Low-volatility vapors and new particle formation over the Southern Ocean during the Antarctic Circumnavigation Expedition

Andrea Baccarini¹, Josef Dommen², Katrianne Lehtipalo³, Silvia Henning⁴, Robin L Modini², Martin Gysel-Beer², U. Baltensperger², and Julia Schmale⁵

¹École polytechnique fédérale de Lausanne - EPFL
²Paul Scherrer Institute
³University of Helsinki
⁴Institute for Tropospheric Research
⁵École Polytechnique Fédérale de Lausanne

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Abstract

During summer, the Southern Ocean is largely unaffected by anthropogenic emissions, which makes this region an ideal place to investigate marine natural aerosol sources and processes. A better understanding of natural aerosol is key to constrain the preindustrial aerosol state and reduce the aerosol radiative forcing uncertainty in global climate models. We report the concentrations of gaseous sulfuric acid, iodic acid, and methanesulfonic acid (MSA) together with a characterization of new particle formation (NPF) events over a large stretch of the Southern Ocean. Measurements were conducted on board the Russian icebreaker *Akademik Tryoshnikov* from January to March 2017. Iodic acid is characterized by a particular diurnal cycle with reduced concentration around noon, suggesting a lower formation yield when solar irradiance is higher. Gaseous MSA does not have a diurnal cycle and measured concentrations in gas and condensed phase are compatible with this species being primarily produced via heterogeneous oxidation of dimethyl sulfide and subsequent partitioning into the gas phase. We also found that NPF in the boundary layer is mainly driven by sulfuric acid but it occurred very rarely over the vast geographical area probed and did not contribute to the CCN budget in a directly observable manner. Despite the near absence of NPF events in the boundary layer, Aitken mode particles were frequently measured, supporting the hypothesis of a free tropospheric source. Iodic acid and MSA were not found to participate in nucleation, however, MSA may contribute to aerosol growth via heterogeneous formation in the aqueous phase.

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Andrea Baccarini^{1,2}, Josef Dommen², Katrianne Lehtipalo^{3,4}, Silvia Henning⁵, 4 Robin L. Modini², Martin Gysel-Beer², Urs Baltensperger² and Julia Schmale¹ 5

¹School of Architecture, Civil and Environmental Engineering, École Polytechnique Fédérale de Lausanne, 6 Lausanne, Switzerland 7

²Laboratory of Atmospheric Chemistry, Paul Scherrer Institute, Villigen PSI, Switzerland ³Institute for Atmospheric and Earth System Research/Physics, University of Helsinki, Helsinki, Finland ⁴Finnish Meteorological Institute, Helsinki, Finland ⁵Institute for Tropospheric Research, Experimental Aerosol and Cloud Microphysics, Leipzig, Germany

Key Points:

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13	•	Production of methanesulfonic acid occurred predominantly in the condensed phase,
14		followed by enhanced volatilization at lower RH
15	•	The observed NPF events were probably driven by sulfuric acid and the environ-
16		mental conditions (mainly temperature and condensation sink)
17	•	Boundary layer new particle formation over the Southern Ocean in summer oc-
18		curred very sporadically and does not contribute to the CCN budget

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 $Corresponding \ author: \ Julia \ Schmale, \ {\tt julia.schmale@epfl.ch}$

19 Abstract

During summer, the Southern Ocean is largely unaffected by anthropogenic emissions, 20 which makes this region an ideal place to investigate marine natural aerosol sources and 21 processes. A better understanding of natural aerosol is key to constrain the preindus-22 trial aerosol state and reduce the aerosol radiative forcing uncertainty in global climate 23 models. We report the concentrations of gaseous sulfuric acid, iodic acid, and methane-24 sulfonic acid (MSA) together with a characterization of new particle formation (NPF) 25 events over a large stretch of the Southern Ocean. Measurements were conducted on board 26 the Russian icebreaker Akademik Tryoshnikov from January to March 2017. Iodic acid 27 is characterized by a particular diurnal cycle with reduced concentration around noon. 28 suggesting a lower formation yield when solar irradiance is higher. Gaseous MSA does 29 not have a diurnal cycle and measured concentrations in gas and condensed phase are 30 compatible with this species being primarily produced via heterogeneous oxidation of 31 dimethyl sulfide and subsequent partitioning into the gas phase. We also found that NPF 32 in the boundary layer is mainly driven by sulfuric acid but it occurred very rarely over 33 the vast geographical area probed and did not contribute to the CCN budget in a di-34 rectly observable manner. Despite the near absence of NPF events in the boundary layer, 35 Aitken mode particles were frequently measured, supporting the hypothesis of a free tro-36 pospheric source. Iodic acid and MSA were not found to participate in nucleation, how-37 ever, MSA may contribute to aerosol growth via heterogeneous formation in the aque-38 ous phase. 30

40 **1** Introduction

Aerosols have a major impact on our climate (Stocker et al., 2014). They scatter 41 and absorb solar radiation and are part of cloud formation processes as cloud conden-42 sation nuclei (CCN) or ice nucleating particles (INP). The combination of aerosol-radiation 43 and aerosol-cloud interactions contributes the largest fraction of uncertainty to the over-44 all radiative forcing budget (Stocker et al., 2014). The present day (PD) aerosol forc-45 ing is calculated against a preindustrial (PI) baseline, which is poorly constrained be-46 cause direct measurements of PI aerosols are impossible. Additionally, the radiative forc-47 ing due to aerosol-cloud interactions (RF_{aci}) is non-linearly dependent on the total aerosol 48 number concentration and is much more sensitive to changes in low concentration regimes, 49 which are more representative of the PI time (Carslaw et al., 2013, 2017). Therefore, 50 the highly uncertain global level and distribution of PI aerosols has a disproportionately 51 large effect on the PD RF_{aci} uncertainty. One way to constrain this uncertainty is to bet-52 ter characterize natural sources of aerosols, which were predominant during the PI time. 53 However, there are very few places on Earth that may still resemble PI-like conditions 54 with minimum anthropogenic influence. Among these locations, the Southern Ocean is 55 probably the region with the highest number of PI-like days during summer (Hamilton 56 et al., 2014). Recently, Regayre et al. (2020) demonstrated that a small set of measure-57 ments over the Southern Ocean can be as effective as a two orders of magnitude larger 58 and more heterogeneous set of data from the Northern Hemisphere in reducing the RF_{aci} 59 in a global climate model. This highlights the value of measurements in pristine and re-60 mote locations. 61

The contribution of anthropogenic activities to the aerosol population over the South-62 ern Ocean is small and generally limited to the more northerly sector (Schmale et al., 63 2019; Uetake et al., 2020). This implies that natural emissions constitute the overwhelm-64 ing share of the aerosol population with sea spray and new particle formation from ma-65 rine emissions presumably being the two main aerosol sources. Other minor sources are 66 volcanic emissions (Schmidt et al., 2012), emissions from sea birds and other animals (Schmale 67 et al., 2013) and blowing snow from ice covered regions (Frey et al., 2020). The concen-68 tration of sea spray aerosol is mainly driven by wind speed and sea state and can vary 69 largely across the Southern Ocean (Quinn et al., 2017; Schmale et al., 2019). Previous 70

measurements reported a contribution between 10% and 100% to the CCN concentra-71 tion, depending also on supersaturation (Quinn et al., 2017; Fossum et al., 2018; Schmale 72 et al., 2019). NPF occurs via the nucleation of low-volatility vapors to form small par-73 ticles, which eventually grow by condensation of the same or other gaseous compounds. 74 Over the Southern Ocean, NPF is thought to happen mainly via sulfuric acid (A. D. Clarke 75 et al., 1998; Yoon & Brimblecombe, 2002; Gordon et al., 2017), which is formed from the 76 oxidation of dimethyl sulfide (DMS), a biogenic compound produced in the water by phy-77 toplankton. During the austral summer the concentration of DMS in the water of the 78 Southern Ocean is the highest of the planet (Lana et al., 2011), with high fluxes into the 79 atmosphere and potentially producing high concentrations of sulfuric acid. However, un-80 der typical boundary layer conditions the concentration of sulfuric acid is too low to form 81 particles alone, and another molecule, such as ammonia, is required to stabilize the nu-82 cleating clusters (Kirkby et al., 2011). Jokinen et al. (2018) reported the first molecu-83 lar characterization of NPF from Aboa station (73.0364°S, 13.4109°W) in Antarctica, 84 showing that new particles are formed via nucleation of sulfuric acid and ammonia. Sources 85 of ammonia over the Southern Ocean are related to animals, mainly bird or seal colonies, 86 which are known to be strong local sources of ammonia (Riddick et al., 2012; Schmale 87 et al., 2013; Riddick et al., 2016). Another potentially important compound for NPF in 88 this region is iodine, which is known to form new particles via iodic acid nucleation (Sipilä 89 et al., 2016; Baccarini et al., 2020; He et al., 2021) and concentrations can be very high 90 in Antarctica. Iodine monoxide concentrations larger than 20 ppt have been reported 91 in coastal Antarctica (Saiz-Lopez et al., 2007; Schönhardt et al., 2008). 92

There are multiple studies investigating new particle formation in Antarctica (Järvinen 93 et al., 2013; Weller et al., 2015; Jokinen et al., 2018; Jang et al., 2019) but observations of NPF over the Southern Ocean have been rare and concentrated around the sea ice re-95 gion (Heintzenberg et al., 2004; Humphries et al., 2015; Dall'Osto et al., 2017). Some 96 field studies have observed a higher concentration of recently formed particles in the free 97 troposphere (A. D. Clarke et al., 1998; Weber et al., 1998; A. D. Clarke & Kapustin, 2002; 98 Sanchez et al., 2021; McCoy et al., 2021) and proposed that NPF may be happening pre-99 dominantly in the free troposphere in the outflow of clouds. Here, formation rates can 100 be higher because of the lower temperatures and smaller condensation sink. Newly formed 101 particles can then be entrained in the boundary layer following for example the passage 102 of cold fronts (Jimi et al., 2008; Gras et al., 2009). These results are also supported by 103 modelling studies showing that typical marine boundary layer conditions are unfavourable 104 for NPF (Katoshevski et al., 1999; Pirjola et al., 2000; Yoon & Brimblecombe, 2002; Ko-105 rhonen et al., 2008; Revell et al., 2019). However, global climate models also tend to un-106 derestimate both the Aitken mode aerosol concentration (Hodshire et al., 2019) and the 107 CCN number (Chambers et al., 2018; Schmale et al., 2019; McCoy et al., 2020) over the 108 Southern Ocean, pointing towards a missing aerosol source or an inaccurate process rep-109 resentation (e.g., too strong deposition velocity) in the models. The underestimation of 110 Aitken mode particles is particularly relevant because NPF is believed to be the largest 111 source of particles in this size range (Seinfeld & Pandis, 2016; MÅrtensson et al., 2010; 112 Spracklen et al., 2007). The reason for this discrepancy is still not known and additional 113 process based measurements over the Southern Ocean are required to better understand 114 the sources and distribution of aerosols. 115

An important process that is often overlooked is DMS oxidation, which is gener-116 ally implemented in models without considering heterogeneous chemistry. This is par-117 ticularly relevant for methanesulfonic acid (MSA), an oxidation product of DMS, which 118 can be more efficiently produced in the aqueous phase than in the gas phase (Hoffmann 119 et al., 2016; Q. Chen et al., 2018) and grow the mass of aerosols activated in cloud droplets 120 (cloud processing). MSA constitutes a large fraction of the secondary aerosol mass over 121 the Southern Ocean, up to 50% compared to the non-sea-salt sulfate aerosol mass (Preunkert 122 et al., 2007; Yan et al., 2019), but its contribution to the CCN budget has not been quan-123 tified so far. 124

In this work, measurements of low-volatility vapors and the observations of NPF 125 events during the Antarctic Circumnavigation Expedition (ACE) will be presented. In 126 particular, we measured the concentration of sulfuric acid, MSA and iodic acid together 127 with naturally charged ions and newly formed aerosol particles. In the Methods (section 128 2) we provide details on the expedition, the instruments used and the methodology adopted 129 to treat the data. The results and discussion (section 3) is divided into three parts. The 130 first part provides a broad overview of the results with a focus on sulfuric acid, iodic acid 131 and MSA distribution over the Southern Ocean. In the second part we provide a detailed 132 analysis on sources and processes controlling the MSA concentration, both in the gas and 133 in the condensed phase. The third part is centered around NPF with a presentation of 134 the events detected during ACE, a characterization of the nucleating vapors and a de-135 scription of the most relevant drivers for the formation of new particles. The conclusions 136 (section 4) summarizes our results and put them into perspective. 137

138 2 Methods

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The Antarctic Circumnavigation Expedition took place between December 2016 and March 2017, sailing around Antarctica across the Southern Ocean on board of the Russian icebreaker *Akademik Tryoshnikov*. The expedition was divided into 3 legs:

- 1. Leg 1: from Cape Town (South Africa) to Hobart (Tasmania) [20 December 2016
 19 January 2017]
 - Leg 2: from Hobart (Tasmania) to Punta Arenas (Chile) [22 January 2017 22 February 2017]
- 1463. Leg 3: from Punta Arenas (Chile) to Cape Town (South Africa) [26 February 2017147- 19 March 2017],

with several stops around islands and other points of interest during the cruise. A detailed description of the voyage is reported in Schmale et al. (2019) and in the cruise report (Walton & Thomas, 2018).

We measured sulfuric acid, MSA and iodic acid using a nitrate chemical ionization 151 Atmospheric Pressure Interface Time-of-Flight Mass Spectrometer (CI-APi-ToF)(Jokinen 152 et al., 2012). The same instrument, an APi-ToF, was also used without the chemical ion-153 ization inlet to characterize the chemical composition of naturally charged ions (Junninen 154 et al., 2010). The concentration and size distribution of newly formed and Aitken mode 155 particles were obtained using a Neutral cluster and Air Ion Spectrometer (NAIS) (Mirme 156 & Mirme, 2013). The particle size distribution (PSD) was measured using a Scanning 157 Mobility Particle Sizer (SMPS) and an Aerodynamic Particle Sizer (APS), together cov-158 ering a size range from 11 nm to 19 μ m. The PSDs obtained from these two different 159 instruments were combined using a mode-fitting technique similar to Modini et al. (2015). 160 The results of the mode-fitting procedure were used to estimate the aerosol number con-161 centration in the different modes and to calculate the condensation sink following Dal Maso 162 et al. (2002). 163

Aerosols and gases were sampled from three different inlets mounted on a container, 164 which was located on the second deck of the ship at a height of about 15 m above the 165 ocean surface. Two of the inlets (the ones used for standard aerosol and trace gas mea-166 surements) consisted of heated 2 m long vertical tubes of 2.54 cm outer diameter and 167 a specifically designed top-cover for isokinetic sampling of particles up to 40 μ m in di-168 ameter, following the Global Atmosphere Watch recommendations for aerosol sampling 169 (Weingartner et al., 1999). A third inlet was specifically designed for short residence time 170 of the sampled air to improve detection of low-volatility vapors and newly formed par-171 ticles. This inlet was a simple 1.5 m long tube of 5 cm inner diameter and a U-shaped 172 bend at the end to prevent rain from entering. It was not heated. The (CI)-APi-ToF and 173 the NAIS were sampling behind this third inlet and were operated only during Leg 2 and 174

Leg 3. A more detailed description of the measurement set-up is provided in the cruise report (Walton & Thomas, 2018).

The CI-APi-ToF was calibrated for sulfuric acid at the end of the campaign with a series of experiments at the Paul Scherrer Institute (PSI) smog chamber (Paulsen et al., 2005), yielding a calibration constant of:

$$C_{ACE} = 6.9 \times 10^9 [-50\% + 100\%]$$
 molecule cm⁻³,

with the calibration uncertainty being indicated in the square brackets. The same calibration constant was used to quantify MSA and iodic acid based on the assumption that
the ionization proceeds at the kinetic limit for species that have a lower proton affinity
than nitric acid as in these cases. A description of the CI-APi-ToF and its calibration
is reported in the Supporting Information (SI) Text S1.

The nitrate CI-APi-ToF is designed to work with a constant addition of nitric acid 182 to the sheath flow in order to produce the reagent ions which are used to ionize the sam-183 ple air. During ACE, the instrument was operated without an active addition of nitric 184 acid due to a technical problem which was identified only at the end of the expedition. 185 Still the background concentration of nitric acid, desorbing from the walls of the inlet 186 lines, was enough to produce a sufficiently high reagent ion concentration like in a reg-187 ularly operated nitrate CI-APi-ToF. This was confirmed by the sulfuric acid calibration, 188 which is comparable with previously reported values (Kürten et al., 2012; Jokinen et al., 189 2012). However, the nitric acid concentration was probably not high enough to take up 190 all the charges produced by the photoionizer. Therefore, reactions with other ions like 191 ${\cal O}_2^-$ and ${\cal CO}_3^-$ also occurred inside the nitrate CI-APi-ToF inlet. These other reactions 192 led to the production of SO_5^- and HSO_4^- from ambient SO_2 , which interfered with the 193 detection of ambient sulfuric acid. A detailed characterization of this issue was performed 194 with experiments at the PSI smog chamber and at the CLOUD chamber at CERN. Un-195 fortunately, because the background production of HSO_4^- was not constant and depended 196 strongly on the instrument settings, such as the inlet flow and voltages, it was not pos-197 sible to correct for it within a reasonable uncertainty. Therefore, all sulfuric acid values 198 reported in this work are uncorrected and should be considered as upper limit estimates. 199

Gases and aerosol particles generated by the ship exhaust and other campaign re-200 lated activities (e.g. helicopter flights) were identified and separated from the background 201 measurement data. As described in Schmale et al. (2019), data were filtered using a method 202 based on particle number, black carbon and CO_2 concentrations leading to a removal of 203 about 50% of the data for the entire expedition. However, there are also species that are 204 not produced by the ship exhaust, like MSA and iodic acid. Figure 1 shows sulfuric acid, 205 MSA and iodic acid data in clean and polluted conditions by means of violin plots, to-206 gether with box and whiskers for a concise statistic summary. A violin plot represents 207 the distribution of the data using a kernel density estimate (Hintze & Nelson, 1998). As 208 expected, sulfuric acid was clearly affected by the ship exhaust with much higher con-209 centrations during polluted conditions, whereas iodic acid was not. 210

A special situation is found for MSA which showed a cluster of high values (larger than 10⁸ molecules cm⁻³) during a polluted period. However, this was a single event where pollution and high MSA occurred coincidentally, but without the pollution causing the high concentration. The event was investigated but it remains unknown why MSA concentrations were so high, because no clear relationship with any external variable was identified. Data from this event were not considered for further analysis because their validity is uncertain.

Even if MSA and iodic acid are not directly emitted by the ship exhaust they could still be affected by the higher aerosol concentration within the exhaust plume which acts as a condensation sink and can reduce the concentration of low-volatility vapors. This effect is not evident from the data distribution shown in Figure 1, however, there are pe-



Figure 1. Violin, and box and whiskers plots of sulfuric acid, MSA and iodic acid divided into clean and polluted conditions. Here, polluted means that the measurements were affected by the exhaust of the research vessel. Polluted periods were identified according to the pollution mask developed by Schmale et al. (2019). The box extends from the first quartile (Q1) to the third quartile (Q3) with a line indicating the median. The whiskers are set to $1.5 \times [Q3-Q1]$.

riods where emissions from the ship reduce the concentration of gaseous MSA and iodic 222 acid. Figure S1 shows an example of this: during pollution (gray shadowing) there are 223 clear spikes in the sulfuric acid and SO_5^- signal (which is produced from SO_2) and in some 224 cases dips in the MSA and iodic acid traces. However, these dips are not always present 225 and generally less pronounced than the pollution spikes, explaining why the overall data 226 distribution seems to be unaffected by pollution. Therefore, given that the effect of pol-227 lution on reducing the concentration of MSA and iodic acid is minor, both polluted and 228 clean data were included in the following analysis (except for the single high-concentration 229 event of MSA mentioned above). 230

We identified NPF events based on the analysis of the particle and ion size distri-231 bution below 10 nm from the NAIS, after excluding the influence from ship exhaust. In 232 particular, only periods with an increase of the sub-10 nm particle concentration larger 233 than a factor of 3 compared to the baseline were considered as NPF events. The sub-234 10 nm particle concentration baseline was calculated using a 2 hour average before and 235 after each potential event. We also excluded cases where the increase in the sub-10 nm 236 concentration could be attributed to a tail of the Aitken mode based on a visual inspec-237 tion of the PSD. 238

²³⁹ **3** Results and discussion

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3.1 Overview of ACE Results

Figure 2 shows an overview map with the expedition track, 6-hour averages of sulfuric acid, MSA and iodic acid concentration measurements and the location of NPF events, which will be described in Section 3.3. The same set of data is also reported in Figures
3.a and 3.b together with the air temperature as hourly averages. Gaps in the data indicate either instrumental problems or periods when the instrument was operated as an
APi-ToF. The time is given in the UTC time-zone here and in the rest of this manuscript,
unless specified differently. Additionally, in Figure 4 the distribution of the data divided
into two latitudinal ranges (above and below 60 °S) is reported. These two latitudinal
bands can be classified as Antarctic and Subantarctic regions (Nowlin & Klinck, 1986).



Figure 2. Map showing the track of the expedition and concentrations of sulfuric acid, MSA and iodic acid. Each marker represents a 6-hour median value with the size being proportional to the concentration on a logarithmic scale. The location of the new particle formation (NPF) events is also reported together with the sea ice concentration (fraction of covered surface) retrieved for January 2017 (Maslanik & Stroeve, 1999). The MSA and iodic acid data were shifted on the map for better visualization. There are no data available for Leg 1 because the CI-APi-ToF and the NAIS were not operated.

Figure 5 illustrates the day and night time data distributions of the trace gases un-250 der consideration by means of violin, and box and whiskers plots. The separation be-251 tween day and night is based on the solar irradiance (SIR) data (night is when SIR is 252 null and day when SIR is larger than 10 Wm^{-2}). Additionally, Figure 6 depicts the di-253 urnal cycles of the data, which were binned according to the local time. We identified 254 the local noon based on the maximum height of the sun above the horizon and the data 255 were shifted accordingly before the diurnal averaging. This procedure was necessary to 256 avoid artefacts due to the eastward movement of the ship, which caused a continuous 257 shift of the local time with respect to UTC. Moreover, the different latitudes at which 258



Figure 3. Time series of sulfuric acid, MSA and iodic acid (left axis). Solid lines represent hourly mean values and the shaded envelopes around these lines represent ± 1 standard deviation. Temperature is shown on the right axis. (a) Leg 2 data and (b) Leg 3 data.

the measurements were performed had an effect on the day duration, which can affect 259 the width of the diurnal profiles. To investigate this effect, we also calculated the diur-260 nal profiles separately for measurements above and below 60 °S as reported in Figure S2. 261 It is evident that the latitudinal variation does not strongly determine the diurnal evo-262 lution of the investigated species but it has an effect on their absolute values (MSA and 263 to a smaller extent also sulfuric acid are higher in more southerly latitudes). The fact 264 that latitude does not have a noticeable effect on the diurnal distribution of the data can 265 probably be explained by solar irradiance being lower at higher latitudes and compen-266 sating for the longer duration of the days (Fig. S2d). 267

The main results, which can be inferred from these overview figures regarding the spatial and temporal distribution of gaseous sulfuric acid, MSA and iodic acid over the Southern Ocean, are:

Sulfuric acid is the only species showing a clear diurnal cycle with higher con-271 centration during midday. This result is consistent with sulfuric acid being predominantly 272 produced via photo-oxidation of SO_2 and is in line with previous measurements in sev-273 eral marine environments (Mauldin et al., 1999; Lucas, 2002; Berresheim et al., 2002) 274 and in Antarctica (Jefferson, Tanner, Eisele, & Berresheim, 1998; Mauldin et al., 2003). 275 On the other hand, its night time values are surprisingly high; this could be an indica-276 tion of a night time production mechanism as previously suggested (Lucas, 2002; Mauldin 277 et al., 2003) or more likely an indication of the instrumental background sulfuric acid 278 production problem described above. Nevertheless, it is safe to assume that the sulfu-279 ric acid increase during day time hours is not driven by the background production be-280 cause SO_2 does not have a diurnal cycle as confirmed by the SO_5^- measurements (Fig. 281 S3). Finally, the sulfuric acid concentration was higher in the region around Antarctica, 282



Figure 4. Violin, and box and whiskers plots of sulfuric acid, MSA and iodic acid separated by latitude. Measurements south and north of 60 °S are representative of Antarctic and Sub-antarctic conditions, respectively. The box extends from the first quartile (Q1) to the third quartile (Q3) with a line indicating the median. The whiskers are set to $1.5 \times [Q3-Q1]$.



Figure 5. Violin, and box and whiskers plots of sulfuric acid, MSA and iodic acid separated by day and night. The separation was done based on the solar irradiance (SIR) value, with night being SIR = 0 Wm⁻² and day SIR > 10 Wm⁻². The box extends from the first quartile (Q1) to the third quartile (Q3) with a line indicating the median. The whiskers are set to $1.5 \times [Q3-Q1]$.

²⁸³ which is a more biologically productive region characterized by higher DMS concentra-



Figure 6. Diurnal profiles of (a) sulfuric acid, (b) MSA and (c) iodic acid. The thick line represents the median and the shaded area the interquartile range. Data were binned using local time determined by the sun height above the horizon. The red line is the solar irradiance median with values on the right axis.

tion in the water (Lana et al., 2011). However, also the SO₅⁻ signal was higher in this region, which may indicate a larger sulfuric acid instrumental background. Therefore, these variations must be interpreted with caution.

MSA does not show any diurnal cycle and the distribution of the data is very sim-287 ilar between day and night, the only difference being the presence of a lower concentra-288 tion mode during night time (Fig. 5). While previous observations already reported that 289 gaseous MSA has a weak to non-existent diurnal cycle (Mauldin et al., 1999; Lucas, 2002), 290 this study is the first to show it on a large regional scale. The absence of a clear MSA 291 diurnal cycle suggests that photochemical production from DMS oxidation is only of mi-292 nor importance, in line with recent modelling work, which proposed that the largest frac-293 tion of MSA is produced in the aqueous phase (Hoffmann et al., 2016; Q. Chen et al., 294 2018). Condensed phase MSA could then be followed by partitioning to the gas phase. 295 The MSA concentration is also higher close to Antarctica, like sulfuric acid, with a dis-296 tribution peaking at around 10^7 molecules cm⁻³ and the median being about 3.7 times 297 higher compared to the Subantarctic region (Figure 4). As described, the region around 298 Antarctica is characterized by higher DMS concentrations which could probably explain 299 the higher MSA concentration. Additionally, higher latitudes correspond to lower tem-300 peratures, which increase the MSA production yield from DMS oxidation compared to 301 SO_2 production (Barnes et al., 2006). We will provide a more detailed analysis of MSA 302 variability and its sources over the Southern Ocean in section 3.2. 303

Iodic acid is characterized by a peculiar diurnal cycle peaking at dawn and dusk with a minimum around noon and very low concentration during night time (the median is below 10^5 molecules cm⁻³). This indicates the presence of a photochemical source

and no production during night. Although the formation mechanism of iodic acid is still 307 not well understood, it is known that iodic acid is formed from the iodine radical, which 308 is photochemically produced from precursor molecules like I₂, HOI or CH₂I₂ (Saiz-Lopez 309 et al., 2012; He et al., 2021; Gómez Martín et al., 2020) and this is consistent with the 310 observations reported here. Figure 7 shows the iodic acid concentration binned by SIR 311 to illustrate the effect of solar radiation. This plot shows that the highest iodic acid con-312 centration is measured when SIR is between 20 and 80 $\mathrm{Wm^{-2}}$ and decreases for higher 313 values up to 1000 Wm^{-2} . The diminished concentration around noon (high SIR) does 314 not have any obvious explanation and it has not been reported before. Two possible hy-315 potheses are (1) that a precursor of iodic acid is reacted away by the OH and/or the HO_2 316 radicals, which have higher concentrations during noon, or (2) that iodic acid or one of 317 its precursors are photolabile and are photolysed during the day. Without a proper un-318 derstanding of iodic acid formation it is not possible to discriminate between the afore-319 mentioned processes. However, Gómez Martín et al. (2020) proposed that iodic acid may 320 be formed from IO or I_2O_3 , where both molecules are photolabile in the near-UV (Saiz-321 Lopez et al., 2012; Lewis et al., 2020) and a reduced concentration of IO during midday 322 has also been predicted (Saiz-Lopez et al., 2014, 2015). Therefore, photolysis is prob-323 ably the reason for the reduced iodic acid concentration at higher SIR values. This phe-324 nomenon may have consequences on the latitudinal and seasonal distribution of iodic acid 325 and its contribution to NPF. He et al. (2021) demonstrated that iodic acid does not re-326 quire the presence of OH to form; ozone and the iodine radical are sufficient. The amount 327 of solar radiation reaching the surface is generally enough to photolyse I_2 even when the 328 atmospheric optical depth is high (e.g. the sun is low over the horizon), meaning that 329 the most favourable conditions for iodic acid formation may be at high latitudes or dur-330 ing early morning/late afternoon. This observation is consistent with recent studies in 331 the Arctic reporting iodic acid NPF in spring and autumn (Baccarini et al., 2020; Beck 332 et al., 2021). Regarding the latitudinal distribution in the Southern Ocean, iodic acid 333 does not show any evident geographical pattern and the data distribution is similar in 334 the Antarctic and Subantarctic regions. It is interesting to note that iodic acid was not 335 enhanced around the coast of Antarctica, despite previous studies showing exception-336 ally high concentrations of other iodine oxides near coastal Antarctica (Saiz-Lopez et al., 337 2007; Schönhardt et al., 2008). This difference is not necessarily a discrepancy consid-338 ering that different iodine oxides, measured in different years and locations, are compared. 339 However, this is a topic that deserves further attention considering the importance of 340 iodic acid for NPF in other locations (Sipilä et al., 2016; Baccarini et al., 2020; Beck et 341 al., 2021). 342

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3.2 Sources and Processes Controlling MSA Concentration

In the previous section we tentatively explained the absence of a diurnal cycle in 344 the concentration of gaseous MSA by the fact that DMS photoxidation is likely not the 345 dominant source of MSA over the Southern Ocean. However, the lifetime of gaseous MSA 346 should also be considered because the atmospheric concentration is controlled by both 347 sources and sinks. MSA is a stable molecule which does not react further under typical 348 tropospheric conditions (Barnes et al., 2006). Therefore, its major sinks are condensa-349 tion to pre-existing aerosol surfaces and dry deposition to the ocean. Previous studies 350 have treated MSA condensation similar to sulfuric acid, assuming kinetic condensation 351 with different accommodation coefficients (De Bruyn et al., 1994; Berresheim et al., 2002; 352 Hanson, 2005; Ammann et al., 2013) varying from about 0.2 to 1 and obtaining a typ-353 ical lifetime of 40 minutes or lower (Berresheim et al., 2014, 2002). If the same approach 354 were used for the ACE data then the median and interquartile (IQR) range of the MSA 355 lifetime for an accommodation coefficient of 0.2 would be 55 (39; 79) minutes, and 23 356 (16; 30) minutes for an accommodation coefficient of 1. In both cases, the lifetime is rel-357 atively short and a decrease in the concentration of MSA during night time would be 358 expected if photooxidation were the dominant source, which was generally not observed. 359



Figure 7. Iodic acid box and whiskers plots as a function of solar irradiance (SIR). Data were binned into different SIR classes as indicated by the axis label. The original data are shown with the small semi-transparent circles. The box extends from the first quartile (Q1) to the third quartile (Q3) with a line indicating the median. The whiskers are set to $1.5 \times [Q3-Q1]$.

We estimated the lifetime of gaseous MSA against dry deposition to the ocean to be around about 23 hours during ACE, which is much longer than the estimated condensation timescales. We calculated dry deposition lifetime by assuming an average boundary layer height of 800 m as reported by Schmale et al. (2019) and a deposition velocity of 1 cm s⁻¹, which is the typical value for nitric acid over the ocean (Seinfeld & Pandis, 2016). Nitric acid and MSA should have a similar deposition velocity as they are equally soluble in water (Seinfeld & Pandis, 2016).

A possible source of gasous MSA which could explain the absence of a diurnal cy-367 cle is evaporation of MSA from the condensed phase. Previous studies already hypoth-368 esized that MSA may evaporate from particles, especially at low relative humidity (RH) 369 (Mauldin et al., 1999; Berresheim et al., 2002). More recently, Hodshire et al. (2019) pro-370 vided a parametrization of MSA equilibrium vapor pressure using the Extended Aerosol 371 Inorganics Model (E-AIM; http://www. aim.env.uea.ac.uk/aim/aim.php, last access: 22 372 April 2021) (S. L. Clegg & Seinfeld, 2006) and showed that MSA could behave both as 373 a non-volatile or semi-volatile species depending on the environmental conditions (tem-374 perature and relative humidity) and aerosol acidity. This is an important result, which 375 can be used to represent more accurately the partitioning of MSA between the gas and 376 the particle phase. However, in the work of Hodshire et al. (2019) only the MSA to am-377 monia ratio was used to evaluate the role of particle acidity without considering the role 378 of other compounds. 379

The gaseous MSA concentration during ACE follows a trend similar to previous studies (Davis et al., 1998; Jefferson, Tanner, Eisele, Davis, et al., 1998; Mauldin et al., 1999; Berresheim et al., 2002) with higher values at lower RH and temperature. Figure 8 shows the gaseous MSA concentration as a function of relative humidity with data separated between day and night for two different parts of the ACE transect. We isolated

these two different periods to reduce confounding factors due to the intrinsic variabil-385 ity of the dataset; they correspond to transects in a defined latitudinal range and with 386 small temperature variations. The first period extends from 4 to 17 February 2017 and 387 includes measurements very close to the Antarctic continent with a temperature median 388 and IQR of -0.8 (-1.3; -0.3) °C. The second period lasts from 4 to 14 March 2017. It 389 is more representative of Subantarctic conditions and was characterized by a tempera-390 ture median and IQR of 1.1 (0.8; 1.4) °C. The number of MSA measurement points con-391 tained in these two periods is similar (80 and 89 hours of measurements, respectively) 392 and corresponds in total to about 2/3 of the entire MSA dataset. The same plot of gaseous 393 MSA as a function of RH for the full ACE dataset is reported in Figure S4. Both fig-394 ure 8 and S4 show a clear increase of gaseous MSA with decreasing RH, most notably 395 during night time. Focusing on Figure 8, in the first period the decrease is evident only 396 for RH greater than 90% and 95% for day and night time, respectively. The second pe-397 riod, instead, is characterized by a more continuous decrease of MSA with increasing RH 398 during night, whereas the trend in the day time data is less clear. 399



Gaseous MSA box and whiskers plot as a function of relative humidity (RH) during Figure 8. two different transects in (a) Leg 2 and (b) Leg 3. Data were separated between day and night and binned into different RH classes as indicated by the axis label. The original data are shown with the small semitransparent circles. The red line in the inset map illustrates the region over which data were collected. The box extends from the first quartile (Q1) to the third quartile (Q3) with a line indicating the median. The whiskers are set to $1.5 \times [Q3-Q1]$. The solid lines in the plots are the predicted MSA gas phase concentrations by partitioning models for different simplified bulk aerosol compositions and as a function of RH (axis on the top). The red line refers to a fully neutralized aerosol system including only sulfate, ammonium and MSA. The purple line relates to a system containing also sea spray aerosol (SSA), in this case the chloride, sodium and sea spray sulfate median concentrations from ACE were used. The green line and shadowed region refer to a system with only 10% of the SSA concentration measured during ACE and varying ammonium concentrations to mimic different degrees of neutralization. The model used for the simulation cannot account for supersaturated solutions when including also sodium and chloride, therefore the two simulations with SSA stop at higher RH because of aerosol efflorescence.

We used E-AIM to investigate if MSA partitioning could explain the increased MSA 400 gaseous concentration at lower RH. The model requires information on the aerosol chem-401 ical composition. For this purpose we used ion chromatography data of daily PM_{10} fil-402 ters (Tatzelt et al., 2020). Non-sea-salt (nss) sulfate and ammonium in the PM_{10} filters 403 were clearly affected by the ship exhaust, and therefore only a subset of the filters (23)404 over a total of 91 filters) with minimum contamination was considered as explained in 405 the SI text S2. Figure S5 reports the concentrations of the major ions in the selected sub-406 set of filters. The mass concentration is dominated by sodium and chloride as expected 407 given the large abundance of sea spray aerosols (SSA) during ACE (Schmale et al., 2019). 408 The nss-sulfate to ammonium ratio points toward a large degree of neutralization (the 409 molar ratio median and IQR are 0.57 and 0.40 - 0.63, respectively). Previous studies 410 in the Southern Ocean and coastal Antarctica have reported generally more acidic aerosols 411 but there is a large range of variability with the nss-sulfate to ammonium ratio varying 412 between 0.5 and 2 and in few cases even larger values (Savoie et al., 1993; Legrand et 413 al., 1998; Quinn et al., 1998; Teinilä et al., 2000; Zorn et al., 2008; Schmale et al., 2013; 414 Xu et al., 2013; Barbaro et al., 2017). It is also important to mention that we do not have 415 any information about the aerosol mixing state but there is probably an external mix-416 ture with SSA being predominantly in the coarse mode and compounds of secondary ori-417 gin (*i.e.* nss-sulfate, ammonium and MSA) in the accumulation mode (Berg et al., 1998; 418 Quinn et al., 1998; Jourdain & Legrand, 2002; Xu et al., 2013). We simulated three dif-419 ferent systems: (I) a system composed only of nss-sulfate, MSA and different concen-420 trations of ammonium, (II) a system dominated by SSA with the sodium and chloride 421 concentration based on the daily PM_{10} filter values, (III) a mixed system composed of 422 sulfate, MSA, ammonium and only 10% of the SSA concentration measured during ACE. 423 Details on the E-AIM simulations are reported in the SI text S2. 424

Based on the E-AIM results we estimated the MSA concentration that would par-425 tition to the gas phase as a function of RH. Figure 8 shows the results for the two tran-426 sects presented before, where the reference value for the condensed phase concentration 427 in each period was taken to be equal to the median concentration from the respective 428 PM_{10} filter data. Additionally, we shifted the simulated gas phase concentration data 429 by an amount equal to the measured gas phase MSA median concentration above 95%430 RH, based on the assumption that at this high RH there would be no repartitioning of 431 MSA from the condensed phase as shown by all simulations. The first system composed 432 of nss-sulfate, ammonium and MSA can reproduce the observed values only for a fully 433 neutralized aerosol; a more acidic aerosol composition would lead to a much higher gas 434 phase MSA concentration (as shown in Figure S6), which is not compatible with our ob-435 servations. On the other hand, for the system with the full SSA aerosol concentration 436 all MSA would stay in the condensed phase with negligible evaporation (in the case of 437 a deliquesced aerosol). The third system produces results that are most in agreement 438 with the observed trend. In this case the nss-sulfate to ammonium ratio has a much smaller 439 influence on MSA partitioning compared to the first system. This result can be explained 440 by the combination of three factors: (i) the overall aerosol acidity is reduced by the SSA 441 components, (ii) SSA is more hygroscopic and takes up more water and (iii) the higher 442 total aerosol mass retains more MSA in the condensed phase. The small effect of the nss-443 sulfate to ammonium ratio on MSA partitioning is consistent with our results consid-444 ering that we observed a comparable increase of MSA at low RH in two very different 445 regions of the Southern Ocean (panel a and b in Fig. 8). The first region being closer 446 to the Antarctic coast and characterized by potentially higher ammonia emission com-447 pared to the second which was characterized by more open ocean conditions where aerosol 448 particles are typically more acidic (Legrand et al., 1998; Zorn et al., 2008). The same 449 450 effect can be observed also in the gas to particle MSA ratio as shown in Figure 9. In fact, the gas to particle MSA ratio during the two transects is essentially equivalent despite 451 the different MSA absolute values. The median and IQR gas to particle ratio in the first 452 period are 0.0047 and (0.0031; 0.0085), while in the second period they are 0.0054 and 453 (0.0028; 0.0084). An aspect that remains unclear is the concentration of gaseous MSA 454

at high RH: for values larger than about 90% the partitioning model would predict a gaseous 455 MSA concentration more than one order of magnitude lower compared to the measure-456 ments. During the day this difference can be explained by gas phase production, which 457 may be the dominant source of gaseous MSA at high RH, but there is no clear expla-458 nation for the night time values. A possible source of error is the choice of MSA ther-459 modynamic properties in E-AIM, which suffers from a large degree of uncertainty, as ex-460 plained in the SI. For example, a reduction in the Henry's law constant would directly 461 affect MSA partitioning, producing higher concentrations in the gas phase. However, this 462 change would affect the gas phase concentration across the entire RH range, leading to 463 unrealistically high values at low RH. The simple approach adopted in this work to de-464 scribe the aerosol chemical composition and mixing state also has an effect on the sim-465 ulation results and probably contributes to this discrepancy. The SSA component, for 466 example, was treated only as a neutral inorganic mixture based on the PM_{10} filter mea-467 surements but it is known that SSA is enriched in organics (Quinn et al., 2014) and is 468 generally characterized by a low pH, even when freshly emitted (Fridlind & Jacobson, 469 2000; Angle et al., 2021). A more acidic aerosol would be characterized by a larger de-470 gree of MSA evaporation from the condensed phase. 471

Our model is clearly a simplification with no pretension to be exhaustive. However, 472 it is based on fundamental thermodynamic calculations and provides support to the hy-473 pothesis of MSA evaporating from the condensed phase at low RH. To our knowledge, 474 the only indications about MSA partitioning from the condensed phase are based on field 475 observations and on thermodynamic modelling similar to those presented in this work, 476 but dedicated experiments are missing. An accurate characterization of MSA equilib-477 rium vapor pressure as a function of aerosol acidity would be highly valuable to improve our understanding of MSA partitioning in a realistic aerosol and its contribution to the 479 total aerosol mass. 480

As described before and shown in Figure 9, we measured a low gas to particle MSA 481 ratio during the entire campaign, around 0.5% on average. These low ratios are in line 482 with previous measurements around coastal Antarctica (Jefferson, Tanner, Eisele, Davis, 483 et al., 1998) and tropical regions (Mauldin et al., 1999; Davis et al., 1999) and seem to 484 support modelling studies predicting that MSA is predominantly formed via aqueous phase 485 oxidation of DMS (Hoffmann et al., 2016; Q. Chen et al., 2018). In order to investigate 486 this hypothesis we calculated the time required to grow the particulate MSA concentra-487 tion from gas phase condensation. This is only a qualitative calculation considering that 488 daily averages were used and that MSA was treated as irreversibly condensing to the par-489 ticles, which is not true as already described. However, our estimate is still valuable because it provides a lower time limit, as condensation cannot be faster than this. Figure 491 S7 shows the result in terms of the number of hours that would be required to grow the 492 observed particulate MSA concentration. Two different accommodation coefficients of 493 0.2 and 1 were used to reproduce the range of values reported in the literature (De Bruyn 494 et al., 1994; Hanson, 2005). Even in the fastest case, when an accommodation coefficient 495 of one is assumed, the typical time required to reproduce the observed particulate MSA 496 is about 3 days, which is equal or even longer than the typical lifetime of an aerosol in 497 the marine BL (e.g. a previous study estimated a lifetime of 2 days for a $0.1 \,\mu\text{m}$ diameter particle over the Indian Ocean (Williams et al., 2002)). 499

In conclusion, measurements of gaseous MSA concentrations conducted during ACE show a lack of a diurnal cycle and an increase at low RH which can be explained by evaporation of MSA from the condensed phase and a low contribution from gas phase oxidation of DMS. Additionally, the low gas to particle MSA ratio consistently supports the hypothesis that MSA is predominantly produced in the aqueous phase as predicted by different modelling studies (Hoffmann et al., 2016; Q. Chen et al., 2018).



Figure 9. MSA concentrations in the gas and particulate phase, gaseous MSA was multiplied by a factor 100 so that the same scale as for particulate MSA could be used. The ratio between gaseous and particulate MSA is reported on the right axis.

3.3 New Particle Formation Over the Southern Ocean

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The frequency of NPF events observed during ACE was low with the events last-507 ing only a few hours and newly formed particles not growing above 10 nm. The only ex-508 ception are two events, which occurred on two consecutive days in the proximity of the 509 Mertz glacier (67.1 °S, 145.0 °E). In this case newly formed particles grew above 10 nm, 510 reaching 20 nm and forming a clear nucleation mode. All the other events were local bursts 511 of newly formed particles, which disappeared shortly after the nucleation onset. Figure 512 10 illustrates the particle and negative ion size distributions corresponding to the 2 in-513 tense NPF events, solar irradiance and the number concentration of particles larger than 514 7 nm. A Roman numeral indicates the event number, in this and all the other figures. 515 Unfortunately, for these events no information concerning the chemical composition of 516 the nucleating vapor is available due to a malfunctioning of the mass spectrometer. Both 517 events have a clear diurnal pattern, with particles being produced during the day and 518 suggesting the involvement of sulfuric acid. However, the first and most intense NPF event 519 starts very early in the morning which could also be compatible with the iodic acid di-520 urnal profiles measured during the campaign (Fig. 6). Hence it is not possible to uniquely 521 determine the NPF mechanism. The ion size distribution shows some peculiar bands be-522 tween 2 and 4 nm, these are probably wind generated ions as similar features have been 523 observed also at other snow-covered sites at high wind speeds (Manninen et al., 2010; 524 X. Chen et al., 2017), but it is not clear if they were involved in the NPF process. The 525 effect of wind is shown in Figure S8, which reports both the negative and positive ion 526 size distribution together with wind speed, relative wind direction and distance to land. 527 It is evident that these ion bands are present only for wind speeds larger than about 10 528 ms^{-1} in close proximity to land, suggesting that blowing snow may be involved as re-529 ported by X. Chen et al. (2017). 530

These two NPF events were interrpted by several short pollution periods. However, the natural origin of nucleation is ensured by the continuous growth of new particles under persistently strong wind conditions with a prevalent wind direction from the clean sector (*i.e.* the bow of the ship as shown in Fig. S8).

Figure 11 shows two other NPF events, which are representative of the local NPF 535 type detected during the expedition. Here, mass spectrometric measurements are avail-536 able. These two events have again a diurnal evolution and the measurement of the neu-537 tral molecules and charged clusters suggest an involvement of sulfuric acid. It is known 538 that, in this temperature range, sulfuric acid alone cannot lead to NPF at these low con-539 centrations and a stabilizing compound is needed (e.g. ammonia or amines) (Kirkby et 540 al., 2011; Almeida et al., 2013). However, the largest cluster that was detected during 541 all NPF events was the sulfuric acid trimer only, without any additional molecule. The 542 trimer alone is not indicative of the full nucleation mechanism and the stabilizing com-543 pound was not identified. Larger clusters were probably not measured because of the low 544 concentrations of the nucleating vapors, which did not produce enough clusters (the sul-545 furic acid trimer was already close to the detection limit of the mass spectrometer). 546

Figures S9 and S10 show the remaining 3 NPF events, which are similar to those described above. The event in Figure S10 is slightly different because it occurred during sunset. However, the real onset of nucleation was not detected in this case (particles were already larger than 4 nm), indicating that the event started during day time and the newly formed particles were then advected to the ship location (or alternatively, the ship transited through the NPF location).

Figure 12 reports the locations of all the detected NPF events and the correspond-553 ing boundary layer 5-day airmass back trajectories calculated with the Lagrangian anal-554 ysis tool LAGRANTO (Sprenger & Wernli, 2015), for additional details the reader is re-555 ferred to Thurnherr et al. (2020). Events are numbered according to Figures 10, 11, S9 556 and S10. All events are characterized by a marine influence with air masses usually com-557 ing from the more productive sea ice region around Antarctica. The only exception is 558 event VI, which happened closer to South America and was not influenced by any sea 559 ice region. 560

The two regional NPF events I and II were exceptional because of the environmen-561 tal conditions encountered. In particular, the temperature and the condensation sink were 562 low during these events with the median temperature being within the first 5 percentiles 563 and the median condensation sink within the first 20 percentiles for both events. At the 564 same time, solar irradiance was above the 75th percentile. Importantly, the combination 565 of these 3 parameters was unique during the entire ACE expedition, which means that 566 there were no other occurrences with similarly low temperature, condensation sink and 567 high solar radiation at the same time. These three parameters are particularly impor-568 tant for NPF because they control the sulfuric acid concentration: more intense solar radiation enhances the OH production increasing the sulfuric acid concentration, while the 570 condensation sink is the main sulfuric acid loss term. They also control the nucleation 571 rates: temperature has a direct effect on the nucleating cluster stability (Kirkby et al., 572 2011). The exceptional combination of these three parameters probably explains why 573 these two events were different from the rest of the campaign and also helps understand-574 ing the difference between ACE and the NPF results reported from Aboa, an Antarc-575 tic research station located about 130 km inland from the Southern Ocean coast (Jokinen 576 et al., 2018). There, Jokinen et al. (2018) reported the frequent occurrence of NPF when 577 the air mass was coming from the surrounding oceanic or sea ice region. This region should 578 be similar in terms of emissions to the area sampled during the most southerly part of 579 the ACE track. However, the frequency and intensity of NPF events recorded in Aboa 580 was much higher. NPF in Aboa is driven by sulfuric acid, which was frequently higher 581 than 10^7 molecules cm⁻³, different from ACE where this threshold was almost never ex-582 ceeded (Figure S11). However, this difference is unlikely driven by DMS emissions only, 583 which are equal or higher along the ACE track than in the region of air mass origin for 584 the Aboa NPF events (Lana et al., 2011; Mahajan et al., 2015). Rather, the higher sul-585



Figure 10. New particle formation events I and II, (a) total particle size distribution (2.5 - 42 nm) and number concentration of particles larger than 7 nm (right axis). (b) negatively charged ion size distribution (0.8 - 42 nm) and solar irradiance time series (right axis). Pollution spikes are highlighted with a different colour map (magma), in this case a less stringent pollution mask was used instead of the default from Schmale et al. (2019) in order to clearly show the evolution of the NPF event.

furic acid reported at Aboa can probably be explained by the lower condensation sink, 586 a factor two lower on average than during ACE (Figure S12), and the higher SIR. Ad-587 ditionally, the temperature measured in Aboa was 2 to 5 degrees lower than the min-588 imum temperature recorded during ACE (with the exception of a single day), and this 589 also enhances NPF. The different temperature and SIR values in Aboa are simply due 590 to the meteorological conditions (the Antarctic continent is colder and less cloudy than 591 the surrounding ocean (King & Turner, 1997)), whereas the lower condensation sink can 592 be explained by the short lifetime of the coarse mode aerosol, which is responsible for 593 a large fraction of the condensation sink over the ocean and would be removed by the 594 time they have reached Aboa. Another important difference is the detection of ammo-595 nia, which was frequently measured by Jokinen et al. (2018) in negative clusters with 596 acids but never observed during ACE. A quantitative comparison of the ammonia con-597 centration is not possible because this molecule was not measured directly during either 598 campaign (ammonia was only detected as a cluster with sulfuric acid). It is possible that 599 during the study of Jokinen et al. (2018) the ammonia concentration was on average higher 600 compared to ACE. However, during ACE a large variety of different locations were ex-601 plored, including places in close proximity to penguin colonies which are known to be 602 strong sources of ammonia (Schmale et al., 2013; Croft et al., 2016). Therefore, it is un-603 likely that ammonia was the only limiting factor for NPF during ACE but it may have 604 contributed together with the other factors described above (temperature, SIR and con-605 densation sink). 606

⁶⁰⁷ Despite the rare occurrence of boundary layer NPF, an Aitken mode was frequently ⁶⁰⁸ detected during ACE contributing to a large fraction of the total particle number con-⁶⁰⁹ centration as reported in Figure S13. The origin of these Aitken mode particles remains



Figure 11. New particle formation events III and IV, (a) total particle size distribution (2.5 - 42 nm) and, on the right axis, number concentration of particles larger than 7 nm and solar irradiance time series. (b) negatively charged ion size distribution (0.8 - 42 nm), on the right axis the concentration of neutral molecules measured with the CI-APi-ToF (solid line) and the negative ions measured with the APi-ToF (round markers) are reported. Only the 4 ions with the largest signal are reported here, the sulfuric acid and MSA monomers are not present because of the instrument mass transmission, which was set to higher masses. Pollution spikes are high-lighted with a different color map (magma), in this case a less stringent pollution mask was used instead of the default from Schmale et al. (2019).

unknown but it is compatible with the hypothesis from previous studies suggesting that
NPF may be prevalently occurring in the free troposphere (Weber et al., 1998; Korhonen et al., 2008; Sanchez et al., 2021; McCoy et al., 2021). However, it is difficult to explain the growth of the freshly formed particles to the typical 30 to 50 nm Aitken mode
diameter (Schmale et al., 2019) considering the low concentration of condensable vapors.
Investigating this topic in detail is beyond the scope of this work, but it clearly deserves
more attention.

617 4 Conclusions

The Southern Ocean is one of the most pristine locations on Earth (Hamilton et 618 al., 2014) and measurements in this region can be valuable to better understand the state 619 of the atmosphere in preindustrial times and constrain the radiative forcing uncertainty 620 in global climate models (Regayre et al., 2020). This work presents an overview of the 621 spatial distribution of sulfuric acid, MSA and iodic acid across the Southern Ocean to-622 gether with ultrafine particle and ion concentration as well as size distribution. These 623 are all quantities that are relevant for new particle formation and growth. Obtaining a 624 better understanding of the processes and the environmental conditions regulating their 625 distribution can, therefore, be valuable to properly represent aerosol sources and prop-626 erties in global climate models. There are studies which previously investigated trace gases 627



Figure 12. Map showing the ACE track, the location of NPF events and the 5-day boundary layer air mass back trajectories for each of the events. The back trajectories are shown using semi-transparent green dots, the density of dots in a specific region is proportional to the amount of trajectories passing over that region. The figure also shows the sea ice concentration (fraction of covered surface) retrieved for January 2017 (Maslanik & Stroeve, 1999).

(sulfuric acid and MSA)(Jefferson, Tanner, Eisele, & Berresheim, 1998; Jefferson, Tanner, Eisele, Davis, et al., 1998), new particle formation (Weber et al., 1998; Weller et al., 2015; Dall'Osto et al., 2017) or both (Jokinen et al., 2018) over the Southern Ocean and coastal Antarctica. However they were focused on single locations. The work presented here is the first comprehensive investigation of trace gases and new particle formation across the Southern Ocean providing a wide geographical coverage and a broader understanding of the processes involved.

Sulfuric acid vapor was characterized by a clear diurnal cycle with maxima at daytime consistent with photochemical production from SO₂. The concentration was lower compared to recent measurements from coastal Antartica (Jokinen et al., 2018), especially considering that only an upper limit was reported here. This had a direct effect on the occurrence of NPF events which were weak in terms of particle production and very sporadic. The lower sulfuric acid vapor concentration was attributed mainly to environmental reasons, such as the high condensation sink and rather low solar irradiance.

Iodic acid also exhibited a diurnal cycle with very low concentrations during night time, as expected from a molecule that is formed from the photochemically produced iodine radical. However, the iodine concentration peaked at dawn and dusk with consistently lower concentration during the central part of the day when solar radiation was stronger. This observation, which has not been reported before, could be related with the photolysis of an iodic acid precursor molecule $(e.g. \text{ IO or } I_2O_3)$ (Saiz-Lopez et al., 2012; Lewis et al., 2020). This result is important because it indicates that iodic acid could eventually reach higher concentrations when solar radiation is lower, like in spring or in autumn, if the iodine flux were comparable. As a consequence, there may be periods of the year when iodic acid may be relevant for NPF also over the Southern Ocean. Additional measurements to investigate this possibility are needed.

Finally, the gaseous MSA concentration is too low to explain the required partic-653 ulate MSA values via a condensation mechanism. This suggests that MSA may be pre-654 dominantly produced in the aqueous phase, as indicated already by previous modelling 655 studies (Q. Chen et al., 2018; Hoffmann et al., 2016). Additionally, gaseous MSA does 656 not show any diurnal cycle and tends to increase under dryer conditions, indicating that 657 the gas phase MSA may be driven by evaporation from the particle phase. This is con-658 sistent with predictions from a thermodynamic model considering a mixture of sea spray, 659 MSA ammonium and sulfate. Both the increase of MSA at lower RH and the absence 660 of a diurnal cycle are in line with previous measurements of MSA in marine environments 661 (Mauldin et al., 1999; Berresheim et al., 2002; Yan et al., 2019). This suggests that a more 662 accurate treatment of MSA production and partitioning in atmospheric chemistry mod-663 els is needed to improve the representation of marine sulfur compounds in the atmosphere. 664

Schmale et al. (2019) reported a large discrepancy in the CCN number concentra-665 tion around the coast of Antarctica when comparing measurements with values modelled 666 using the Global Model of Aerosol Processes (GLOMAP) (Mann et al., 2010). This area 667 corresponds also to the strongest MSA signal detected during the entire expedition (both 668 in the gas and in the particle phase as shown in Figure 9). The area is also known to ex-669 hibit one of the largest DMS concentrations (both in the water and in the atmosphere) 670 in the world during summer (Lana et al., 2011; Mahajan et al., 2015). GLOMAP (as many 671 other global climate models) only includes homogeneous production of MSA in the gas 672 phase, whereas it does not consider condensation of this MSA nor heterogeneous pro-673 duction which could contribute to the underestimation of the CCN concentration around 674 the coast of Antarctica. Future studies should focus on the MSA partitioning and aque-675 ous phase production to understand its contribution to the concentration of CCN and 676 their properties. 677

The results obtained during ACE clearly show that discernible NPF in the bound-678 ary layer is rare across the Southern Ocean in summer and only in exceptional cases it 679 contributes to the aerosol Aitken mode population. Sulfuric acid was the main nucle-680 ating compound for the observed NPF events. A base, such as ammonia or amines, would 681 also be required to stabilize the nucleating clusters given the low sulfuric acid concen-682 tration (Kirkby et al., 2011; Almeida et al., 2013) but no stabilizing compound was iden-683 tified. We also found that environmental conditions, mainly temperature and the con-684 densation sink, are critical in determining the occurrence of NPF and are likely respon-685 sible for the different observations compared to previous studies between the open ocean 686 and coastal Antarctica in terms of boundary layer NPF (A. D. Clarke et al., 1998; Heintzen-687 berg et al., 2004; Jimi et al., 2008; Gras et al., 2009; Weller et al., 2015; Jokinen et al., 688 2018). The low relevance of boundary layer NPF together with the frequent detection 689 of Aitken mode aerosols is compatible with new particles being formed in the free tro-690 posphere and then transported downward as shown also by other studies (A. D. Clarke 691 et al., 1998; Weber et al., 1998; A. D. Clarke & Kapustin, 2002; Sanchez et al., 2021). 692 This hypothesis cannot be confirmed with our dataset because we lack information con-693 cerning the vertical distribution of aerosol particles. Future expeditions in the region should specifically address this topic, investigating aerosol sources both in the boundary layer 695 and in the free troposphere while trying to understand their exchange processes. 696

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Tota availability The gaseous sulfuric acid and methanesulfonic acid data are available on the Zenodo database (https://zenodo.org/) (Baccarini et al., 2019b, 2019a) as well as the PM_{10} filter data (Tatzelt et al., 2020). The remaining dataset will be made available upon acceptance of the publication.

Author contribution A.B., J.D., M.GB. and J.S. conceived and designed the experimental set-up. A.B., J.D, K.L., S.H., R.L.M. and J.S. prepared the instruments and operated them during the campaign. A.B., J.D., R.L.M. and J.S. analysed the data. A.B. prepared the plots and wrote the manuscript with major contributions from J.D., U.B. and J.S. All authors interpreted the results and commented on the manuscript.

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Supporting Information for "Low-volatility vapors and new particle formation over the Southern Ocean during the Antarctic Circumnavigation Expedition"

Andrea Baccarini^{1,2}, Josef Dommen², Katrianne Lehtipalo^{3,4}, Silvia

Henning⁵, Robin L. Modini², Martin Gysel-Beer², Urs Baltensperger²and

Julia Schmale¹

¹School of Architecture, Civil and Environmental Engineering, École Polytechnique Fédérale de Lausanne, Lausanne, Switzerland

 $^2 {\rm Laboratory}$ of Atmospheric Chemistry, Paul Scherrer Institute, Villigen PSI, Switzerland

³Institute for Atmospheric and Earth System Research/Physics, University of Helsinki, Helsinki, Finland

 $^4{\rm Finnish}$ Meteorological Institute, Helsinki, Finland

 $^5 \mathrm{Institute}$ for Tropospheric Research, Experimental Aerosol and Cloud Microphysics, Leipzig, Germany

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- 1. Text S1 to S2 $\,$
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Introduction

This supporting information contains first a text section describing the operation and calibration of a nitrate chemical ionization Atmospheric Pressure Interface Timeof-Flight Mass Spectrometer (CI-APi-ToF)(Jokinen et al., 2012). A second text

section describes how the extended Aerosol Inorganic Model (E-AIM; http://www. aim.env.uea.ac.uk/aim/aim.php, last access: 22 April 2021) (S. L. Clegg & Seinfeld, 2006) was used to model the gas-particle partitioning of methanesulfonic acid (MSA) using an aerosol mixture representative of the Southern Ocean aerosol.

Figures S1 to S13 provide additional information to the results shown in the main text and are referenced therein, whereas Figures S14 and S15 are related to Text S1 and S2. Tables S1 and S2 are related to Text S2 and provide details regarding the E-AIM simulation.

Text S1. Nitrate CIMS operation and calibration

Sulfuric acid, MSA and iodic acid were measured with a nitrate CIMS (Jokinen et al., 2012). These molecules are detected either as clustered with the nitrate monomer or as deprotonated species. The concentration of a species X is calculated as:

$$[X] = C \frac{X^{-} + XNO_{3}^{-}}{\sum_{n=0}^{2} (HNO_{3})_{n}NO_{3}^{-}},$$
(1)

with X- being the deprotonated species and C the calibration factor which can be experimentally determined. The instrument is typically calibrated by injecting a known amount of sulfuric acid which can be produced in different ways, for this work we used a series of experiments at the PSI smog chamber (SC). The sensitivity of the instrument can be assumed to be constant (*i.e.* the same calibration factor C can be applied), under the assumption that each collision leads to a stable cluster. This is the case for sulfuric acid, MSA and iodic acid because they have a lower proton affinity than nitric acid (Jokinen et al., 2012; Eisele & Tanner, 1993; Sipilä et al., 2016).

The CI inlet is generally operated with an electric field driving the nitrate ions inside the sample flow. However, for a part of ACE no voltage was applied to the inlet due to a

technical issue. In this case, diffusion and turbulence in the inlet drive the mixing of the nitrate with the sample flow. A different sensitivity could be expected compared to a CI inlet operated in the standard way. However, as shown below, the difference was found to be smaller than the calibration uncertainty. Therefore, the same calibration constant was used for the entire ACE campaign.

The calibration of the nitrate CIMS was performed at the PSI SC, a $27m^3$ flexible Teflon bag inside a wooden enclosure where temperature can be controlled. Four xenon arc lamps and a set of 80 UV-A light tubes are used to initiate photochemistry inside the chamber, further details on the chamber can be found in Paulsen et al. (2005). Experiments were carried out injecting SO₂ at a mixing ratio between 0 and 15 ppb, ozone between 0 and 250 ppb and trimethylbenzene (TMB) between 0 and 10 ppb. The temperature inside the chamber was kept fixed at 20 ± 2 °C and the RH was varied between 20% and 40%. The reaction of SO₂ with OH was used to produce sulfuric acid, whose concentration inside the chamber can be described by a simple kinetic model:

$$\frac{d[\mathrm{H}_2\mathrm{SO}_4]}{dt} = k_{\mathrm{OH} + \mathrm{SO}_2}[\mathrm{OH}][\mathrm{SO}_2] - (W + CS)[\mathrm{H}_2\mathrm{SO}_4], \tag{2}$$

with $k_{\text{OH} + \text{SO}_2}$ being the reaction rate constant (Wine et al., 1984), W the sulfuric acid wall loss rate inside the chamber and CS the condensation sink. The wall loss rate was estimated to be $(3.5 \pm 0.9) \times 10^{-3} \text{ s}^{-1}$. The CS represents the sulfuric acid loss term to the particles and can be calculated from the particle number size distribution following Dal Maso et al. (2002).

Equation 2 can be used to calculate the concentration of sulfuric acid inside the chamber at the steady state as:

$$[H_2SO_4] = \frac{k_{OH + SO_2}[OH][SO_2]}{W + CS}.$$
(3)

The OH concentration is estimated from the decay of TMB which reacts with it at a known reaction rate constant (Kramp & Paulson, 1998).

The calibration result is reported in Figure S14, where the ordinate shows the sulfuric acid concentration calculated from Equation 3 and the abscissa the average of the measured sulfuric acid signal normalized by the reagent ion concentration. The uncertainty of the calculated sulfuric acid concentration was obtained by standard error propagation from Equation 3, whereas the uncertainty on the CI-APi-ToF measurements was calculated as the standard error of the mean. A weighted least squares regression was applied to estimate the calibration constant C as described in Equation 1. The CI inlet is usually operated with an electric field to force the reagent ions into the sample flow. During the calibration the instrument was run also without the electric field in order to simulate the CI-APi-ToF operating conditions during ACE. The results presented in Figure S14 clearly show that there is no systematic difference between the two operating modes of the inlet. Therefore a single calibration value was used:

$$C_{ACE} = 6.9 \times 10^9 [-50\% + 100\%] \text{ molecule cm}^{-3}, \tag{4}$$

where the calibration coefficient was already corrected for diffusional losses of sulfuric acid in the inlet line. An uncertainty of [-50% + 100%] was associated with the calibration factor, which is a standard estimate for sulfuric acid measurements in the field. This estimate is larger than the linear regression confidence interval and accounts for the intrinsic variability of field conditions, which are difficult to quantify.

Text S2. E-AIM calculation

E-AIM was used to estimate MSA partitioning over the Southern Ocean aerosol. MSA is not included among the default species available in the E-AIM library and needs to

be created by the user defining some of its fundamental thermodynamic properties. A part from the most basic properties, such as the molar mass and the molar volume, MSA thermodynamic properties are not readily available and model studies often use very different values. Table S1 reports all the values that were used in this study and the corresponding references, as a general criterion we decided to use values based on experimental results rather than modelling or ab initio calculation. By reporting all the information required to include MSA in E-AIM we hope to provide a useful reference for the community and to foster discussion, which thermodynamic properties are the most appropriate values for MSA.. All the values in Table S1 were taken directly from the cited reference with minimal adaption (*e.g.* unit of measure), with the only exception of the surface tension parameters. In this case we had to recreate the surface tension data from Myhre, D'Anna, Nicolaisen, and Nielsen (2004) and fit them with the function defined by Dutcher, Wexler, and Clegg (2010) to obtain the parameters in the form required by E-AIM.

Concerning the specific details of the E-AIM simulations, we fixed the temperature to 273.15 K and varied relative humidity (RH) between 60% and 100%. These are representative values for the environmental conditions encountered during the two transects described in the main text. E-AIM model II was used for the simulations containing only sulfate, ammonium and MSA whereas E-AIM model IV was used when also sodium and chloride were included. Aerosol components were forced to stay in the liquid form as the efflorescence RH is below typical values encountered in the marine boundary layer. However, E-AIM model IV is unable to account for supersaturated solutions, for this reason only RH values above $\sim 75\%$ could be considered when using this model.

The aerosol composition used for the E-AIM simulation was based on ion chromatography (IC) analysis of daily PM_{10} filters. These filters were collected using a high-volume sampler on the upper deck of the ship. The sampler had an automatic system to stop the sample flow when the wind was coming from the direction of the ship chimney. However, this system proved to be not sufficient to prevent sampling of the exhaust plume as evident from the high elemental carbon (EC) concentrations measured on some of the filters (larger than $1 \,\mu \text{g m}^{-3}$). We used EC as a proxy for contamination from the ship exhaust on the PM_{10} filters because ambient concentration of black carbon in the Southern Ocean is generally very low (less than 40 ng m⁻³)(Schmale et al., 2019). Figure S15 shows the ammonium and the non-sea-salt (nss) sulfate concentrations measured on the PM_{10} filters as a function of EC, there is an evident positive correlation indicating that both ammonium and nss-sulfate are probably affected by the ship exhaust. However, nss-sulfate and ammonium seem to become independent from the ship exhaust for EC values below about $0.6 - 0.7 \,\mu \text{g m}^{-3}$. Under this condition, the concentration of these two compounds is dominated by natural sources and not by the ship exhaust. Hence, we decided to consider only the filters characterized by an EC concentration below $0.65 \,\mu \text{g m}^{-3}$ to minimize the influence of the ship exhaust. Additionally, we also excluded the filters with a total sampled volume below 240 m³, which is one third of the maximum possible volume. Such a low sampling volume indicates that the samplers were often turned off because of the wind coming from the direction of the ship chimney. Figure S5 shows the concentration of the ions relevant for the E-AIM calculation from this filter selection.

In order to understand the effect of aerosol composition on MSA partitioning we run three different sets of simulations using E-AIM. For the first set of simulations we decided

to fix the nss-sulfate concentration based on the median concentration measured during ACE and varied the ammonium concentration to obtain different ratios (2:1, 1:1) and 1:2). The MSA concentration was fixed based on the median MSA to nss-sulfate ratio (Fig.S5). Figure S6 shows the model predictions in terms of the MSA vapour pressure and the gas fraction due to partitioning from the condensed phase. The second set of simulations includes also sea spray, in the form of sodium, chloride and sea-salt (ss) sulfate. For the concentrations of ss-sulfate and chloride we used the median values from ACE, whereas the sodium concentration was increased to achieve neutrality of the SSA and account for the presence of other cations which cannot be included in E-AIM (*i.e.* magnesium and calcium). Finally, for the third set we considered only 10% of the sea spray concentration, together with nss-sulfate ratio from 0.25 to 2. Results are shown in the main text. Table S2 shows the concentrations of aerosol constituents used for each model run.

The predicted MSA gas phase concentrations shown in the main text (Fig. 8) was calculated considering the estimated gas phase fraction from each E-AIM simulation multiplied by the average particulate MSA concentration measured during the two periods under analysis (0.21 μ g m⁻³ and 0.09 μ g m⁻³ for the first and the second period, respectively).

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Figures



Figure S1. Time series showing the effect of pollution on the major compounds measured with the nitrate CI-APi-ToF, the polluted periods are highlighted with a gray shadow according to the pollution mask described in Schmale et al. (2019). SO_5^- is generated inside the inlet of the CI-APi-ToF and is a proxy for ambient SO_2 .



Figure S2. Diurnal profiles of (a) sulfuric acid, (b) MSA, (c) iodic acid and (d) solar irradiance for the entire campaign and separated by latitude. Here, high and low latitude indicates measurements above and below 60 °S, respectively.



Figure S3. Diurnal cycle of SO_5^- . The thick line represents the median and the shaded area the interquartile range. The red line shows the solar irradiance median with values on the right axis.

April 22, 2021, 11:45pm

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Figure S4. Gaseous MSA box and whiskers plot as a function of relative humidity (RH). Data were separated between day and night and binned into different RH classes as indicated by the axis label. The original data are shown with the small semi-transparent circles. The box extends from the first quartile (Q1) to the third quartile (Q3) with a line indicating the median. The whiskers are set to $1.5 \times [Q3-Q1]$.



Figure S5. Box and whiskers plot of the ion chromatography data from PM_{10} daily aerosol filter. Only a subset of filters with minimum contamination from the ship exhaust was selected. (a) Concentration of the ions used for the thermodynamic modelling, (b) molar ratio of non-seasalt (nss) sulfate to ammonium and nss-sulfate to MSA.

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Figure S6. E-AIM results for MSA partitioning over an aerosol mixture composed of sulfate, MSA and ammonium as a function of relative humidity. (a) MSA equilibrium vapour pressure (C*), (b) fraction of MSA in the gas phase due to partitioning from the condensed phase. The sulfate, MSA and ammonium concentrations used for this simulation are reported in table S2 (run 1.1, 1.2 and 1.3).



Figure S7. Number of hours required to reproduce the observed particulate MSA concentration assuming kinetic condensation of gaseous MSA with two different accommodation coefficients. This condensation time was calculated based on the daily average values and the figure shows the number of occurrences as a histogram and a box and whiskers plot on top. The box extends from the first quartile (Q1) to the third quartile (Q3) with a line indicating the median. The whiskers are set to $1.5 \times [Q3-Q1]$.

Hours



Figure S8. Wind induced ions, (a) negative ion size distribution and, on the right axis, wind speed and relative wind direction (the bow of the ship corresponds equivalently to 0° or 360°).
(b) positive ion size distribution, on the right axis the distance to land is shown. Pollution periods are highlighted with a different color map (magma).



Figure S9. New particle formation event, (a) total particle size distribution and, on the right axis, number concentration of particles larger than 7 nm and solar irradiance time series. (b) negatively charged ion size distribution, on the right axis the concentration of negative ions measured with the APi-ToF (round markers) is reported. Only the 4 ions with the largest signal are reported here, the sulfuric acid and MSA monomers are not presented because of the instrument mass transmission, which was set to higher masses. Pollution spikes are highlighted with a different color map (magma).



Figure S10. New particle formation event, (a) total particle size distribution and, on the right axis, number concentration of particles larger than 7 nm and solar irradiance time series. (b) negatively charged ion size distribution, on the right axis the concentration of neutral molecules measured with the CIMS (solid line) and the negative ions measured with the APi-ToF (round markers) are reported. Only the 4 ions with the largest signal are reported here, the sulfuric acid and MSA monomers are not presented because of the instrument mass transmission, which was set to higher masses.



Figure S11. Comparison of the sulfuric acid measured during ACE and at Aboa. The figure shows a histogram reporting the frequency of the observation with a kernel density estimate (thick line) and box and whiskers plot on top. The box extends from the first quartile (Q1) to the third quartile (Q3) with a line indicating the median. The whiskers are set to $1.5 \times [Q3-Q1]$.

6.0

log(Concentration [molecules cm⁻³])

6.5

7.0

8.0

7.5

4.5

5.0

5.5



0.0 -4.5 -4.0 -3.5 -3.0 -2.5 -2.0log(Condensation Sink [s⁻¹])

Figure S12. Comparison of the condensation sink measured during ACE and at Aboa. The figure shows a histogram reporting the frequency of the observation with a kernel density estimate (thick line) and box and whiskers plot on top. In this case the condensation sink from ACE was calculated using the same size range as at Aboa to improve the accuracy of the comparison (from 10 nm to 900 nm). The box extends from the first quartile (Q1) to the third quartile (Q3) with a line indicating the median. The whiskers are set to $1.5 \times [Q3-Q1]$.





Figure S13. Aitken mode particles during ACE: (a) total Aitken mode particle number concentration from the fit of the SMPS size distribution and (b) ratio of the Aitken mode particle number to the total number particle concentration. The thick line is a 3 hours mean and the shadowed region indicates a 1 standard deviation interval.



Figure S14. Sulfuric acid calibration with smog chamber experiments. Each data point represents a different experiment with the CI-APi-ToF normalized sulfuric acid signal on the x-axis and the corresponding concentration of sulfuric acid inside the chamber on the y-axis. Error bars on the x-axis are equal to the standard error of the mean whereas errors on the y-axis are equal to the propagated uncertainty via Equation 3. Orange and blue markers indicate measurements where the CI inlet was operated with and without voltage, respectively.



Figure S15. Concentration of nss-sulfate and ammonium as a function of elemental carbon. The vertical dashed line was drawn at $0.65 \,\mu \text{g} \text{ m}^{-3}$ of carbon and indicates the concentration below which the effect of ship pollution is not evident anymore on nss-sulfate and ammonium. Elemental carbon measurements below detection limit were fixed to a value equal to $0.01 \,\mu \text{g} \text{ m}^{-3}$.

Tables

Property	Unit of measure	Value	Reference
Molar mass	$g \text{ mol}^{-1}$	96.1	Barnes et al. (2006)
Molar Volume	$\mathrm{cm}^3 \mathrm{\ mol}^{-1}$	64.93	Barnes et al. (2006)
First dissociation constant	$mol kg^{-1}$	73	Clarke and Woodward (1966)
Enthalpy of dissociation	kJ mol $^{-1}$	0	Not determined
Henry's law constant	$mol kg^{-1} atm^{-1}$	$8.9 imes 10^{11}$	S. Clegg and Brimblecombe (1985)
Henry's law enthalpy change	$kJ mol^{-1}$	14.644	De Bruyn et al. (1994)
Surface tension: c_1	${ m mN}~{ m m}^{-1}$	138.23	Myhre et al. (2004)
Surface tension: c_2	${ m mN}~{ m m}^{-1}~{ m K}^{-1}$	-0.284	Myhre et al. (2004)
Surface tension: a_{ws}	${ m mN}~{ m m}^{-1}$	147.86	Myhre et al. (2004)
Surface tension: b_{ws}	$mN m^{-1} K^{-1}$	-0.275	Myhre et al. (2004)
Surface tension: a_{sw}	${ m mN}~{ m m}^{-1}$	-167.117	Myhre et al. (2004)
Surface tension: a_{sw}	${ m mN}~{ m m}^{-1}~{ m K}^{-1}$	0.400	Myhre et al. (2004)

 ${\bf Table \ S1.} \quad {\rm MSA \ thermodynamic \ properties}$

 Table S2.
 Concentration of aerosol constituents used for the E-AIM simulations

Run number	nss-sulfate	Ammonium	MSA	ss sulfate	Chloride	Sodium
	$\rm nmol~m^{-3}$	$\mathrm{nmol}\ \mathrm{m}^{-3}$	$\mathrm{nmol}\ \mathrm{m}^{-3}$	$\rm nmol~m^{-3}$	$\mathrm{nmol}~\mathrm{m}^{-3}$	$nmol m^{-3}$
1.1	2.3	4.6	1	0	0	0
1.2	2.3	2.3	1	0	0	0
1.3	2.3	1.15	1	0	0	0
2.1	2.3	3.9	1	7.2	134	148
3.1	2.3	4.6	1	0.7	13.4	14.8
3.2	2.3	2.3	1	0.7	13.4	14.8
3.3	2.3	1.15	1	0.7	13.4	14.8
3.4	2.3	0.58	1	0.7	13.4	14.8