ratios (a few pptv) during spring and relatively high ones (a few tens pptv with some peaks around 100 to 300 pptv) during summer. They concluded that the Arctic Ocean is the potential source of DMS(g), and DMS(g) ocean–atmosphere exchange is more important in early summer due to sea ice melt.

Observations of the NASA DC-8 during ARCTAS (https: //www-air.larc.nasa.gov/cgibin/ArcView/arctas) showed low DMS mixing ratios in spring (below the detection limit to a few pptv in the boundary layer and a maximum of 1 pptv in the free troposphere) (Simpson et al., 2010; Lathem et al., 2013).

Kupiszewski et al. (2013) measured atmospheric DMS(g) on board a helicopter and observed large variability in DMS(g) mixing ratios over the central Arctic Ocean during summer. The median (mean) values were around 7 (34) pptv near the surface < 200 m, 11 (22) pptv for altitudes between 200 and 1000 m, and 4 (5) pptv above 1000 m.

Lunden et al. (2010) presented model results for the vertical distribution of DMS(g) in the Arctic (north of 70° N) during summer. They reported a variable vertical profile for DMS(g) concentrations above open water, with the highest concentrations near the surface (around 115 and 365 pptv for the median and 95th percentiles, respectively) and an exponential decrease with height. In contrast, over the pack ice, DMS(g) concentrations were higher above the local boundary layer than at the surface. Also, Lunden et al. (2010) showed that DMS(g) can be mixed downward by turbulence into the local boundary layer to provide a DMS source over the pack ice. In addition, they compared modelling results with measurements from the Arctic Ocean Expedition 2001 (AOE-2001; Leck et al., 2004; Tjernström et al., 2004) and reported that DMS(g) was present above the local boundary layer in both the model and observations.

For our study, atmospheric DMS(g) samples were collected onto Tenax tubes during Polar 6 aircraft flights in the Arctic. We compared these DMS(g) measurements to GEOS-Chem (www.geos-chem.org) chemical transport model simulations and conducted sensitivity simulations to examine the local vs. long-range transport (LRT) DMS(g) sources for both the spring and summer. In addition, FLEXPART (FLEXible PARTicle dispersion model) was applied in back trajectory mode in order to investigate the DMS(g) source regions based on potential emission sensitivity simulations. Field and sampling locations, as well as measurement and modelling methods are described in Sect. 2. Section 3 includes DMS(g) measurement data. Section 4 presents discussion of results, and comparison of measurement with modelling results (GEOS-Chem and FLEXPART) are in Sect. 5. The summary and conclusion of this study are reported in Sect. 6.

2 Field description and methods

2.1 Measurements

2.1.1 DMS

DMS(g) was collected aboard the research aircraft *Polar 6* in the Arctic during July 2014 and April 2015, as part of the NETCARE (Network on Climate and Aerosols: Addressing Key Uncertainties in Remote Canadian Environments) project. The *Polar 6* aircraft routes and sampling locations from 12 to 21 July 2014 and from 5 to 20 April 2015 are shown in Figs. 1 and 2, respectively. The *Polar 6* campaign was based in Resolute Bay, Nunavut, and covered the Lancaster Sound area in July 2014. In April 2015, the flights started from Longyearbyen, Spitzbergen, and passed through Alert and Eureka, Nunavut, and Inuvik, Northwest Territories. DMS sampling locations, altitude, latitude and longitude are reported in Table 1.

Atmospheric DMS(g) was collected on cartridges packed with Tenax TA[®]. Mass flow was controlled at approximately $200 \pm 20 \text{ mL min}^{-1}$, and a KI-treated 47 mm quartz Whatman filter was fitted at the intake of the cartridge to remove all oxidants. Two Teflon valves were placed before and after the Tenax tube to control the sampling period, and Teflon tubing was used to transfer the sample from outside the aircraft to the sampler. The samples were stored in an insulated container with a freezer pack after collection and in a freezer after the flight. Sampling collection time was 300 ± 5 s (for few samples the sampling time was shorter or longer than 300 s, leading to different volume of samples).

A glass gas chromatograph (GC) inlet liner was used to pack 170 \pm 2 mg of Tenax. The Tenax packed in glass tubes was cleaned by heating to 200 °C in an oven with a constant He flow of around 15 mL min⁻¹ for 5 h. The DMS samples were analysed using a Hewlett Packard 5890 gas chromatograph (GC) fitted with a Sievers Model 355 sulfur chemiluminescence detector (SCD). Two DMS(g) certified standards from Praxair (1 and 50 ppmv) were used to calibrate the GC-SCD and to determine the accuracy of the measurements by checking the standards against each other (for example, 1 μ L of 50 ppmv vs. 50 μ L of 1 ppmv). Collection and analysis of samples were based on methods described by Sharma (1997), Sharma et al. (1999) and Rempillo et al. (2011). Precision of analysis was ±12 pptv and was determined based on the SD (σ) of triplicate measurements of DMS(g) standards. The detection limit for this method is approximately 7 pptv.

Additional tests were performed to determine if there was significant loss of DMS(g) over time after collection. An experiment was performed to determine how long Tenax is able to store DMS(g) with no significant loss of concentration. This experiment was conducted in triplicate by the loading of $50 \,\mu$ L of 1 ppmv DMS(g) standard and by storing it in a freezer at -25 °C. In general, Tenax storage tests at -25 °C showed that DMS losses were approximately 5 and 15 % after 10 and 20 days, respectively (Fig. 3). The DMS(g) mixing ratios summarised in Table 1 are adjusted according to the result of this test.

2.1.2 Meteorological measurements

Meteorological measurements were performed by an AIMMS-20 (Aircraft Integrated Meteorological Measurement System) instrument, manufactured by Aventech Research Inc., Barrie, Ontario, Canada. This instrument was used to measure the three-dimensional, aircraft-relative flow vector (true air speed, angle of attack, and sideslip), temperature, relative humidity, turbulence and horizontal/vertical wind speeds. Accuracy and resolution were 0.30 and 0.01 °C, respectively, for temperature and 2.0 and 0.1 %, respectively, for relative humidity. More details of the instrument and corresponding aircraft measurements were recently published in other studies from the same campaign (e.g. Leaitch et al., 2016; Aliabadi et al., 2016b; Willis et al., 2016).

2.2 Model description

2.2.1 GEOS-Chem chemical transport model

The GEOS-Chem chemical transport model was used to interpret the vertical profile of DMS(g). We used GEOS-Chem version 9-02 at $2 \times 2.5^{\circ}$ resolution with 47 vertical layers between the surface and 0.01 hPa. The assimilated meteorology is taken from the National Aeronautics and Space Administration (NASA) Global Modeling and Assimilation Office (GMAO) Goddard Earth Observing System version 5.7.2 (GEOS-FP) assimilated meteorology product, which includes both hourly surface fields and 3-hourly 3-D fields. Our simulations used 2014 and 2015 meteorology following a 1-month spin-up prior to the simulation of July 2014 and April 2015.

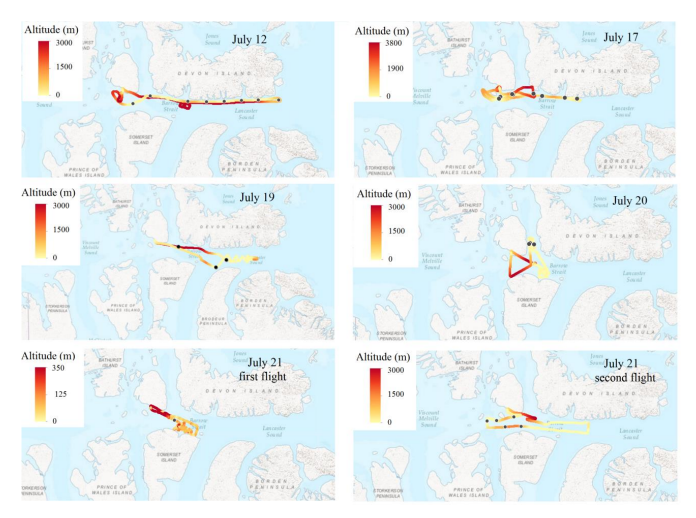


Figure 1. Polar 6 aircraft routes from 12 to 21 July 2014. Colour bars indicate altitudes, and sampling locations are shown with black dots.

The GEOS-Chem model includes a detailed oxidantaerosol tropospheric chemistry mechanism as originally described by Bey et al. (2001). DMS(g) emissions are based on the Liss and Merlivat (1986) sea-air flux formulation and oceanic DMS(g) concentrations from Lana et al. (2011). In our simulations, DMS(g) emissions occurred only in the fraction of the grid box that is covered by seawater and also free of sea ice. Simulated DMS(g) oxidation occurs by reaction with OH and NO3. The model also includes natural and anthropogenic sources of SO₂ and NH₃ (Fisher et al., 2011). Oxidation of SO₂ occurs in clouds by reaction with H₂O₂ and O₃ and in the gas phase with OH (Alexander et al., 2009). Reaction rates and the yields of SO₂ and MSA from DMS(g) oxidation are determined by De-More et al. (1997) and Chatfield and Crutzen (1990), respectively. The simulated aerosol species include sulfate-nitrateammonium (Park et al., 2004, 2006), carbonaceous aerosols (Park et al., 2003; Liao et al., 2007), dust (Fairlie et al., 2007, 2010) and sea salt (Alexander et al., 2005). The sulfatenitrate-ammonium chemistry uses the ISORROPIA II thermodynamic model (Fountoukis et al., 2007), which partitions ammonia and nitric acid between the gas and aerosol phases. Climatological biomass burning emissions are from the Global Fire Emissions Dataset (GFED3).

The GEOS-Chem model has been extensively applied to study the Arctic atmosphere, with regard to aerosol acidity (Wentworth et al., 2016; Fisher et al., 2011), carbonaceous aerosol (Wang et al., 2011), aerosol number (Leaitch et al., 2013; Croft et al., 2016a, b), aerosol absorption (Breider et al., 2014), mercury (Fisher et al., 2012) and recently surface-layer DMS(g) (Mungall et al., 2016).

2.2.2 FLEXPART-ECMWF

For this study, the Lagrangian particle distribution model, FLEXPART (Stohl et al., 2005; website: https://www.flexpart.eu/), is driven by global meteorological analysis data from the European Centre for Medium-Range Weather Forecasts (ECMWF) for July 2014 and April 2015. For the ECMWF data a horizontal grid spacing of 0.25° was used along with 137 hybrid sigma-pressure levels in the vertical

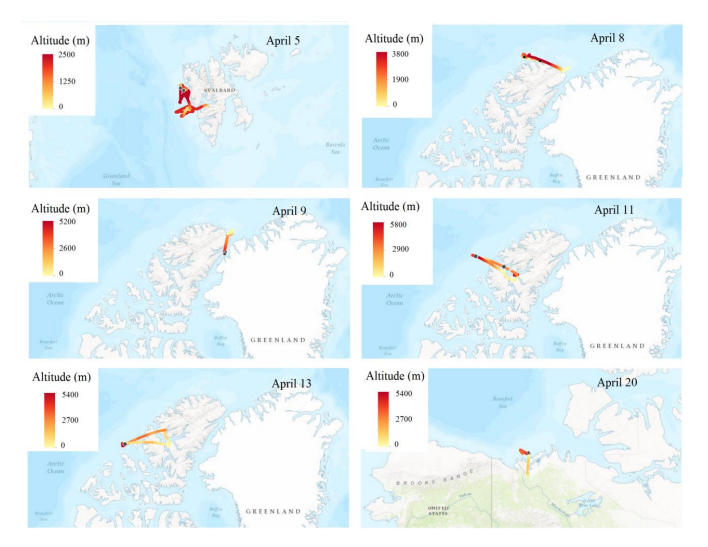


Figure 2. Polar 6 aircraft routes from 5 to 20 April 2015. Colour bars indicate altitudes, and sampling locations are shown with black dots.

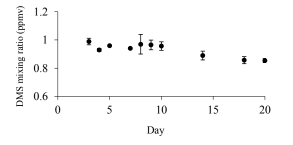


Figure 3. DMS mixing ratios vs. Tenax storage days. Error bars indicate the SD for each test.

from the surface up to 0.01 hPa. FLEXPART was operated in backward mode to estimate potential emission sources and transport pathways influencing *Polar 6* DMS(g) measurements in summer 2014 and spring 2015. For this, plumes of a passive tracer with properties of air (i.e. molar mass of dry air, no removal) were released along the flight paths every minute. These plumes were traced back for several days to study the potential origin of the air masses sampled during the flights with the *Polar* 6.

3 DMS measurement and discussion

DMS(g) concentrations as a function of altitude are shown in Fig. 4 for the July 2014 and April 2015 flights. The campaign-mean DMS(g) mixing ratios were 20 ± 6 pptv (maximum of 114 pptv) for July 2014 and 116 ± 8 pptv (maximum of 157 pptv) for April 2015.

The 2014 sampling locations focused on the Lancaster Sound, Nunavut region, in July 2014, whereas sampling in April 2015 occurred over a broad region of the Arctic: Longyearbyen, Spitzbergen; Alert and Eureka, Nunavut; and Inuvik, Northwest Territories. Observations on individual flights in July 2014 indicate either decreasing DMS(g) mixing ratios with increasing altitude or relatively uni-

Sample no.	DMS (pptv)	Sampling day	Analysis day	Latitude (°)	Longitude (°)	Altitude (m)
1	17	12 Jul 2014	25 Jul 2014	74.45	-79.87	872 (240) ^b
2	39	12 Jul 2014	25 Jul 2014	74.45	-81.85	78
3	26	12 Jul 2014	25 Jul 2014	74.44	-83.47	275
4	14	12 Jul 2014	25 Jul 2014	74.41	-85.11	69
5	24	12 Jul 2014	25 Jul 2014	74.4	-86.85	280
6	9	12 Jul 2014	25 Jul 2014	74.41	-88.63	880
7	11	12 Jul 2014	25 Jul 2014	74.55	-92.28	303
8	below detection	12 Jul 2014	25 Jul 2014	74.36	-93.97	57
9	below detection	17 Jul 2014	27 Jul 2014	74.47	-94.88	63
10	8	17 Jul 2014	27 Jul 2014	74.45	-88.64	159
11	8	17 Jul 2014	27 Jul 2014	74.42	-87.44	149
12	below detection	17 Jul 2014	27 Jul 2014	74.42	-95.01	58
13	below detection	17 Jul 2014	27 Jul 2014	74.54	-93.71	884
14	below detection	17 Jul 2014	27 Jul 2014	74.56	-91.66	2862
15	below detection	19 Jul 2014	27 Jul 2014	74.51	-92.20	2862
16	below detection	19 Jul 2014	27 Jul 2014	74.10	-86.53	165
17	9	19 Jul 2014	27 Jul 2014	73.86	-87.75	46
18	44	20 Jul 2014	25 Jul 2014	74.95	-93.09	67 (384) ^b
19	below detection	20 Jul 2014	25 Jul 2014	74.92	-92.72	97
20	below detection	20 Jul 2014	25 Jul 2014	74.93	-93.18	143
21	22	21 Jul 2014	25 Jul 2014	74.39	-92.70	79 (213) ^b
22	below detection	21 Jul 2014	25 Jul 2014	74.29	-93.62	51
23	below detection	21 Jul 2014	25 Jul 2014	74.28	-95.15	61
24	13	21 Jul 2014	25 Jul 2014	74.43	-96.91	61
25	114	21 Jul 2014	25 Jul 2014	74.45	-95.98	61
26	9	21 Jul 2014	25 Jul 2014	74.54	-94.35	65
27	127	5 Apr 2015	7 May 2015	78.91	10.31	2390
28	147	5 Apr 2015	7 May 2015	78.94	10.82	515
29	66	8 Apr 2015	7 May 2015	83.07	-71.99	1986
30	72	8 Apr 2015	7 May 2015	83.24	-78.59	76
31	122 ^a	9 Apr 2015	7 May 2015	81.43	-63.39	78
32	134	11 Apr 2015	8 May 2015	80.77	-87.85	2733
33	112 ^a	11 Apr 2015	8 May 2015	81.50	-99.72	68
34	125	11 Apr 2015	8 May 2015	81.57	-100.72	1244
35	75	13 Apr 2015	8 May 2015	80.06	-104.08	5015
36	123	13 Apr 2015	8 May 2015	80.09	-104.05	2651
37	135 ^a	13 Apr 2015	8 May 2015	80.13	-103.95	79
38	157	20 Apr 2015	8 May 2015	70.00	-133.16	59

Table 1. DMS mixing ratio values, sampling/analysis dates and sampling locations for July 2014 and April 2015.

^a Examples of DMS samples concurrent with ozone depletion events (< 1 ppbv) during April. ^b Numbers in parentheses show the boundary layer heights in Resolute Bay from Aliabadi et al. (2016b).

form DMS(g) mixing ratios (independent of altitude below 3 km). During spring of the following year (April 2015), DMS(g) mixing ratios on individual flights were more uniform with altitude below 4 km and in some cases increased with altitude.

Figure S1 in the Supplement shows the ice fraction for July 2014 flights. During July 2014, the highest DMS(g) mixing ratios were measured near ice edges and above open waters (e.g. samples > 40 pptv; 12, 20 and 21 July). That and the decrease in atmospheric DMS(g) with altitude suggest that the atmospheric DMS(g) was locally sourced (Lancaster Sound and Baffin Bay) during the month of July, con-

sistent with the findings of Mungall et al. (2016) from the icebreaker CCGS *Amundsen*. Mungall et al. (2016) also suggested LRT of DMS from marine regions outside Baffin Bay and the Lancaster Sound area and observed that an episode of elevated DMS(g) mixing ratios with values of 400 pptv or above occurred on 18–20 July. The airborne measurement, showed a decline in DMS(g) mixing ratios with height during 17 July and relatively low DMS mixing ratios during 19 and 20 July (see Table 1). The decline in DMS(g) mixing ratios with height may be due to a combination of weak vertical mixing and photochemical reactions.

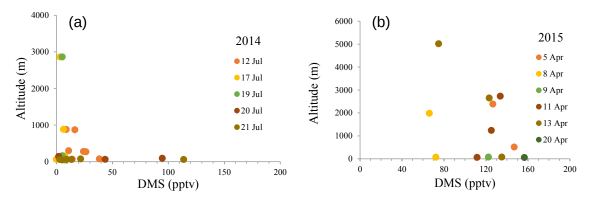


Figure 4. Sampling altitudes (m) vs. DMS mixing ratios (pptv) for July 2014 (a) and April 2015 (b).

Previous observations of seasonal variations in DMS(aq) in the Arctic Ocean found that the maximum DMS(aq) occurred in July and August (e.g. Bates et al., 1987; Leck and Persson, 1996; Levasseur, 2013). After the August peak, DMS(aq) declined due to lower biological activity (Leck and Persson, 1996). From DMS concentrations in both the surface ocean and in the atmosphere just above the ocean surface (median DMS(g) of 186 pptv), Mungall et al. (2016) estimated the air-sea DMS(g) flux as ranging from 0.02- $12 \,\mu\text{mol}\,\text{m}^{-2}\,\text{d}^{-1}$ in July 2014 in the same location as the present measurements (Lancaster Sound). For the same campaign, Ghahremaninezhad et al. (2016) showed that the dominant source for fine aerosol and SO₂ measured onboard the Amundsen at the same location and about 30 m above the ocean's surface was biogenic sulfur, arising from DMS(g) oxidation. Atmospheric oxidation of DMS(g) is expected to proceed more readily in the summertime Arctic atmosphere than in spring, due to higher temperatures and more sunlight. However, relatively high DMS mixing ratios (> 15 pptv) were observed for 12 July at high altitudes ($> 800 \,\mathrm{m}$), and FLEXPART results show influence from a local source, Lancaster Sound, for that day (mentioned in Sect. 4.2). On this day, NETCARE results do not follow the usual DMS vertical pattern of high DMS at the surface declining with altitude to near zero above the marine boundary layer (MBL). Instead, high concentrations aloft on 12 July imply convective transport into the free troposphere and potentially an extended photochemical lifetime due to reduced water vapour or limited sunlight.

During April, DMS(g) samples were collected above ice and snow surfaces, and heat fluxes were negligible. Figure S3 shows the ice fraction during the April 2015 campaign. The higher DMS(g) mixing ratios in April, in the free troposphere over ice-covered regions (Fig. 4b), stability of the Arctic atmosphere, and limited vertical mixing, suggest that DMS(g) can be transported to the sampling locations from other regions within the Arctic and/or from lower latitudes (except for 4 April when DMS(g) sampling was above open water). These results contrast with results from Ferek et al. (1995) where lower DMS(g) mixing ratios (a few pptv) were found over the Arctic Ocean near Barrow during spring 1992. Andrea et al. (1988) presented vertical profiles of DMS(g) mixing ratios measured over the northeast Pacific Ocean during May 1985 (with a maximum ~ 30 pptv in the mixed layer and also 3600 m). They found that DMS(g) mixing ratios depend on the stability of the atmosphere and air mass sources and that long-range transport at mid-tropospheric levels was important in remote areas of the Northern Hemisphere.

The relatively larger observed DMS(g) away from openwater sources in springtime relative to summer suggests longer DMS(g) lifetimes in April than July, possibly due to lower OH mixing ratios enabling more long-range transport of DMS(g) (Li et al., 1993). Lower water vapour and higher DMS mixing ratios during the spring compared with the summer (Fig. S2) suggests that more of the April DMS(g) originated from open-water sources further away from the observations point than in summertime. The greater ice cover and increased presence of DMS(g) at higher altitudes during April suggest an origin from further south than in summertime. More water vapour will initially accompany that DMS(g), but the Arctic is cold in April, especially aloft, and the low water vapour indicates significant loss via cloud processes during transport. Some of the water vapour loss will occur via the ice phase, and DMS oxidation in the aqueous phase was likely relatively insignificant during this time (Henry's Law constant for DMS is relatively small: $0.14 \text{ mol } \text{L}^{-1} \text{ atm}^{-1})$ or the DMS(g) values at their origin were much higher than the present observations.

Ozone depletion during spring was observed within the boundary layer (Fig. S2) and is well documented in the literature (e.g. Barrie et al., 1988). Ozone depletion may further decrease OH near the surface and enhance DMS(g) lifetimes in the boundary layer due to reduced oxidation rates, contributing to the relatively larger springtime DMS(g) in our measurements. However, if DMS is present in the ozone-depleted boundary layer, halogen oxides, such as BrO radical, can be more important during winter and spring

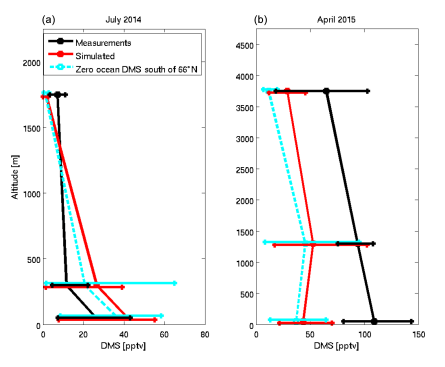


Figure 5. The campaign-mean vertical profile of DMS from the GEOS-Chem simulation (red line) and measurements (black line) for July 2014 and April 2015. Simulations for zero ocean DMS at latitudes south of 66° N (SimZeroBelow66) are shown as cyan dashed line. The 20th and 80th percentiles are shown by horizontal bars

than summer and could oxidise DMS(g) (von Glasow and Crutzen, 2004; Chen et al., 2016).

DMS(g) vertical profiles are sensitive to the boundary layer height. For the July 2014 campaign, Aliabadi et al. (2016b) reported an average boundary layer height of 275 ± 164 m. They showed that the profiles of the potential temperature exhibited a positive vertical gradient throughout the aircraft campaign (their Fig. 4). In addition, using vertical profiles of wind speed, they derived a positive gradient Richardson number (Ri) with a median of 2.5 (their Fig. 7) throughout the aircraft campaign. The magnitude of the positive gradient Richardson number is an indicator of the strength of thermal stability in the atmospheric boundary layer. Due to the strong thermally stable conditions during the field campaign, mixing was weaker compared to wellmixed boundary layers at midlatitudes. As a result, the summertime measurements show a strong decrease in DMS(g)above the boundary layer. Although there is no reference for the April 2015 campaign boundary layer, we expect similar boundary layer characteristics in the stable Arctic boundary layer at high latitudes due to the even more reduced thermal forcing with large sun angles in the month of April compared to the month of July. The springtime measurements show a more uniform vertical profile suggesting transport in the free troposphere from open-water sources that were a relatively farther distance from the observation point in springtime than in summer.

Aerosol number concentrations and size distributions during the July 2014 study are discussed by Willis et al. (2016) and Burkart et al. (2017), who show that increases in the number concentrations of smaller particles (5-20 nm), believed to reflect new particle formation (NPF), occurred principally near the surface during 12 July 2014. The highest levels of DMS(g) during the July study also occurred near the surface (Fig. 4a), and both Willis et al. (2016) and Burkart et al. (2017) noted increased MSA near the surface associated with two case studies of NPF. In the clean conditions of the Arctic summer (e.g. CO in Fig. S2), the low-level DMS may contribute to NPF. The springtime Arctic differs in that the aerosol mass near the surface is much higher, resulting in a higher condensation sink that, in addition to other potential factors, inhibits NPF. During the springtime flights, there was no evidence for NPF near the surface and only a few instances aloft. Unfortunately, no sampling for DMS coincided with those few events, and we cannot say if they were connected with the DMS(g) aloft.

4 Chemical transport model simulations and discussion

4.1 GEOS-Chem

We simulated the vertical profile of DMS(g) mixing ratios with the GEOS-Chem chemical transport model, and the

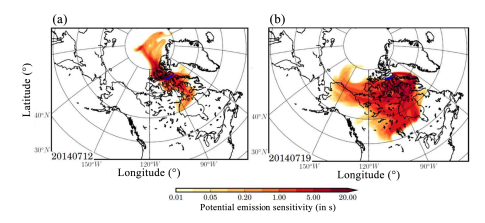


Figure 6. FLEXPART-ECMWF potential emissions sensitivity simulation plots for 4-day back trajectories for column from 0 to 1000 m on (a) 12 July (20:40:00 UTC) and (b) 19 July (17:00:00 UTC) 2014. The colour bars indicate air mass residence time (s) before arriving at the aircraft location. The blue lines show *Polar 6* aircraft routes.

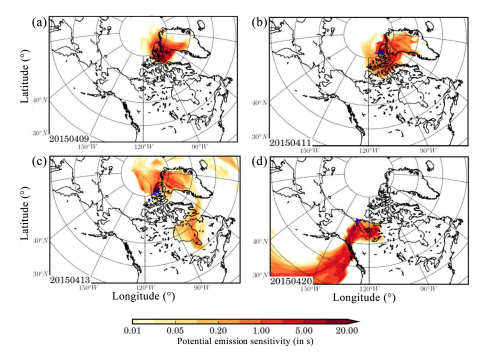


Figure 7. FLEXPART-ECMWF potential emissions sensitivity simulation plots for 4-day back trajectories for column from 0 to 1000 m on (a) 9 April (14:45:00 UTC), (b) 11 April (18:55:00 UTC), (c) 13 April (18:27:00 UTC) and (d) 20 April (22:26:00 UTC) 2015. The colour bars indicate air mass residence time (seconds) before arriving at the aircraft location. The blue lines show *Polar 6* aircraft routes.

model was co-sampled along the *Polar 6* aircraft tracks. Recently, GEOS-Chem was used to interpret DMS(g) measurements in the Arctic surface-layer atmosphere (Mungall et al., 2016). However, despite the significant influence of DMS(g) on the Arctic climate relative to lower latitudes and the importance of where DMS(g) oxidation occurs vertically (Woodhouse et al., 2013), measurements of DMS(g) vertical profiles are rare in the Arctic atmosphere.

Figure 5 shows the campaign-mean vertical profile of DMS(g) for the co-sampled GEOS-Chem simulation and our measurements for both July 2014 and April 2015. Caution

should be used in interpreting the model-measurement comparisons since these comparisons are conducted over a very limited number of measurement periods and the spatial and temporal resolution of these measurements is a challenge for a global model to simulate. In July 2014, both the measurements and simulation show a strong decrease in DMS(g) mixing ratios with altitude in the lowest 300 m. Aliabadi et al. (2016a, b) estimated the boundary layer height as 275 ± 164 m, using data from radiosondes launched at Resolute Bay and the Amundsen icebreaker, during the 2014 campaign. Aliabadi et al. (2016b) indicated that the magnitude of

Table 2. Simulated campaign-mean percent contribution of DMS from oceans north of 66° N to the GEOS-Chem-simulated DMS at the sampling locations for the July 2014 and April 2015 flight tracks.

Altitude	July 2014	April 2015	
0–100 m	98	88	
100–500 m	97	90	
500–3000 m	91	61	

turbulent fluxes of momentum, heat and the associated diffusion coefficients are significantly reduced above the boundary layer height during the 2014 campaign. Thus, we find the strongest vertical gradient between the boundary layer and above. In the boundary layer, the GEOS-Chem simulation over-predicts the measurements but is within a factor of 2 to 3. Hoffmann et al. (2016) showed that DMS chemistry should be considered in the aqueous phase as well as the gas phase to improve modelling predictions. This chemistry is not included in our model but could contribute to the model's over-prediction of the measurements at these lower altitudes. Above 1500 m, the simulation under-predicts the measurements. Overall, the simulations and observations agree within their respective variabilities.

The April 2015 campaign mean shows a more gradual decrease with altitude (Fig. 5b). Mixing ratios are also greater than during the July campaign. Both the simulated and the measured DMS(g) profiles during spring ($\sim 30 \text{ to} > 50 \text{ pptv}$) show more variability at all altitudes below 4 km than in summer (~ 20 to 40 pptv at low altitudes and < 10 pptv at higher altitudes). Ozone depletion not represented by the simulation is one potential explanation for the underestimated DMS(g) in the simulations since the oxidation rates may be too high. Surface layers depleted of ozone were observed on several occasions during April 2015: three of five samples collected at 60 m above ice surfaces were concurrent with measured ozone depletion events (< 1 ppbv) during the April campaign (shown in Table 1). If the DMS(g) oxidation potential is reduced by ozone depletion, the lifetime of DMS(g) in the region of ozone depletion may increase. Another reason for underestimation by the model may be errors in the simulated source strength. The monthly mean seawater DMS field used in our simulations is based on very limited observations from this region (Lana et al., 2011). Datasets of seawater DMS with higher spatial and temporal resolution are needed but are still under development.

We conducted a sensitivity simulation to identify the latitude-dependent contribution of the oceans to the simulated DMS(g) at the sampling points along the flight tracks. In Fig. 5, the "SimZeroBelow66" simulation has no ocean DMS(g) for all latitudes south of 66° N. This simulation compared with the standard simulation suggests that a large

majority of the campaign-mean DMS(g) for both April and July arises from the oceans north of 66° N.

As given in Table 2, SimZeroBelow66 simulates 97 % or more of the DMS(g) below 500 m during July coming from waters north of 66° N. The fractional contribution from north of 66° is about 90% for April and at the same altitudes although different regions were sampled at that time. The simulations attribute about 60 and 90% of the DMS(g) at altitudes of 500 to 3000 m to seawater north of 66° N in April and July, respectively. This 30% difference indicates a greater contribution from long-range transport from lower latitudes in the springtime.

4.2 FLEXPART

FLEXPART-ECMWF modelling was used to explore the origin of air samples measured along the *Polar 6* flight tracks. Figures 6 and 7 show the potential source regions of these air samples 4 days before the releases along the flight path. More specifically, the response function is shown to all releases of a passive tracer, which in this case has properties of dry air. If this response function were folded with an emission flux of the tracer, the concentration of this tracer at the release location along the flight paths could be calculated. We chose to show the potential emission sensitivity after 4 days. Sharma et al. (1999) showed that atmospheric DMS(g) lifetime was 2.5 to 8 days in the high Arctic. More details about FLEX-PART and the potential emissions sensitivity (PES) can be found in Stohl et al. (2005) and references therein.

Figure 6 shows two examples of FLEXPART-ECMWF PES for 4-day back trajectories in July 2014: an influence from a broad area, especially Lancaster Sound (local region), and northwards on 12 July (Fig. 6a), and Hudson Bay, and Baffin Bay (south) on 19 July (Fig. 6b). A more detailed analysis of PES reveals that the measured air mass descended from > 1500 m on 19 July, which may explain the low DMS(g) mixing ratios.

Figure 7 shows some examples of FLEXPART-ECMWF PES simulations for 4-day back trajectories during April 2015. For the flights near Alert and Eureka on 9 and 11 April, some DMS may have originated from ice-free areas of the Nares Strait and Baffin Bay (Fig. 7a and b, respectively). For the 13 April flight, the Norwegian Sea, North Atlantic Ocean and Hudson Bay are additional potential source regions (Fig. 7c). The highest DMS, measured on 20 April near Inuvik is associated with the north Pacific Ocean (Fig. 7d).

Assuming a DMS atmospheric lifetime of 1 to 4 days, these results suggest that the DMS(g) measured during July 2014 originated primarily from the local region over Baffin Bay and the Canadian Arctic Archipelago. For spring 2015, the DMS(g) sampled was from a range of sources, including Baffin Bay, possibly the Norwegian Sea, the North Atlantic Ocean and the north Pacific Ocean.

5 Conclusions

Atmospheric samples for DMS(g) measurements were collected at different altitudes aboard the *Polar 6* aircraft expeditions during July 2014 and April 2015, as part of the NETCARE project. In this study, we present vertical profile measurements of DMS(g), together with model simulations to interpret these profiles. This study includes a very limited spatial and temporal extent, and further vertical profile measurements of Arctic DMS(g) are recommended. Vertical variations in DMS(g) mixing ratios are important since DMS(g) can influence aerosol concentrations via new particle formation and growth. In addition, further DMS vertical measurements could be useful to have a robust comparison with global models such as GEOS-Chem.

Our measured vertical profiles of DMS(g) suggest differences between the main sources and lifetime of DMS(g) during the Arctic summer and spring. For the summertime flights near Lancaster Sound, Nunavut, Canada, DMS(g) mixing ratios were higher near the surface (maximum > 110 pptv) and lower at higher altitudes up to 3 km. The highest mixing ratios were found above ice edges and open waters suggesting that the Arctic Ocean in the vicinity of the aircraft was the main source of DMS(g). Oxidation and/or limited vertical mixing could contribute to the decline in DMS(g) mixing ratios with altitude. During the springtime pan-Arctic flights from Svalbard to the Canadian Arctic Archipelago ending near Inuvik, Northwest Territories, the measured DMS(g) mixing ratios were unusually high (> 100 pptv), and more uniform with altitude than during summer. DMS(g) mixing ratios in samples collected in the free troposphere (> 2000 m) during April ranged from 60 to 134 pptv. Transport of DMS(g) to the high Arctic from other regions of the Arctic and/or lower latitudes along with a reduced oxidising potential in springtime relative to summer may explain these observations.

The DMS(g) vertical profile along the flight tracks was simulated with the GEOS-Chem chemical transport model. The measurement and simulated co-sampled campaign-mean DMS(g) vertical profile agreed within a factor of 3 for both July 2014 and April 2015. A sensitivity test indicated that the oceans north of 66° N contributed about 97 and 90 % of simulated DMS(g) at altitudes below 500 m at the measurement sampling times in July and April, respectively. For the April flights, about 60 % of the simulated DMS at altitudes between 500 and 3000 m was attributed to water north of 66° N. Potential emission sensitivity from FLEXPART analysis for the aircraft tracks showed that local sources (Lancaster Sound and Baffin Bay) primarily contributed to air sampled during July 2014. On the other hand, LRT from the northern tip of Greenland of air that originated over the waters to the northwest of Greenland as well as the north Pacific Ocean were important contributors to air masses sampled during April 2015.

In short, this study suggests a dominant role of the Arctic Ocean for DMS(g) in the Arctic during summer and a significant contribution from LRT to DMS(g) in spring.

Data availability. Data are available by email request (alnorman@ucalgary.ca). The GEOS-Chem model is freely available for download from www.geos-chem.org (Breider et al., 2017).

Competing interests. The authors declare that they have no conflict of interest.

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