Characteristics of methanesulfonic acid, non-sea-salt sulfate and organic carbon aerosols over the Amundsen Sea, Antarctica

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Abstract. To investigate the characteristics of particulate methanesulfonic acid (MSA(p)), non-sea-salt sulfate (nss SO42−) and organic carbon (OC) aerosols, aerosol and seawater samples were collected over the Southern Ocean (43–70° S) and the Amundsen Sea (70–75° S) during the ANA06B cruise conducted in the austral summer of 2016 aboard the Korean icebreaker IBR/V Araon. Over the Southern Ocean, the atmospheric MSA(p) concentration was low (0.10 ± 0.002 µg m−3), whereas its concentration increased sharply up to 0.57 µg m−3 in the Amundsen Sea where Phaeocystis antarctica (P. antarctica), a producer of dimethylsulfide (DMS), was the dominant phytoplankton species. Unlike MSA(p), the mean nss SO42− concentration in the Amundsen Sea was comparable to that in the Southern Ocean. Water-soluble organic carbon (WSOC) concentrations over the Southern Ocean and the Amundsen Sea varied from 0.048 to 0.16 and 0.070 to 0.18 µgC m−3, with averages of 0.087 ± 0.038 and 0.097 ± 0.038 µgC m−3, respectively. For water-insoluble organic carbon (WIOC), its mean concentrations over the Southern Ocean and the Amundsen Sea were 0.25 ± 0.13 and 0.26 ± 0.10 µgC m−3, varying from 0.083 to 0.49 and 0.12 to 0.38 µgC m−3, respectively. WIOC was the dominant organic carbon species in both the Southern Ocean and the Amundsen Sea, accounting for 73%–75% of the total aerosol organic carbon. WSOC/Na+ and WIOC/Na+ ratios in the fine-mode aerosol particles were higher, especially in the Amundsen Sea where biological productivity was much higher than the Southern Ocean. The fluorescence properties of water-soluble organic aerosols investigated using a fluorescence excitation–emission matrix coupled with parallel factor analysis (EEM–PARAFAC) revealed that protein-like components were dominant in our marine aerosol samples, representing 69%–91% of the total intensity. Protein-like components also showed a significant positive relationship with the relative biomass of diatoms; however, they were negatively correlated with the relative biomass of P. antarctica. These results suggest that the protein-like component is most likely produced as a result of biological processes of diatoms in the Amundsen Sea.

1 Introduction

Marine aerosols have been recognized to play an essential role in global climate due to their impact on the radiation budget and cloud microphysics by scattering solar radiation and acting as cloud condensation nuclei (CCN) (Andreae and Crutzen, 1997; O’Dowd et al., 2004). Considerable efforts have been devoted to investigating the physicochemical characteristics of marine aerosols to elucidate the source and formation of CCN in the marine environment because of their importance for understanding the cloud-mediated effects of aerosols on climate (e.g., Hegg et al., 1991; O’Dowd et al., 1997; Vallina et al., 2006; Furutani et al., 2008; Meskhidze and Nenes, 2010; Mochida et al., 2011; Gras and Keywood, 2017).
The biogeochemical cycle of sulfur between the marine atmosphere and the ocean has received much attention, especially after the proposal by Charlson et al. (1987), who postulated that the most significant source of CCN in the marine environment is non-sea-salt sulfate (nss SO$_4^{2-}$) derived from the atmospheric oxidation of dimethylsulfide (DMS). DMS produced by marine phytoplankton is the dominant sulfur species in ocean surface waters and is transported to the atmosphere through the sea-to-air flux (Bates et al., 1992). After emission to the atmosphere, DMS is oxidized by the hydroxyl (OH), nitrate (NO$_3$) and bromine oxide (BrO) radicals to form either gaseous methanesulfonic acid (MSA$_{(g)}$) or sulfur dioxide (SO$_2$). While MSA$_{(g)}$ rapidly condenses onto existing particles (forming particulate MSA, MSA$_{(p)}$), SO$_2$ is further oxidized to nss SO$_4^{2-}$ (e.g., Andreae et al., 1985; Gondwe et al., 2004; Read et al., 2008). The conversion of DMS into nss SO$_4^{2-}$ aerosols is an essential process because of the potential interaction of sulfur aerosols with incoming solar radiation and their role in cloud microphysics, which could result in a negative climate feedback mechanism (Legrand and Pasteur, 1998). However, Quinn and Bates (2011) questioned the role of sulfur-containing aerosols derived from DMS in the climate feedback, but it is still clear that DMS emissions contribute significantly to sulfur-containing aerosols acting as CCN over the oceans (Sanchez et al., 2018).

In addition to atmospheric sulfur species, marine organic aerosols have recently drawn increasing attention due to their potentially significant contribution to the CCN budget over the remote ocean (e.g., O’Dowd et al., 2004; Quinn and Bates, 2011; Gantt and Meskhidze, 2013). Ocean surface waters are enriched with small particulate organic materials including phytoplankton, bacteria, viruses, fragments of larger organisms and organic detritus (Quinn and Bates, 2011), as well as dissolved organic matter released or exuded by phytoplankton during growth, predation by grazing organisms and viral lysis (Biersmith and Benner, 1998). The organic fraction in surface waters can be broadly characterized as lipids, amino and fatty acids, monosaccharides and polysaccharides, and humic substances (Benner et al., 1992; Mochida et al., 2002; Gantt and Meskhidze, 2013), which can be emitted into the marine atmosphere with sea-salt particles through bubble bursting processes (Russell et al., 2010; Gantt et al., 2011). Previous studies have highlighted the significance of ocean-derived organic aerosol as a critical component of the aerosol–cloud–climate feedback system (e.g., Kawamura et al., 1996, 2010; Kanakidou et al., 2005; Russell et al., 2010; Miyazaki et al., 2011; Quinn and Bates, 2011; Fu et al., 2015; Wilson et al., 2015; Miyazaki et al., 2016). For example, O’Dowd et al. (2004) revealed clear differences in ocean-derived organic carbon (OC) concentrations during periods of high and low biological activity, suggesting that OC derived from biological activity can enhance the CCN concentration in the marine atmosphere and that ocean-derived OC is a significant component of the aerosol–cloud–climate feedback system involving marine biota.

Although the importance of marine biogenic source contributions to the CCN concentration has motivated numerous studies, quantitative measurements of the size-dependent composition of marine aerosols over high southern latitudes, especially the Antarctic, remain sparse due to inaccessibility. Because of the difficulty in conducting a field observation, the sources and evolution of aerosols over the Antarctic are still a subject of many open questions (Giordano et al., 2017). It is therefore necessary to fill the data gap in the knowledge of marine aerosols in the Antarctic to improve our understanding of the effect of the ocean ecosystem on the marine aerosol–cloud–climate system.

Polynyas, recurring areas of seasonally open water surrounded by sea ice in high-latitude regions, often exhibit high primary productivity (e.g., Arrigo et al., 2003, 2012; Yager et al., 2012; Hahm et al., 2014) because they are the first polar marine systems to be exposed to the increasing springtime solar radiation (Arrigo and van Dijken, 2003; Criscitiello et al., 2013). Coastal polynyas surrounding Antarctica exhibit massive phytoplankton blooms during the austral summer (December–February), with most peaking in January. The productive polynyas are located in the Ross and Amundsen seas, while annual production in the polynyas of East Antarctica is generally low (Arrigo and van Dijken, 2003). The Amundsen Sea, located in West Antarctica, hosts two coastal polynyas: the Amundsen Sea Polynya (ASP) and Pine Island Bay Polynya (PIBP) (Arrigo and van Dijken, 2003). The ASP is the most productive polynya (per unit area) among 37 identified coastal polynya systems in the Antarctic (Arrigo and van Dijken, 2003) due to the combined effects of enhanced light conditions (Park et al., 2017), the abundance of macronutrients, the supply of iron (Fe) from melting sea ice and/or glaciers, and continental shelf sediments resulting from the intrusion of relatively warm, salty and nutrient-rich Circumpolar Deep Water (CDW) (Arrigo et al., 2012; Dutrieux et al., 2014; Sherrell et al., 2015). Consequently, the Amundsen Sea is an ideal place to monitor a direct link between biological production and local emissions of sulfur compounds and organics, not only because of its remoteness from anthropogenic activity but also because it is an area of exceptionally high seasonal primary production. However, little is known about the distributions of atmospheric sulfur and organic species in the Amundsen Sea.

To understand the influence of marine biological activities on atmospheric marine aerosols in the Amundsen Sea, we have investigated the characteristics of MSA$_{(p)}$, nss SO$_4^{2-}$, water-soluble organic carbon (WSOC) and water-insoluble organic carbon (WIOC) in marine aerosols collected over the Amundsen Sea, Antarctica. We have also carried out a hydrographic survey to examine the link between biological production and atmospheric sulfur and organic species in the Amundsen Sea. The objectives of this study are to (1) investigate the distributions of MSA$_{(p)}$ and nss SO$_4^{2-}$ over the
Amundsen Sea, (2) examine the factors influencing the atmospheric MSA$_{(p)}$ concentration in the Amundsen Sea, (3) investigate the distributions of atmospheric WSOC and WIOC over the Amundsen Sea, and (4) estimate the source of atmospheric water-soluble organic aerosols using a fluorescence technique.

2 Method

Aerosol and seawater samples were collected during the ANA06B cruise conducted over the Southern Ocean and the Amundsen Sea, Antarctica, aboard the Korean icebreaker IBR/V Araon (Fig. 1). The cruise started from Christchurch, New Zealand, on 6 January 2016, sailed over the Amundsen Sea for 23 d (14 January–5 February) and returned to Christchurch, New Zealand, on 24 February 2016. In this study, the Southern Ocean and the Amundsen Sea are defined as the regions between 43 and 70° S and between 70 and 75° S, respectively. Although the cruise track covered the Southern Ocean (43–70° S) and the Amundsen Sea (70–75° S), a significant portion of our cruise was allocated to the ASP and near-ice-shelf surveys (<2 km from ice shelves) adjacent to the Dotson, the Getz and the Pine Island ice shelves.

2.1 Aerosol collection

Two high-volume aerosol samplers (HV-1000R, Sibata Scientific Technology Ltd.) were installed on the front of the upper deck (20 m a.s.l.) and used to collect marine aerosols on pre-combusted (at 550 °C per deck (20 m a.s.l.) and used to collect marine aerosols on

A wind sector controller was used to avoid contamination from the ship’s exhaust during the cruise (Jung et al., 2013, 2014). The wind sector controller was set to collect marine aerosol samples only when the relative wind directions were within plus or minus 100° relative to the ship’s bow and the relative wind speeds were over 1 m s$^{-1}$. The flow rate was 1000 L min$^{-1}$ and the total sampling time was 3–4 d, representing a total sampling air volume of 1300–4500 m$^3$. After sampling, the filters were stored frozen at −24 °C before chemical analysis. Four procedural blanks were obtained by placing quartz filters in the aerosol sampler for 5 min under idle conditions (i.e., no airflow through the filters) and processed as other aerosol samples. Meteorological variables (e.g., wind speed and wind direction) were continuously monitored by weather monitoring systems installed on the research vessel during the cruise.

2.2 Chemical analysis

2.2.1 Ionic species

The quartz filters, on which aerosols were collected, were cut into four equivalent subsamples. The subsamples were placed in acid-cleaned polypropylene bottles with the dusty side facing up; 50 mL of Milli-Q water (> 18MΩ cm$^{-1}$; Millipore Co.) was added to the bottles, and the bottles were covered using polypropylene screw caps. The subsamples were sonicated for 30 min. The extraction solution was then filtered through a 13 mm diameter, 0.45 µm pore size membrane filter (PFE syringe filter, Millipore Co.). The filtrates were analyzed by ion chromatography (IC; ICS 2000, Thermo Scientific Dionex) for anions (Cl$^-$, MSA, NO$_3^-$ and SO$_4^{2-}$) and cations (Na$^+$, NH$_4^+$, K$^+$, Mg$^{2+}$ and Ca$^{2+}$). The instrumental detection limits were the following: Cl$^-$ 0.05 µg L$^{-1}$, MSA 0.02 µg L$^{-1}$, NO$_3^-$ 0.02 µg L$^{-1}$, SO$_4^{2-}$ 0.02 µg L$^{-1}$, Na$^+$ 0.02 µg L$^{-1}$, NH$_4^+$ 0.14 µg L$^{-1}$, K$^+$ 0.16 µg L$^{-1}$, Mg$^{2+}$ 0.08 µg L$^{-1}$ and Ca$^{2+}$ 0.20 µg L$^{-1}$ (Hong et al., 2015). From replicate injections, the analytical precision was estimated to be < 5 %. The concentrations of nss SO$_4^{2-}$ were calculated as total SO$_4^{2-}$ minus Na$^+$ concentration times 0.2516, the SO$_4^{2-}$/Na$^+$ mass ratio in bulk seawater (Millero and Sohn, 1992).

2.2.2 Water-soluble organic carbon

Other subsamples of the quartz filters were ultrasonically extracted using the same method for ionic species measurements. The filtrates were analyzed by a total organic carbon (TOC) analyzer (model TOC-L, Shimadzu Inc.) for the determination of total dissolved organic carbon (TDOC), which was defined as water-soluble organic carbon (WSOC) in this study. In the analytical system, inorganic carbon was removed by acidifying the samples to pH 2 by 2 M HCl and sparging for 8 min before analysis of the total carbon content. Carbon dioxide (CO$_2$) derived from the conversion of TDOC by high-temperature (680 °C) catalytic oxidation was measured by a nondispersive infrared (NDIR) detector to quantify TDOC (Miyazaki et al., 2011). Milli-Q water and consensus reference material (CRM; 42–45 µM C for DOC; deep Florida Strait water obtained from the University of Miami) were measured every sixth analysis to check the accuracy of the measurements. The procedural mean blank for TOC was 180 µg C L$^{-1}$, which represented 14 % of the mean WSOC concentration in aerosols. The detection limit, calculated as 3 times the standard deviation of the procedural blanks, was 21 µg C L$^{-1}$. The relative standard deviation of the WSOC analysis for the reproducibility test (at least three measurements per sample) was less than 3 %.
2.2.3 Organic and elemental carbon

Concentrations of OC and elemental carbon (EC) were measured using the thermal optical transmission (TOT) method on a Sunset Laboratory OC/EC analyzer (model 4, USA) (Birch and Cary, 1996). The analytical procedures for OC and EC measurements are described in detail elsewhere (e.g., Miyazaki et al., 2011; Niu et al., 2012, 2013). In brief, a filter punch of 1.54 cm$^2$ was placed in the oven and heated in a completely pure helium environment up to 850 °C to convert all OC into CO$_2$. For EC measurement, the oven was cooled to 550 °C and then heated until the oven temperature rose back to 850 °C under a 2% oxygen-containing helium environment. All CO$_2$ derived from the conversion of both...
OC and EC was measured by an NDIR detector. The equivalent OC concentration from the field blank accounted for \( \sim 15\% \) of the average OC concentrations of the actual samples. Based on field blank uncertainties, detection limits for OC and EC were 0.1 and 0.02 \( \mu g \) \( C \) cm\(^{-2} \), respectively. In this study, WIOC was defined as the difference between OC and WSOC (i.e., WIOC = OC – WSOC) (Miyazaki et al., 2011). Using the propagating errors of each parameter, the uncertainty of the WIOC concentration was estimated to be 5.8%.

### 2.2.4 Optical measurements and excitation–emission matrix coupled with parallel factor analysis

Absorption spectra of atmospheric WSOC were obtained from 240 to 800 nm on a Shimadzu 1800 ultraviolet–visible (UV–Vis) spectrophotometer (Shimadzu Inc.). Three-dimensional fluorescence excitation–emission matrices (EEMs) were scanned using a Hitachi F-7000 luminescence spectrometer (Hitachi Inc.) at the excitation–emission (Ex–Em) wavelengths of 250–500 and 280–550 nm. The excitation and emission scans were set at 5 and 1 nm steps, respectively. The UV–Vis spectra were used for inner filter correction for the EEMs according to McKnight et al. (2001). Further details on the EEM measurements and the procedure of post-acquisition corrections are available in previous studies (Chen et al., 2010, 2017, 2018). The procedure for Raman unit (RU) normalization can be found elsewhere (Lawaetz and Stedmon, 2009). Parallel factor (PARAFAC) modeling was performed using MATLAB7.0.4 with the DOMFluor toolbox. All corrected EEMs of aerosol samples were used for modeling. The number of fluorescent components was determined based on split-half validation. The biological index (BIX), an index of recent biological and autochthonous contribution, was calculated according to Huguet et al. (2009).

### 2.3 Hydrographic data

#### 2.3.1 Chlorophyll \( a \)

**In situ measurement**

Seawater samples were collected from four to five layers in the upper 100 m at 46 stations in the Amundsen Sea using a conductivity–temperature–depth (CTD) and a rosette system holding 24–10 L Niskin bottles (Sea-Bird Electronics, SBE 911 plus) (Fig. 1). The seawater sample for chlorophyll \( a \) (Chl \( a \)) analysis was filtered through a GF/F filter (47 mm; Whatman), which was then extracted with 90 \% acetone for 24 h. Chl \( a \) was measured onboard using a fluorometer (Trilogy, Turner Designs, USA) (Lee et al., 2016).

**Satellite observation**

A monthly composite of the sea surface Chl \( a \) concentration (mg m\(^{-3} \)) was obtained from Moderate Resolution Imaging Spectroradiometer (MODIS) Aqua data available from the Goddard Space Flight Center, NASA. The spatial resolution of the data was approximately 9 km per pixel (http://oceandata.sci.gsfc.nasa.gov, last access: 30 November 2017).

#### 2.3.2 Dissolved organic carbon

A seawater sample was drawn from the Niskin bottle by gravity filtration through an inline pre-combusted (at 550 °C for 6 h) Whatman GF/F filter held in an acid-cleaned (0.1 M HCl) polycarbonate 47 mm filter holder (PP-47, ADVANTEC). The filter holder was attached directly to the Niskin bottle spigot. The filtrate was collected in an acid-cleaned glass bottle and then distributed into two pre-combusted 20 mL glass ampoules with a sterilized serological pipette. Each ampoule was sealed with a torch, quick-frozen and preserved at \( -24°C \) until analysis in our land laboratory. Analysis of dissolved organic carbon (DOC) was performed by high-temperature combustion using a Shimadzu TOC-L analyzer. Milli-Q water (blank) and consensus reference material (CRM; 42–45 \( \mu g \) C for DOC; deep Florida Strait water obtained from the University of Miami) were measured every sixth analysis to check the accuracy of the measurements. Analytical errors based on the standard deviations for replicated measurements (at least three measurements per sample) were within 5 \% for DOC.

#### 2.3.3 Particulate organic carbon

For the determination of particulate organic carbon (POC), a seawater sample was drawn from the Niskin bottle into an amber polyethylene bottle. Known volumes (500 mL–1 L) of seawater were filtered onto pre-combusted Whatman GF/F filters (25 mm) under gentle vacuum at \(< 0.1 \) MPa. The filter samples were stored at \(-80^°C \) until analysis in our land laboratory. Before POC analysis, the filter samples were freeze-dried and then exposed to HCl fumes for 24 h in a desiccator to remove inorganic carbon from the samples. Measurement of POC was carried out with a CHN elemental analyzer (vario MACRO cube, Elementar, Germany). Acetanilide was used as a standard. The precision of these measurements was \( \pm 4\% \).

#### 2.3.4 Phytoplankton taxonomic composition

Phytoplankton taxonomic composition was assessed using high-performance liquid chromatography (HPLC) analysis of accessory photosynthetic pigment concentrations. For the analysis of photosynthetic pigments, 1–2 L of seawater sample was filtered onto 47 mm Whatman GF/F filters and stored at \(-80^°C \) until analysis in our land laboratory. The filters were extracted with 3 mL of 100 \% acetone, ultrasonicated for 30 s and maintained under 4 °C in the dark for 15 h. Debris was removed by filtering through 0.45 \( \mu m \) Teflon syringe filters. Before injection, the extracts were diluted with dis-
tilled water (1 mL of extract plus 0.3 mL of distilled water) to avoid the peak distortion of the first eluting pigments. Pigments were assessed by HPLC analysis following the method of Zapata et al. (2000). Before analysis, the instrument (Agilent series 1200 chromatographic system, Germany) was calibrated using standard pigments (DHI, Denmark). A C8 column (250 mm × 4.6 mm, 5 µm particle size; Agilent XDB-C8, USA) was used for pigment separation. HPLC pigment data were processed using CHEMTAX (CHEMical TAXonomy), a matrix factorization program to calculate the absolute Chl a biomass of major algal groups. A total of 12 pigments were chosen for CHEMTAX analysis, and 7 pigment algal groups were defined according to Wright et al. (2010), including Phaeocystis antarctica (P. antarctica), diatoms and cryptophytes.

3 Results and discussion

3.1 Atmospheric MSA$_{(p)}$ over the Southern Ocean and the Amundsen Sea

The concentration of MSA$_{(p)}$ in bulk (fine + coarse) aerosols during the cruise ranged from 0.038 to 0.57 µg m$^{-3}$, with an average of 0.22 ± 0.14 µg m$^{-3}$ (mean ± 1 standard deviation). About ∼80% (median values for all data) of MSA$_{(p)}$ existed in the fine-mode aerosols. Over the Southern Ocean (43–70°S, samples A1–A2), the MSA$_{(p)}$ concentration was relatively low (mean: 0.10 ± 0.002 µg m$^{-3}$), whereas its concentration over the Amundsen Sea (70–75°S, samples A3–A9) increased sharply up to 0.35 µg m$^{-3}$ and showed high spatial variability (range: 0.096–0.57 µg m$^{-3}$, mean: 0.28 ± 0.15 µg m$^{-3}$), with the highest value observed in the ASP (Fig. 2a). As expected, however, the MSA$_{(p)}$ concentration gradually decreased from 0.29 to 0.038 µg m$^{-3}$ with distance from the ASP (i.e., from the Amundsen Sea to the Southern Ocean; samples A10–A14, mean: 0.18 ± 0.11 µg m$^{-3}$). The MSA$_{(p)}$ concentration from this study was in reasonable agreement with previously published results obtained over the regions near the Antarctic continent during the austral summer when air masses originate from the western Antarctic coastal regions (60–70°S, 0.05–0.26 µg m$^{-3}$; Chen et al., 2012). However, our mean MSA$_{(p)}$ concentration in the Amundsen Sea was about 1.8–4.4 times higher than those observed at Antarctic research stations during the austral summer, such as Palmer Station (64.77°S, 64.05°W; 0.122 ± 0.127 µg m$^{-3}$; December 1990–March 1991; Savoie et al., 1993), Halley Station (75°35′S, 26°19′W; 0.14 µg m$^{-3}$, January–February 2005; Read et al., 2008), Neumayer Station (70°39′S, 8°15′W; 0.154 ± 0.077 µg m$^{-3}$, monthly mean in January from 1983 to 1995; Minikin et al., 1998) and Dumont d’Urville Station (66°40′S, 140°1′E; 0.063 ± 0.019 µg m$^{-3}$, monthly mean in January from 1991 to 1995; Minikin et al., 1998). Although the mean MSA$_{(p)}$ concentration was much higher than those observed at the Antarctic research stations, the maximum MSA$_{(p)}$ in the Amundsen Sea was comparable to the highest concentration measured at Palmer Station (0.57 µg m$^{-3}$; Savoie et al., 1993), implying that the source strength for MSA$_{(p)}$ is greater in the Amundsen Sea. These differences in atmospheric MSA$_{(p)}$ concentrations among coastal Antarctic regions presumably reflect regional differences in factors controlling MSA$_{(p)}$ abundance in the marine atmosphere, such as spatial variations of phytoplankton bloom, phytoplankton taxonomic composition, emission flux of DMS, atmospheric oxidative capacity, and atmospheric transport and removal (Minikin et al., 1998; Gondwe et al., 2004; Chen et al., 2012; Criscitiello et al., 2013; Jung et al., 2014).

Figure 2. Concentrations of particulate methanesulfonic acid (MSA$_{(p)}$) (a) and non-sea-salt sulfate (nss SO$_4^{2-}$) (b) against sample ID in aerosols collected over the Southern Ocean (43–70°S) and the Amundsen Sea (70–75°S). The black solid lines with circles indicate the latitude of the halfway point between each aerosol sampling start point and endpoint. Double arrows show the sampling locations of aerosol samples.
3.2 Factors influencing atmospheric MSA\(_{(p)}\) concentration over the Amundsen Sea

Various factors appear to influence the atmospheric MSA\(_{(p)}\) concentration in the Amundsen Sea. According to Arrigo and van Dijken (2003), the 5-year (1997–2002) average Chl \(a\) level in the ASP during the month of January (6.98 ± 3.32 mg m\(^{-3}\)) is more than 2 times higher than that in the Ross Sea Polynya (2.67 ± 0.82 mg m\(^{-3}\)). Indeed, satellite ocean color images (http://oceancolor.sci.gsfc.nasa.gov, last access: 30 November 2017) exhibited persistently high Chl \(a\) levels in the ASP during the sampling period (Fig. 3), implying a strong influence of biogenic sources on atmospheric MSA\(_{(p)}\) in the Amundsen Sea.

In addition to the high productivity, phytoplankton taxonomic assemblages could be a significant factor influencing the atmospheric MSA\(_{(p)}\) abundance. In general, phytoplankton taxonomic abundances in the Amundsen Sea are...
dominated by the haptophyte *P. antarctica* (e.g., Lee et al., 2016; Yager et al., 2016), which produces large amounts of DMS in Antarctic waters (Liss et al., 1994; Schoemann et al., 2005). During the cruise, *P. antarctica* was the dominant phytoplankton species in the upper 50 m, accounting for 42 ± 19 % of phytoplankton biomass (Chl a), with lesser abundances of diatoms (39 ± 17 %) found throughout the polynya and sea ice zone (Fig. S1 in the Supplement). In addition, extremely high concentrations (> 150 nM) and fluxes (85 ± 119 µmol m⁻² d⁻¹) of DMS have been observed in the region where phytoplankton assemblages were dominated by *P. antarctica* during the cruise (Kim et al., 2017), which is consistent with previous results observed in the Amundsen Sea (Tortell et al., 2012).

We investigated the relationships between the atmospheric MSAₚ concentration and the variables mentioned above to examine factors influencing the atmospheric MSAₚ concentration in the Amundsen Sea. The atmospheric MSAₚ concentration showed no relationship with either the in situ sea surface Chl a concentration (r = 0.029, p > 0.05) (Fig. S2) or the relative biomass of *P. antarctica* (r = 0.30, p > 0.05). This suggests that the Chl a concentration and phytoplankton taxonomic composition are not direct factors determining the atmospheric MSAₚ concentration over the Amundsen Sea.

To investigate the relationship between atmospheric MSAₚ and the DMS flux, we used the DMS flux data reported by Kim et al. (2017), who calculated sea–air DMS fluxes using sea surface DMS concentrations and shipboard wind speed data monitored during our cruise. Details of the measurement of the sea surface DMS concentration and the sea–air DMS flux calculation are given by Kim et al. (2017). The DMS flux (reported by Kim et al., 2017) averaged for the duration of each aerosol sampling showed a somewhat similar variation trend to that of the atmospheric MSAₚ concentration (Fig. 4a), but no correlation was found between atmospheric MSAₚ and the DMS flux (r = 0.18, p > 0.05; Fig. 4b). DMS fluxes typically rely on gas transfer velocity (k), which is frequently parameterized as a function of wind speed (Wanninkhof, 2014). The measurement and parameterization of the gas transfer velocity are more challenging and subject to greater uncertainty, particularly at high wind speeds (Smith et al., 2018). Wanninkhof et al. (2014) reported that there is considerable uncertainty in k, especially under strong wind conditions. About 20 % uncertainty was estimated at global mean wind speed (7.3 m s⁻¹). When we applied four different k values (units of k: cm h⁻¹) calculated from the equations suggested by Wanninkhof (1992), Wanninkhof and McGillis (1999), Nightingale et al. (2000), and Wanninkhof (2014), respectively, to the equation for the sea–air DMS flux calculation reported by Kim et al. (2017), the uncertainty in the DMS flux in the Amundsen Sea region was about 25 % (1 standard deviation of the four different mean DMS fluxes). As the gas transfer velocity increases with increasing wind speed, the DMS flux can be overestimated, especially in higher latitudes where DMS is commonly found at high concentrations in surface water and where both low temperatures and high winds are typical (McGillis et al., 2000). In addition to the uncertainty in the DMS flux, the insignificant relationship between atmospheric MSAₚ and the DMS flux could result from various complexities in the rate of oxidation of DMS to form atmospheric MSAₚ and long-range transport of atmospheric MSAₚ from biogenically active regions given that the lifetime of DMS is approximately 1–2 d (Kloster et al., 2006; Read et al., 2008).
Although we found no significant relationship between the atmospheric MSA\(_{(p)}\) concentration, in situ sea surface Chl \(a\) concentration, the relative biomass of *P. antarctica* and the local sea–air DMS flux, the higher atmospheric MSA\(_{(p)}\) concentrations observed over the Amundsen Sea compared to those over the Southern Ocean and in coastal Antarctic regions most likely resulted from a complex linkage between these factors.

### 3.3 Atmospheric nss SO\(_4^{2-}\) over the Southern Ocean and the Amundsen Sea

The concentration of nss SO\(_4^{2-}\) in bulk (fine + coarse) aerosols during the cruise ranged from 0.30 to 0.87 \(\mu g\) m\(^{-3}\), with \(\sim 79\%\) (median values for all data) of nss SO\(_4^{2-}\) being present on fine-mode aerosols (Fig. 2b). The mean concentration of nss SO\(_4^{2-}\) during the cruise was 0.61 \(\pm\) 0.17 \(\mu g\) m\(^{-3}\). Over the Southern Ocean (43–70° S; samples A1–A2 and A11–A14), the nss SO\(_4^{2-}\) concentration ranged from 0.30 to 0.87 \(\mu g\) m\(^{-3}\) (mean: 0.60 \(\pm\) 0.23 \(\mu g\) m\(^{-3}\)), whereas its concentration over the Amundsen Sea (70–75° S; samples A3–A10) varied from 0.49 to 0.81 \(\mu g\) m\(^{-3}\) (mean: 0.62 \(\pm\) 0.10 \(\mu g\) m\(^{-3}\)). Surprisingly, our mean nss SO\(_4^{2-}\) concentration in the Amundsen Sea was about 1.5 and 2.5 times higher than those observed at American Samoa (14.25° S, 170.58° W; 0.41 \(\pm\) 0.17 \(\mu g\) m\(^{-3}\), seasonal average for December–February from 1990 to 1992; Savoie et al., 1994) and over the South Pacific (8–55° S; mean: 0.25 \(\pm\) 0.17 \(\mu g\) m\(^{-3}\), range: 0.094–0.62 \(\mu g\) m\(^{-3}\), January–March 2009; Jung et al., 2014), respectively. In addition, the mean nss SO\(_4^{2-}\) concentration in the Amundsen Sea was also a factor of 1.6–4.4 higher than those observed at Palmer Station (0.24 \(\pm\) 0.16 \(\mu g\) m\(^{-3}\), December 1990–March 1991; Savoie et al., 1993), Halley Station (0.14 \(\pm\) 0.017 \(\mu g\) m\(^{-3}\), monthly mean in January from 1991 to 1993; Legrand and Pasteur, 1998), Neumayer Station (0.38 \(\pm\) 0.13 \(\mu g\) m\(^{-3}\), monthly mean in January from 1993 to 1995; Minikin et al., 1998) and Dumont d’Urville Station (0.34 \(\pm\) 0.039 \(\mu g\) m\(^{-3}\), monthly mean in January from 1991 to 1995; Minikin et al., 1998) during the austral summer.

Unlike MSA\(_{(p)}\), the mean nss SO\(_4^{2-}\) concentration in the Amundsen Sea was comparable to that in the Southern Ocean, although the variation trend of nss SO\(_4^{2-}\) in the Amundsen Sea was similar to that of MSA\(_{(p)}\), suggesting that nss SO\(_4^{2-}\) was affected by marine sources and large-scale transport (Korhonen et al., 2008). Indeed, nss SO\(_4^{2-}\) showed a strong correlation (\(r = 0.98, p < 0.01\)) with MSA\(_{(p)}\) in the Amundsen Sea, whereas no relationship was found between them in the Southern Ocean (\(r = 0.51, p > 0.05\)) (Fig. S3), suggesting that the local emission of DMS is a significant source of nss SO\(_4^{2-}\) in the Amundsen Sea. It is worth mentioning that nss SO\(_4^{2-}\) can be formed from the homogeneous nucleation of new particles involving H\(_2\)SO\(_4\) or from the condensation of gas-phase DMS oxidation products onto existing particles (e.g., Covert et al., 1992; Quinn and Bates, 2011). Recently, Sanchez et al. (2018) identified two types of SO\(_4^{2-}\) particles using an event-trigger aerosol mass spectrometer (ET-AMS) and reported that 63\% of SO\(_4^{2-}\) was derived from newly formed particles in the free troposphere and 38\% of SO\(_4^{2-}\) was formed from the condensation of DMS products onto existing particles in clean marine conditions, revealing the importance of phytoplankton-produced DMS emissions for CCN in the Atlantic. In this study, it is hard to distinguish nss SO\(_4^{2-}\) derived from new particle formation from nss SO\(_4^{2-}\) formed by the condensation of DMS products onto existing particles because of the limitations related to the method to collect data. However, the significant correlation of nss SO\(_4^{2-}\) with MSA\(_{(p)}\) in the Amundsen Sea suggests that DMS emissions from the Amundsen Sea play a crucial role in the formation of nss SO\(_4^{2-}\) aerosol particles that can act as CCN.

On the other hand, a possible explanation for the insignificant relationship between nss SO\(_4^{2-}\) and MSA\(_{(p)}\) in the Southern Ocean is the entrainment of nucleated nss SO\(_4^{2-}\) particles into the marine boundary layer of the Southern Ocean from the free troposphere by turbulent diffusion and large-scale transport (Korhonen et al., 2008; Woodhouse et al., 2010). Sanchez et al. (2018) reported that the lack of correlation between SO\(_4^{2-}\) particles and atmospheric DMS (or its oxidation products) could result from the competition for DMS and its oxidation products with the competing sinks of condensation onto existing particles and vertical transport to the free troposphere. The lack of correlation between nss SO\(_4^{2-}\) and MSA\(_{(p)}\) in the Southern Ocean could therefore have resulted from the input of nss SO\(_4^{2-}\) from the free troposphere. Although our dataset is not sufficiently complete to allow for a meaningful analysis of this likely explanation, the result for nss SO\(_4^{2-}\) concentrations from this study could be valuable for filling the data gap, especially for the Amundsen Sea during the austral summer, and could also be helpful for the validation of the modeling of sulfur-containing aerosols.

### 3.4 MSA\(_{(p)}\) / nss SO\(_4^{2-}\) ratios over the Southern Ocean and the Amundsen Sea

There have been several field studies investigating the MSA\(_{(p)}\) / nss SO\(_4^{2-}\) ratio in the Southern Ocean during the austral summer (range: 0.32–0.53 in the South Pacific at 40–45° S, Berresheim et al., 1990; range: 0.12–0.24 for Cape Grim at 40°41’S, 144°41’E in Ayers et al., 1991; range: 0.17–0.32 in the South Pacific at 30–60° S, Bates et al., 1992; range: 0.096–0.49 in the South Pacific at 40–56° S, Jung et al., 2014).

During the cruise, the MSA\(_{(p)}\) / nss SO\(_4^{2-}\) ratio in bulk aerosols varied from 0.12 to 0.70 (mean: 0.35 \(\pm\) 0.17), with lower ratios in marine aerosols collected over the Southern Ocean (range: 0.12–0.51, mean: 0.26 \(\pm\) 0.14) and higher values over the Amundsen Sea (range: 0.20–0.70, mean: 0.44 \(\pm\) 0.16), showing a similar variation trend (\(r = 0.92,0.98,\)).
Figure 5. MSA\(_{(p)}\) / nss SO\(_4^{2-}\) ratio against aerosol sample ID. Double arrows show the sampling locations of aerosol samples. The black solid line with circles indicates the latitude of the halfway point between each aerosol sampling start point and endpoint.

\(p < 0.01\) to that of MSA\(_{(p)}\) (Fig. 5). This result suggests that the atmospheric MSA\(_{(p)}\) plays a key role in the variation in MSA\(_{(p)}\) / nss SO\(_4^{2-}\) ratio over the Southern Ocean and the Amundsen Sea during the austral summer since atmospheric nss SO\(_4^{2-}\) concentrations in the Southern Ocean were quite comparable to the values in the Amundsen Sea.

3.5 Atmospheric WSOC and WIOC over the Southern Ocean and the Amundsen Sea

The concentration of WSOC in bulk (fine + coarse) aerosols during the cruise ranged from 0.048 to 0.18 µgC m\(^{-3}\), with an average of 0.092 ± 0.037 µgC m\(^{-3}\) (Fig. 6a). The WSOC concentrations over the Southern Ocean (43–70° S; samples A1–A2 and A11–A14) and the Amundsen Sea (70–75° S; samples A3–A9) varied from 0.048 to 0.16 and 0.070 to 0.18 µgC m\(^{-3}\), with averages of 0.087 ± 0.038 and 0.097 ± 0.038 µgC m\(^{-3}\), respectively. For WIOC, its mean concentrations over the Southern Ocean and the Amundsen Sea were 0.25 ± 0.13 and 0.26 ± 0.10 µgC m\(^{-3}\), varying from 0.083 to 0.49 and 0.12 to 0.38 µgC m\(^{-3}\), respectively (Fig. 6b). We expected much higher WSOC and WIOC concentrations in the Amundsen Sea than the Southern Ocean because of the extremely high Chl \(a\) concentration in the Amundsen Sea (Fig. 3). However, no significant differences in mean WSOC and WIOC concentrations were found between the Southern Ocean and the Amundsen Sea, suggesting that the Chl \(a\) concentration is not a direct factor controlling the atmospheric OC concentration in our study area (Quinn et al., 2014), although a significant correlation between atmospheric OC and Chl \(a\) concentrations was observed in the Austrl Ocean (Amsterdam Island; 37°48′ S, 77°34′ E; Sciare et al., 2009).

Figure 6. Concentrations of water-soluble organic carbon (WSOC) (a), water-insoluble organic carbon (WIOC) (b) and organic carbon (i.e., OC = WSOC + WIOC) (c) against sample ID in aerosols collected over the Southern Ocean (43–70° S) and the Amundsen Sea (70–75° S). Black solid lines with circles indicate the latitude of the halfway point between each aerosol sampling start point and endpoint. Double arrows in (a) show the sampling locations of aerosol samples.
Both WSOC and WIOC mainly existed in fine-mode particles, and the percentages of WSOC and WIOC present in fine aerosol particles were ~ 93% and ~ 74%, respectively (median value for all data). During the cruise, WIOC was the dominant OC species in both the Southern Ocean and the Amundsen Sea, accounting for 75% and 73% of total aerosol organic carbon, respectively (Fig. 6c). These results were consistent with previous studies. For example, O’Dowd et al. (2004) observed a dominant water-insoluble OC fraction (~45%) in fine marine aerosol collected during periods of phytoplankton bloom in the North Atlantic. Moreover, Facchini et al. (2008a) reported that OC observed through a bubble bursting experiment and a field measurement (at Mace Head) was mainly water-insoluble, accounting for 77 ± 5% of the primary marine aerosol fraction in the submicron size range, and that WIOC consisted of colloids and aggregates exuded by phytoplankton. Given that atmospheric WIOC is mechanically produced through bubble bursting processes from hydrophobic organic matter that accumulates in the ocean surface (Facchini et al., 2008a; Gantt et al., 2011; Miyazaki et al., 2011), the dominance of WIOC suggests that water-insoluble organic matter exuded by phytoplankton is more accumulated in sea surface water and emitted into the marine atmosphere via bubble bursting and breaking waves by local wind.

Our mean values of WSOC and WIOC in the Amundsen Sea were comparable to the results by Sciare et al. (2009), who reported that the mean concentrations of WSOC and WIOC observed at Amsterdam Island (37°48’S, 77°34’E) during the austral summer (January) were 0.083 ± 0.028 and 0.19 ± 0.062 µgC m⁻³, respectively. It is worth noting that atmospheric WSOC and WIOC show seasonal variations, with maximum values during austral summer (January) and minimum concentrations during winter (Sciare et al., 2009). These variations are attributable to pronounced seasonal variations in biogenic marine productivity. Given that our study was carried out during the austral summer, the concentrations of WIOC and WSOC from this study would be considered maximum values in the Amundsen Sea. Although our results were supported by previous studies as mentioned above, spatial variability in WSOC and WIOC concentrations has been observed over various oceanic regions by previous studies. For instance, Fu et al. (2015) reported that atmospheric OC species concentrations observed at Alert in the Canadian High Arctic were 0.186 µgC m⁻³ (range: 0.041–0.30 µgC m⁻³) for WSOC and 0.068 µgC m⁻³ (range: 0.022–0.12 µgC m⁻³) for WIOC. The results by Fu et al. (2015) were 1.9 times higher and 3.8 times lower than our mean concentrations of WSOC and WIOC in the Amundsen Sea, respectively. In addition, Miyazaki et al. (2011) reported that mean concentrations of WSOC and WIOC observed over the western North Pacific (40°–44°N) were 0.65 ± 0.27 and 0.56 ± 0.19 µgC m⁻³, which were a factor of 6.7 and 2.2 higher than those in the Amundsen Sea, respectively. These differences in WSOC and WIOC concentrations among the oceanic regions presumably reflect regional differences in factors influencing atmospheric WSOC and WIOC concentrations, such as source strength for volatile organic compounds emitted from biogenic sources (BVOCs), atmospheric oxidative capacity (e.g., OH, NO₃ and ozone), meteorological conditions (e.g., wind speed), DOC and POC concentrations in sea surface water, and atmospheric transport and removal (e.g., Facchini et al., 1999; Kanakidou et al., 2005; Sun et al., 2006).

3.6 WIOC/Na⁺ and WSOC/Na⁺ ratios and relationships of WIOC and WSOC with Na⁺ over the Southern Ocean and the Amundsen Sea

Breaking waves on the ocean surface generate air bubbles that scavenge organic matter from seawater. When injected into the atmosphere, these bubbles burst, yielding sea spray aerosols enriched in organic matter relative to seawater (Quinn et al., 2014). Sea spray aerosols have been defined as the hydrated droplets encapsulating dissolved sea salt and entrained organic matter (O’Dowd et al., 2008). Previous studies have revealed that organic matter is enriched in sea spray aerosol particles produced by bubble bursting processes in the fine and ultrafine aerosol size fractions, suggesting that sea spray aerosol particles have an important role in transferring organic matter from the sea surface to the atmosphere (Facchini et al., 2008a; O’Dowd et al., 2008). During the cruise, ~76% of Na⁺, a tracer of sea spray, was associated with coarse-mode particles (Fig. 7a). Moreover, statistically significant relationships were found between mean wind speed and Na⁺ concentrations in fine-mode (r = 0.54, p < 0.05) and coarse (r = 0.69, p < 0.01) aerosols, reflecting the fact that Na⁺ was formed from bubble bursting by local wind. Although Na⁺ was predominantly associated with coarse-mode particles, WSOC/Na⁺ and WIOC/Na⁺ in the fine-mode aerosol particles were higher than those in the coarse-mode aerosol particles, especially in the Amundsen Sea where biological productivity was much higher than the Southern Ocean (Fig. 7b). In the Southern Ocean, the WSOC/Na⁺ ratio in the fine-mode particles varied from 0.045 to 0.40, whereas the WSOC/Na⁺ ratio in the Amundsen Sea ranged from 0.17 to 0.89. The average WSOC/Na⁺ ratio in the Amundsen Sea (0.40 ± 0.24) was substantially higher than that in the Southern Ocean (0.16 ± 0.12). For the WIOC/Na⁺ ratio in the fine-mode particles, similar results were observed. The WIOC/Na⁺ ratios in the Southern Ocean and the Amundsen Sea varied from 0.38 to 0.97 and 0.26 to 2.4, with averages of 0.35 ± 0.31 and 0.91 ± 0.73, respectively; however, the WIOC/Na⁺ ratio in the fine-mode aerosol particles was much higher than WSOC/Na⁺, suggesting that bubble bursting at the ocean surface is a major source of atmospheric WIOC and that WIOC is more accumulated in the sea surface water. These results show that the higher marine biological activities in the Amundsen Sea can be a significant factor leading to higher OC/Na⁺ ratios, in-
indicating a linkage between OC emissions and biological activities (Fig. 3).

Because sea spray aerosols are emitted as a result of wind-driven bubble bursting, correlations of organic aerosol mass concentrations with sea spray aerosols (i.e., Na$^+$), whose concentrations are related to local wind speeds, have been used to attribute their origin to marine sources because submicron Na$^+$ is known to form primary aerosols from evaporating seawater droplets (Russell et al., 2010). In this study, we investigated relationships of WIOC and WSOC concentrations with the Na$^+$ concentration in the fine modes since both WIOC and WSOC were primarily associated with the fine-mode particles (Fig. 6a and b). The submicron WIOC showed no statistically significant relationships with submicron Na$^+$ over the Southern Ocean and the Amundsen Sea (Fig. 8a), although the WIOC/Na$^+$ ratio in the fine-mode aerosol particles was much higher (Fig. 7b). Similarly, Boreddy et al. (2018) found no correlation between sea salt and WIOC in the western North Pacific. The increase in sea-salt particle flux under higher wind speed conditions shifts the sea spray aerosol size distribution towards larger sizes and accelerates their dry deposition and gravitational settling from the atmosphere (de Leeuw et al., 2011). Thus, these insignificant relationships between WIOC and Na$^+$ in the fine modes could result from the differences in local wind speeds and local biological activities because wind speed, a key factor determining sea spray aerosols, controls the local flux rather than the local concentration of marine particles (Monahan and O’Muircheartaigh, 1986). Our results are further supported by the study of Ceburnis et al. (2008), who found that WIOC and sea salt exhibited upward fluxes observed through gradient flux measurements, suggesting a primary production mechanism for WIOC. Although we found no significant relationships between WIOC and Na$^+$ in the fine modes, the high WIOC/Na$^+$ ratio in the fine-mode aerosol particles (Fig. 7b) indicates that WIOC was predominantly of primary origin (Ceburnis et al., 2008). However, WIOC production by secondary processes cannot be completely excluded (Ceburnis et al., 2016), but we have no evidence of that.

Unlike the relationship between WIOC and Na$^+$, submicron WSOC showed a strong correlation ($r = 0.94$, $p < 0.01$) with submicron Na$^+$ in the Amundsen Sea (Fig. 8b). In addition, we also found a significant correlation ($r = 0.93$, $p < 0.01$) between WSOC and MSA$_{(p)}$ concentrations in the Amundsen Sea (Fig. 8c). However, in the Southern Ocean, WSOC showed no significant relationship with submicron Na$^+$ or MSA$_{(p)}$. MSA$_{(p)}$ is produced by the atmospheric oxidation of DMS, which is released as a gas phase from marine biological activities and thus can be used as an indicator of secondary aerosols of marine biogenic origin (Miyazaki et al., 2011). Consequently, the strong correlations between WSOC, Na$^+$ and MSA$_{(p)}$ in the Amundsen Sea implies that the Amundsen Sea, which has the most productive polynya in the Antarctic, is a strong source region of BVOCs and that WSOC was formed by the condensation of BVOCs released from the sea surface onto preexisting submicron sea spray aerosols through gas-to-particle conversion due to a higher surface-to-volume ratio of submicron aerosols (Romakkaniemi et al., 2011). On the other hand, the poor correlations between WSOC, Na$^+$ and MSA$_{(p)}$ in the Southern Ocean implies differences in the local source strength of BVOCs and that the presence of DMS in seawater and its subsequent oxidation to MSA$_{(p)}$ were not necessarily linked to the formation of submicron WSOC over the Southern Ocean (Miyazaki et al., 2016).
3.7 Fluorescence properties of water-soluble organic aerosols over the Southern Ocean and the Amundsen Sea

A fluorescence excitation–emission matrix coupled with parallel factor analysis (EEM–PARAFAC) has been widely used to investigate the sources and optical properties of dissolved organic matter in terrestrial and oceanic systems (e.g., Coble, 1996, 2007; Stedmon et al., 2003; Yamashita et al., 2011; Retelletti Brogi et al., 2018). Moreover, recent field studies demonstrated that EEM–PARAFAC provides useful information for characterizing atmospheric OC in aerosols and rainwater (e.g., Fu et al., 2015; Miyazaki et al., 2018; Yang et al., 2019).

As described in Sect. 3.6, our results strongly suggested that the submicron WSOC observed in the Amundsen Sea was formed by the condensation of BVOCs onto preexisting submicron sea spray aerosols, showing strong correlations with Na$^+$ and MSA$_{(p)}$. To further elucidate the sources of water-soluble organic aerosols, we investigated the fluorescence properties of submicron aerosols using EEM–PARAFAC. Fluorophores in water-soluble organic aerosols were divided into three primary types on the basis of their peak position (Fig. 9). The spectral features of component 1 (C1, Ex–Em: 300 and 344 nm) were similar to the component previously identified in coastal and oceanic waters as well as the polar ocean and was thought to be a phytoplankton-derived (or ice algae-derived) protein-like component (Stedmon et al., 2007, 2011, Retelletti Brogi et al., 2018). Component 2 (C2, Ex–Em: 276 and 326 nm) was assigned as a tryptophan-like fluorophore, which has been considered to be a labile component produced as a result of biological production in marine environments (Coble et al., 1998; Yamashita and Tanoue 2003). Component 3 (C3, Ex–Em: <260 and 458 nm) spectra were characterized as representing terrestrial humic-like fluorophores (Coble et al., 1996; Yamashita and Tanoue, 2003; Chen et al., 2010, 2018). During the cruise, the C1 fluorescence intensity was much higher and variable, varying between 0.0133 and 0.139 RU (Fig. 10a). In comparison, the fluorescence intensity of C2 (range: 0.0195–0.0518 RU) and C3 (range: 0.00857–0.0351 RU) was much less variable. Among the three components, the protein-like C1 was the dominant fluorescence component in our marine aerosol samples, accounting for 31 %–73 % of the total intensity, and the relative contributions of tryptophan-like C2 and terrestrial humic-like C3 were found to represent 17 %–50 % and 8 %–31 %, respectively (Fig. 10b). In our marine aerosol samples, protein-like components (i.e., C1 and C2) represented 69 %–91 % of the total intensity.

Despite the extremely high Chl $a$ concentration in the Amundsen Sea (Fig. 3), we found no significant difference in the average values of protein-like C1 and tryptophan-like C2 fluorescence intensity between the Amundsen Sea and the Southern Ocean. However, relatively much higher values of C1 and C2 fluorescence intensity were observed when
Figure 9. EEM contour plots of the three fluorescent components C1–C3 (a–c) identified using EEM–PARAFAC in the fine-mode aerosol particles \((D < 2.5 \, \mu m)\) collected over the Southern Ocean and the Amundsen Sea, as well as the excitation and emission spectra of C1–C3 (d–f).

Figure 10. Variations of fluorescence intensities (a), relative contributions of C1–C3 and biological index (black solid square line) (b) in the fine-mode aerosol particles \((D < 2.5 \, \mu m)\) collected over the Southern Ocean and the Amundsen Sea. The black solid line with circles in (a) indicates the latitude of the halfway point between each aerosol sampling start point and endpoint. Double arrows in (a) show the sampling locations of aerosol samples. Panels (c) and (d) show the relationship of the fluorescence intensity of C1 to the relative biomass of diatoms and Phaeocystis antarctica \((P. \, \text{antarctica})\) observed in the Amundsen Sea, respectively.
the ship approached the Amundsen Sea (i.e., samples A3 and A4), passing through the sea ice zone (Figs. 1b and 10a). The C1 and C2 fluorescence intensity values sharply decreased and remained relatively low in the Amundsen Sea and then gradually increased from the Amundsen Sea to the Southern Ocean. As mentioned in Sect. 3.2, *P. antarctica* was the dominant phytoplankton species in the Amundsen Sea, whereas diatoms formed a major group in the marginal sea ice zone (Lee et al., 2016).

Ice algae, commonly found in polar sea ice and surrounding waters, are largely dominated by diatoms (Roberts et al., 2007), which are an important contributor to aerosols by emitting aerosol-forming volatile (e.g., alkyl-amines) and non-volatile (e.g., mycosporine-like amino acids) organic nitrogen in the Antarctic sea ice region (Dall’Osto et al., 2017). Previous studies (e.g., Facchini et al., 2008b; Miyazaki et al., 2011) provided the evidence that volatile emissions of alkylamine from marine algae can represent an important source of marine secondary organic aerosol. Moreover, Dall’Osto et al. (2017) observed that the fluorescence signal for the protein-like component was positively correlated with organic nitrogen originating from melted Antarctic sea ice floes, indicating that the protein-like component was associated with organic nitrogen derived from the microbiota of sea ice and the sea-ice-influenced ocean. Although we have no aerosol water-soluble organic nitrogen dataset, our results provide additional evidence that marine algae can influence the fluorescent property of marine aerosols. Interestingly, we found that the fluorescence intensity of C1 showed a significant positive relationship with the relative biomass of diatoms (*r* = 0.89, *p* < 0.01); however, it was negatively correlated with the relative biomass of *P. antarctica* (*r* = −0.79, *p* < 0.05) (Fig. 10c and d). Given the dominance of diatoms in the marginal sea ice zone during the cruise and the significant positive relationship between the fluorescence intensity of protein-like C1 and the relative biomass of diatoms, it is plausible that the biological processes of diatoms are an important factor in controlling the abundance of the protein-like component in water-soluble organic aerosols over the Southern Ocean and the Amundsen Sea, although further studies are necessary to clarify this point.

The biological index (BIX) has been used to estimate the contribution of autochthonous biological activity (Fu et al., 2015; Miyazaki et al., 2018). An increase in BIX is related to an increase in the contribution of microbially derived organics. High BIX values (> 1) have been shown to correspond to a predominantly biological or microbial origin of dissolved organic matter and to the presence of organic matter freshly released into water, whereas low values (< 0.6) indicate little biological material (Huguet et al., 2009). In this study, the BIX values ranged from 1.17 to 3.61, with an average of 2.23 ± 0.807 (Fig. 10b). The high BIX values also support the idea that the fluorescence properties of WSOC were influenced by marine biological activities.

4 Conclusions

Characteristics of atmospheric sulfur (i.e., MSA$_{(p)}$ and nss SO$_4^{2-}$) and OC (i.e., WSOC and WIOC) species in marine aerosols, and the environmental factors influencing their distributions, were investigated over the Southern Ocean and the Amundsen Sea during the austral summer. In the Amundsen Sea, the atmospheric MSA$_{(p)}$ concentration drastically increased (up to 0.57 µg m$^{-3}$), suggesting significant influences of marine biological activities on atmospheric MSA$_{(p)}$. The higher MSA$_{(p)}$ concentration was attributed to exceptionally high seasonal primary production during the austral summer, the dominance of *P. antarctica* with respect to phytoplankton biomass and extremely high DMS concentrations produced by DMS-producing algae species (e.g., *P. antarctica*) in the Amundsen Sea.

WIOC was the dominant OC species in both the Southern Ocean and the Amundsen Sea, accounting for 75 % and 73 %, respectively. Despite the extremely high Chl a concentration in the Amundsen Sea, no significant differences in mean WSOC and WIOC concentrations were found between the Southern Ocean and the Amundsen Sea. However, higher WSOC/Na$^+$ and WIOC/Na$^+$ ratios were observed in the submicron aerosol particles, especially in the Amundsen Sea where biological productivity was much higher than the Southern Ocean.

It is worth noting that the simultaneous measurements of chemical species in marine aerosols as well as the chemical and biological properties of seawater in the Amundsen Sea allowed for a better understanding of the effect of the ocean ecosystem on OC species. Moreover, the fluorescence properties of water-soluble organic aerosols revealed that protein-like components are most likely produced as a result of the biological processes of diatoms.

West Antarctica is one of the fastest-warming regions globally (Bromwich et al., 2013). Ice shelves and glaciers in the Amundsen Sea have been shrinking at a remarkable rate (Rignot et al., 2008). Moreover, sea ice coverage is decreasing fast in the western Antarctic (Stammerjohn et al., 2012). Because ocean buoyancy, stratification and trace metal distribution are affected by these changes, the regional oceanography, phytoplankton community structure and biogeochemical cycles of sulfur and carbon in the Amundsen Sea are likely affected as well (Yager et al., 2012). Further studies are therefore required to more clearly understand the biogeochemical cycles of sulfur and carbon between the ocean and the marine atmosphere; they should focus on the long-term monitoring of atmospheric sulfur and OC species in the Amundsen Sea.
Data availability. The data used in this study are available on request to the correspondence author Jinyoung Jung (jinyoungjung@kopri.re.kr).

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Author contributions. JJ designed the research, carried out the experiments, processed the data and wrote the paper. SBH, MC and LJ analyzed the aerosol samples. YL and EJY provided the marine biological data. JOC and JP helped in obtaining satellite products. JH, KP, DH and EJY contributed to the scientific discussion and paper correction. TWK and SHL organized the field campaign.

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

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