A seasonal analysis of aerosol NO₃⁻ sources and NO_x oxidation pathways in the Southern Ocean marine boundary layer

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9 Abstract. Nitrogen oxides, collectively referred to as NOx (NO + NO2), are an important component of 10 atmospheric chemistry involved in the production and destruction of various oxidants that contribute to the 11 oxidative capacity of the troposphere. The primary sink for NOx is atmospheric nitrate, which has an influence on 12 climate and the biogeochemical cycling of reactive nitrogen. NOx sources and NOx to NO3⁻ formation pathways 13 remain poorly constrained in the remote marine boundary layer of the Southern Ocean (SO), particularly outside 14 of the more frequently sampled summer months. This study presents seasonally resolved measurements of the 15 isotopic composition (δ^{15} N, δ^{18} O and Δ^{17} O) of atmospheric nitrate in coarse mode (> 1µm) aerosols, collected 16 between South Africa and the sea ice edge in summer, winter and spring. Similar latitudinal trends in δ^{15} N-NO₃ were observed in summer and spring, suggesting similar NOx sources. Based on $\delta^{15}N\text{-}NO_3$, the dominating 17 18 primarymain NOx sources were likely a combination of lightning, biomass burning and/or soil emissions at the 19 low latitudes, as well as oceanic alkyl nitrates and snowpack emissions from continental Antarctica or the sea ice 20 at the low, mid and high latitudes, respectively. Snowpack emissions associated with photolysis were derived 21 from both the Antarctic snowpack as well as from snow on sea ice. A combination of natural NOx sources, likely 22 transported from the lower latitude Atlantic contribute to the background level NO3⁻ observed in winter, with the 23 potential for a stratospheric $NO_{\underline{a}_{+}}$ source evidenced by one sample of Antarctic origin. <u>Greater values of $\delta^{18}O_{-}$ </u> 24 NO3- in spring and winter compared to summer, suggest an increased influence of oxidation pathways that 25 incorporate oxygen atoms from O3 into the end product NO3- (i.e., N2O5, DMS and XOhalogen oxides (XO)) Low 26 summertime δ^{48} O-NO₂⁻ (< ~70‰) are consistent with daytime processes involving oxidation by OH dominating 27 nitrate formation, while higher winter and springtime δ^{18} O-NO₂⁻ (>~60‰) indicate an increased influence of O₃ 28 $\frac{1}{2}$ oxidation (i.e., N₂O₅, DMS, BrO). Significant linear relationships between δ^{18} O and Δ^{17} O suggest isotopic mixing 29 between H2O(v) and O3 in winter, and isotopic mixing between H2O(v) and O3/XO in spring with the addition of a 30 third endmember (atmospheric O2) becoming relevant in spring. The onset of sunlight in spring, coupled with 31 large sea ice extent, can activate chlorine chemistry with the potential to increase peroxy radical concentrations, 32 contributing to oxidant chemistry in the marine boundary layer.

33 1 Introduction

The atmosphere of the Southern Ocean is geographically remote from major anthropogenic influences. Although
there is evidence of microplastics and at times long-range transport of anthropogenic pollution (Jacobi et al., 2000;
Obbard 2018;), the Southern Ocean marine boundary layer (MBL) is one of the few regions dominated by natural
sources, and as such at times it can serve as a proxy for the pre-industrial atmosphere. The pre-industrial

atmosphere is used as a baseline for comparing the magnitude of anthropogenic impacts on climate (e.g., Haywood
 and Boucher, 2000; Hamilton et al., 2014; Schmale et al., 2019).

40 Nitrogen oxides ($NO_x = NO + NO_2$) are an important part of biogeochemical cycling and influence the oxidative

41 capacity of the troposphere as they are involved in the production and destruction of ozone and hydroxyl radicals

42 (Lawrence and Crutzen, 1999; Finlayson-Pitts and Pitts, 2000). The primary sink for NO_x is atmospheric nitrate

 $\label{eq:solution} 43 \qquad (NO_3`), which impacts both air quality and climate by influencing particulate matter load and Earth's radiative$

44 heat budget (IPCC, 2013; (Park and Kim, 2005)).

45 The logistical difficulties of measurement campaigns to the remote Southern Ocean, particularly in winter, have 46 resulted in a lack of observational data from this region including that of NOx sources and sinks (Paton-Walsh et 47 al., 2022). Consequently, the seasonality of NO_x cycling remains poorly constrained in the Southern Ocean MBL. Globally, fossil fuel combustion is the primary NO_x source (van der A et al., 2008), far exceeding natural 48 49 emissions like biomass burning (Finlayson-Pitts and Pitts, 2000), soil processes (Davidson and Kingerlee, 1997) 50 and lightning (Schumann and Huntrieser, 2007). However, regional budgets of NOx sources can have a variety of 51 anthropogenic and natural contributors. In the summertime Southern Ocean MBL, natural NOx sources are the 52 mainprimary contributors to atmospheric NO₃⁻ formation (Morin et al., 2009; Burger et al., 2022). Along the 53 South African coastline, these natural NOx sources include a combination of lightning, biomass burning and soil 54 emissions (Morin et al., 2009). In coastal Antarctica, or near to the marginal ice zone, NOx emitted from snow 55 cover serves as the mainprimary precursor to atmospheric NO3⁻ (Savarino et al., 2007; Morin et al., 2009; Shi et 56 al., 2021; Burger et al., 2022). Over the mid-latitude region of the Southern Ocean, sea surface emissions of a 57 group of nitrogen gases referred to as alkyl nitrates (RONO2), have recently been proposed as a NOx source leading 58 to NO3⁻ formation in the MBL (Fisher et al., 2018; Burger et al., 2022). During winter, NO_x sources to the Antarctic 59 troposphere primarily include long-range transported peroxyacetyl nitrates (PAN) and stratospheric inputs (Savarino et al., 2007; Lee et al., 2014; Walters et al., 2019). To our knowledge, however, there are no 60 61 observational data regarding NOx sources from the Southern Ocean MBL during winter, and few observations in 62 spring.

In addition to there being multiple NO_x sources across the Southern Ocean MBL, several different oxidation
 pathways can be responsible for NO_x to NO₃⁻ conversion, varying with chemistry and time of day (Savarino et al.,
 2007). Once emitted, NO is rapidly oxidised by ozone (O₃) (R1), peroxy radicals (RO₂ or HO₂) (R2), and/or

- 66 halogen oxides (XO; where X = Br, Cl, or I) (R3), to NO₂.
- $67 \qquad \underline{\text{NO}} + \underline{\text{O}}_3 \rightarrow \underline{\text{NO}}_2 + \underline{\text{O}}_2 (\underline{\text{R1}})$
- 68 $\underline{\text{NO} + \text{RO}_2 \text{ (or HO}_2)} \rightarrow \underline{\text{NO}_2 + \text{RO (or OH) (R2)}}$
- $69 \qquad \underline{\text{NO} + \text{XO}} \rightarrow \text{NO}_2 + X \text{ (R3)}$
- 70 $\underline{NO_2 + O_2 + hv} \rightarrow NO + O_3 (R4)$
- 71 <u>Under sunlit conditions, NO₂ is readily photolyzed to regenerate NO and O₃ (R4). The recycling of NO_x between</u>
- **NO** and NO₂ happens much faster than NO_3 oxidation to NO_3 during the day (Michalski et al., 2003). On a
- 73 global scale, NO is primarily oxidised to NO₂ by ozone (O₃), followed by although over the open ocean oxidation

74	by peroxy radicals (HO2 and RO2, while NO to NO2 oxidation via XO is relatively minor) can also occur		
75	(Alexander et al., 2020)		
76	During summer in the Southern Ocean MBL, NO ₂ is subsequently oxidised primarily by hydroxyl radicals (OH)		
77	to form HNO_3 (R5)		
<i>``</i>			
78	$\underline{\mathrm{NO}}_{2} + \underline{\mathrm{OH}} + \underline{\mathrm{M}} \rightarrow \underline{\mathrm{HNO}}_{3} + \underline{\mathrm{M}} (\underline{\mathrm{RS}})$		
79	In winter, under dark conditions, when the photolytic production of OH stops, NO ₂ is oxidised primarily by O ₃ to		
80	form nitrate radicals (NO ₃) (R6). NO ₃ can then react with NO ₂ to form dinitrogen pentoxide (N ₂ O ₅) followed by		
81	hydrolysis on a wet particle surface to form $HNO_3 \cdot (R7-R8)$.		
01	injurity sits on a wet particle surface to form $11003 - 300 - 300$		
82	$\underline{\mathrm{NO}}_{2} + \underline{\mathrm{O}}_{3} \underline{\mathrm{NO}}_{3} + \underline{\mathrm{O}}_{2} \ (\underline{\mathrm{R6}})$		
83	$NO_3 + NO_2 + M \rightleftharpoons N_2O_{S(\ell)} + M (R7)$	_	Formattad Subscript
00			Formatted: Subscript
84	$\underline{N_2O_{S(e)} + H_2O_{(l)} + surface \rightarrow 2HNO_{S(aq)}(R8)}$		Formatted: Subscript
85	Alternatively, HNO ₃ can be formed by the reaction of NO ₃ with hydrocarbons (HC) (e.g., dimethylsulphide	\searrow	Formatted: Subscript
86	(DMS)) (R9).		Formatted: Subscript
00	$(DNG)/(\overline{NZ})$.		
87	<u>NO₃ + HC or DMS \rightarrow HNO₃ + products (R9)</u>		
88	Lastly, halogen in places with elevated halogen concentrations, NO2 can be oxidised by reactive halogens (e.g.,		
89	bromine oxide (BrO)), to from HNO ₃ -chemistry may result in NO_3^- formation via the production and subsequent		
90	hydrolysis of halogen nitrates (R10-R11), as has been suggested for coastal Antarctica in summer (Baugitte et al.,		
91	2012).		
51	2012).		
92	$\underline{\mathrm{XO}} + \mathrm{NO}_2 \rightarrow \underline{\mathrm{XNO}}_3 (\text{R10})$		
93	$XNO_3 + H_2O_{(l)} + surface \rightarrow HNO_{3(ac)} + HOX (R11)$		Formattad Subscript
55		\leq	Formatted: Subscript
94	The nitrogen (N) and oxygen (O) isotopic composition of atmospheric NO_3^- provides information regarding NO_x		Formatted: Subscript
95	sources and NO $_3$ ⁻ formation pathways (i.e., NO oxidation to NO $_2$ and NO $_2$ oxidation to NO $_3$ ⁻). This technique has		
96	been applied in polluted (Elliot et al., 2007; Zong et al., 2017), open ocean (Hastings et al., 2003; Altieri et al.,		
97	2013; Kamezaki et al., 2019; Burger et al., 2022) and polar environments (Walters et al., 2019). Stable isotope		
98	ratios are reported as the ratio of the heavy to light isotopologues of a sample relative to the constant isotopic ratio		
99	of a reference standard, using delta (δ) notation in units of "per mil" (‰) following Eq. (1):		
100	$\delta = \left(\left(R_{sample} / R_{standard} \right) - 1 \right) \times 1000 \tag{1}$		
101	where R represents the ratio of ¹⁵ N/ ¹⁴ N, ¹⁸ O/ ¹⁶ O or ¹⁷ O/ ¹⁶ O in the sample and in the reference standard,		
102			

- 103 (Bolhke et al., 2003).
- 104 The N isotopic composition of atmospheric NO₃⁻ (δ^{15} N-NO₃⁻) largely reflects the δ^{15} N of different precursor NO_x 105 emissions (e.g., Elliott et al., 2019 and references therein), but can be influenced by isotopic fractionation during

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106 NO_x cycling and NO_x to NO_3^- conversion (Walters and Michalski 2015; Walters et al., 2016; Li et al., 2021). $\delta^{15}N^{-1}$ 107 NO_3 is therefore useful for constraining NO_x sources. For example, the $\delta^{15}N$ of lightning NO_x is approximately 108 0% (Hoering, 1957), biomass burning may produce NOx with a 815N range of -7 to 12% (Fibiger and Hastings, 109 2016), while soils lead to NO_x emissions with relatively low δ^{15} N signatures (-44.2 to -14.0%; Miller et al., 2018). 110 More relevant to the remote Southern Ocean is lightning NO_x which has a δ^{15} N signature of approximately 0‰ 111 (Hoering, 1957). Thisand is distinct from the snowpack NO_x source, which typically has a very low δ^{15} N signature 112 of ~ - 48‰ (Berhanu et al., 2014; Berhanu et al., 2015) on the order of -50 to -20‰ (Wagenbach et al., 1998; 113 Winton et al., 2020), depending on the degree of snowpack NO3^{- 15}N enrichment (Shi et al., 2018). Savarino et 114 al., (2007) derived an Antarctic stratospheric NO_x source signature of $19 \pm 3\%$. Additionally, Burger et al. (2022) 115 estimated the δ^{15} N signature of NO_x produced by surface ocean RONO₂ emissions over the mid-latitude Southern 116 Ocean to be $-21.8 \pm 27.6\%$.

117 The O isotopic composition of atmospheric NO₃⁻ (δ^{18} O- and Δ^{17} O-NO₃⁻) reflects the oxidants responsible for NO₃⁻ 118 formation, as atmospheric oxidants have distinct O isotope signatures (Michalski et al., 2011). \delta18O-NO3- and 119 Δ^{17} O-NO₃⁻ are thus useful for identifying pathways of NO₃⁻ production (Michalski et al., 2003; Hastings et al., 120 2003; Alexander et al., 2020). O3 possesses a distinctively large 17O excess as a result of non-mass-dependent 121 isotope fractionation. This ¹⁷O excess is expressed as $\Delta^{17}O = \delta^{17}O - 0.52 \text{ x } \delta^{18}O$ (Berhanu et al., 2012). Non-mass 122 dependent fractionation occurs in the troposphere and is thought to originate from related to asymmetric molecules 123 of excited ozone (O_3^*) that lose excess energy via stabilisation to product O_3UV photolysis breaking oxygen 124 molecules to form O₃ (Heidenreich & Thiemens, 1986; Ireland et al., 2020). As a result, O₃ possesses a uniquely 125 high terminal $\Delta^{17}O = 39.2 \pm 2\%$ (Vicars and Savarino, 2014) that can be transferred to NO₃ during oxidation 126 reactions between NOx and O3 (Thiemens 2006; Savarino et al., 2008; Michalski and Bhattacharya, 2009), or NOx 127 and other oxidants like XO where the oxygen atom originated from O₃ (Savarino et al., 2016). Δ^{17} O-NO₃⁻ therefore 128 can serves as a proxy for the influence of O3 and/or XO during NO3⁻ formation (Berhanu et al., 2012).

129 O_3 also has a uniquely high terminal $\delta^{18}O = 126.3 \pm 11.9\%$ (Vicars and Savarino, 2014) compared to other 130 oxidants that have a Δ^{17} O of 0‰ and much lower δ^{18} O signatures (Michalski et al., 2003; Michalski et al., 2011). 131 For example, atmospheric O_2 has a $\delta^{18}O$ signature of 23.9‰ and the $\delta^{18}O$ of OH and H₂O are negative (Michalski 132 et al., 2011). As such, a higher δ^{18} O or Δ^{17} O for atmospheric NO₃⁻ reflects the increased influence of O₃ and/or 133 <u>XO</u> on NO₃⁻ formation, while a lower δ^{18} O or Δ^{17} O occurs when there is an increased contribution from other 134 oxidants lack of exchange of O atoms with O3 (Hastings et al., 2003; Fang et al., 2011; Altieri et al., 2013). 135 Oxidation by peroxy radicals would also result in a lower $\delta^{18}O$ and $\Delta^{17}O$ signature for atmospheric nitrate because 136 the O atom in peroxy radicals derives from atmospherice O2.

137 <u>Antarctic t</u>-Fropospheric oxidation chemistry has been well characterised using Δ^{17} O at coastal (Savarino et al., 138 2007; Ishino et al., 2017) and interior Antarctic sites (Frey et al., 2009; Savarino et al., 2016; Walters et al., 2019). 139 A distinct seasonal cycle in Δ^{17} O-NO₃⁻ is generally observed whereby a higher relative contribution from O₃ 140 oxidation and/or stratospheric input occurs during winter, and more HO_x + RO_x oxidation occurs during summer. 141 The Atlantic Southern Ocean is less constrained in terms of oxidation chemistry, with growing evidence that the 142 atmospheric oxidant budget is poorly understood in unpolluted low-NO_x environments (Beygi et al., 2011). This study presents the first seasonally resolved data set of coarse mode (> 1 μ m) atmospheric NO₃⁻ concentration and isotopic composition from the Atlantic Southern Ocean MBL. Using air mass back trajectories and observed aerosol δ^{15} N-NO₃⁻, δ^{18} O-NO₃⁻ and Δ^{17} O-NO₃⁻ between Cape Town, South Africa and the marginal ice zone this work aims to identify how the main sources and formation pathways of NO₃⁻ vary over the remote Southern Ocean from winter through spring and summer.

148 2 Methods

149 2.1 Sample collection

150 Samples were collected on board the Research Vessel (R/V) SA Agulhas II during three voyages to and from the 151 marginal ice zone in summer (7th to 21st December 2018 & 27th February to 15th March 2019), winter (19th July to 152 12th August 2019) and spring (13th October to 19th November 2019) (Fig. 1). The summer samples presented here 153 are the same as those in Burger et al. (2022). The winter and spring samples were collected and analysed as in 154 Burger et al. (2022), with any methodological modifications noted below. Briefly, all voyages departed from Cape 155 Town (33.9° S, 18.4° E) and sailed southward along the Good Hope transect (0°E), until reaching Penguin Bukta 156 (71.4° S, 2.5° W) in summer and the northern extent of the sea ice in winter (approximately 58.1° S) and spring 157 (approximately 59.3° S). The ship then returned to Cape Town, sailing north via the Good Hope transect, with a 158 deviation to South Georgia in the summer. In spring an additional ice edge transect was conducted during which 159 the ship sailed from 0 to approximately 22° E and back, before returning to Cape Town.

Size-segregated aerosols were collected on the ninth floor above the bridge (approximately 20 m above sea level), using a high-volume air sampler (HV-AS; Tisch Environmental). Air was pumped at an average flow rate of 1.3 m³ min⁻¹ through a five-stage cascade impactor (TE-235; Tisch Environmental), loaded with pre-combusted (400°C for 4 hours) glass fibre filters. Given that aerosol nitrate in the MBL is predominantly present in the coarse mode (> 1 μ m), only filter stages 1 through 4 were analysed. The aerodynamical diameter of particles captured by filter stages 1, 2, 3 and 4 are > 7 μ m, 3 to 7 μ m, 1.5 to 3 μ m and 1 to 1.5 μ m, respectively.

166 A sector collector was used to restrict HV-AS activity to avoid contamination of the filters with ship stack 167 emissions (Campbell Scientific Africa). The HV-AS only operated if the winds were blowing at an angle less than 168 120° or greater than 240° from the bow of the ship during winter, and less than 75° or greater than 190° from the 169 bow of the ship during spring. These criteria were altered based on the dominant wind direction during each 170 voyage to ensure sufficient sample collection while avoiding contamination. In addition to wind direction, the 171 wind speed had to exceed 0 m s⁻¹ for ten minutes for the HV-AS to begin sampling. Filters were removed from 172 the cascade impactor inside a laminar flow cabinet (Air Science), placed in individual zip-sealed bags and stored 173 at -20°C until analysis.

An attempt was made to ensure that there were at least 24 hours of in-sector sampling before removing filters from the cascade impactor, to ensure atmospheric NO_3^- concentrations were sufficient for isotope analysis (Sect.

176 2.3). However, this was not always possible as on occasion filters had to be removed early due to unusual ship

177 manoeuvres that could have resulted in sample contamination by ship stack emissions if left unattended. Sampling

duration ranged from 11 to 36 hours in winter and 7 to 41 hours in spring (Table S1).

179 During each voyage, a field blank was collected by fitting the cascade impactor with a set of filters and loading

180 the HV-AS in the same manner that atmospheric samples were deployed. The cascade impactor was then

181 immediately removed without turning on the HV-AS pump. The field blanks were removed from the cascade

182 impactor and stored in the same manner as the atmospheric samples. All chemical analysis performed on samples

183 was performed on the field blanks to assess any possible contamination during filter deployment or laboratory

184 procedures.

185 2.2 Sample analysis

186 <u>Once back on land, f</u>Filters were extracted using ultra-clean deionised water (DI; 18 M Ω) under a laminar flow 187 cabinet (Air Science). The extraction ratio was approximately 30 cm² to 100 cm² of filter in 30 mL of DI. Extracts

cabinet (Air Science). The extraction ratio was approximately 30 cm² to 100 cm² of filter in 30 mL of DI. Extracts
were immediately sonicated for one hour and then stored at 4°C for at least 12 hours. Thereafter, extracts were
filtered (0.2 µm) using an acid washed syringe into clean 30 mL HDPE bottles and stored at -20°C until analysis

190 (Burger et al., 2022).

191 2.2.1 NO₃⁻ concentration analysis

192 [NO3-] was determined using a Thermo Scientific Dionex Aquion Ion Chromatography (IC) system equipped with 193 an autosampler. The anion IC contained an AG22 RFIC 4 x 50 mm guard column and AG22 RFIC 4 x 250 mm 194 analytical column. A six-point standard curve was run on each day of analysis (Dionex Seven Anion-II Standard) 195 and an R² value > 0.999 was required for sample analysis to proceed. Final [NO₃⁻] were corrected by subtracting 196 the field blanks, which represented on average_32% and 59% of the [NO₃] in winter and spring, respectively. 197 Where the field blank had a [NO3-] greater than that of the sample, the sample [NO3-] was assumed to be zero. 198 Samples were measured for $[NO_3]$ only once to preserve sample volume for isotopic analysis (Sect. 2.2.23), 199 motivated by the small difference between repeated sample measurements from the summertime dataset $(SD_p =$ 200 0.3 µmol L⁻¹). It is important to note that given the low [NO₂] of the field blanks (<1.5 µM), no isotopic analysis 201 could be performed using blank filters.

202 2.2.24 Isotopic analysis

203 The isotopic composition of atmospheric NO₃⁻ (δ^{15} N, δ^{18} O, and Δ^{17} O-NO₃⁻) was measured using the denitrifier 204 method (Sigman et al., 2001; Casciotti et al., 2002; Kaiser et al., 2007). In brief, a natural strain of denitrifying 205 bacteria, Pseudomonas aureofaciens, that lack the terminal nitrous oxide (N2O) reductase enzyme were used to 206 convert aqueous NO3⁻ quantitatively to N2O gas. The product N2O was analyzed by Gas-Chromatograph IRMS 207 (Thermo- Scientific Delta V Plus) for simultaneous isotopic determination of ¹⁵N/¹⁴N and ¹⁸O/¹⁶O (Sigman et al., 208 2001; Casciotti et al., 2002). The ¹⁵N/¹⁴N of samples was corrected for the contribution of ¹⁷O to the peak at mass 209 45 using Δ^{17} O determined for each sample, with values ranging from 21.7% to 44.4%. International reference 210 materials (Table S2) IAEA-N3 and USGS34 were used to normalize isotopic values to air (δ^{15} N), and IAEA-N3, 211 USGS34 and USGS-35 were used to normalize to VSMOW (δ^{18} O) scales. The pooled standard deviation of 212 sample replicates and duplicates for $\delta^{15}N$ was 0.19% (n = 16) and for $\delta^{18}O$ was 0.27% (n = 16). The pooled 213 standard deviations of sample references IAEA-N3, USGS-34, USGS-35 for δ^{15} N and for δ^{18} O are reported for 214 each season in Table S3.

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For winter and spring samples, Δ^{17} O-NO₃⁻ was characterized by using a separate 50 nmol aliquot to convert NO₃⁻ to N₂O, thermally decomposing the N₂O to N₂ and O₂ in a gold furnace at 770°C and analyzing the isotopic composition of O₂ for determination of ¹⁸O/¹⁶O and ¹⁷O/¹⁶O (Kaiser et al., 2007; Fibiger et al., 2013). The product O₂ was referenced to USGS34 and USGS35, and a 50/50 mix of USGS34 and USGS35 was also quantified within runs serving as a quality control check. The pooled standard deviations for Δ^{17} O were 0.84‰ (n=21), 0.90‰ (n=21), and 0.61‰ (n=18) for USGS34, USGS35, and the 50/50 mix, respectively. The pooled standard deviation of sample replicates and duplicates was 0.63‰ in winter and 0.31‰ in spring.

223 It is important to note that given the low $[NO_3^-]$ of the field blanks (< 1.5 μ M), no isotopic analysis could be

performed on the blank filters and therefore the blank was not subtracted from the isotope results. However, we

note that there was no relationship found between the blank percent contribution and δ^{15} N- or δ^{18} O-NO₃⁻ for

226 <u>spring and winter. This indicates that the measured signal is not driven by the blank contribution.</u>

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228 2.2.3 Sea water sampling and NO2⁻ concentration analysis,

Seawater samples were collected in triplicate every two hours from the ships underway system (position at depth approximately 5 m) for the analysis of surface ocean nitrite concentrations ([NO₂⁻]). Seawater samples for NO₂⁻
determination where immediately frozen at -20°C and stored in dark conditions until analysis. [NO₂⁻] was analysed using the colorimetric method of Grasshof et al. (1983) using a Thermo Scientific Genesys 30 visible spectrophotometer (detection limit of 0.05 µmol L⁻¹). The majority of seawater [NO₂⁻] analysis was conducted while at sea.

235 2.3 Air mass back trajectory analysis

236 Airmass back trajectories (AMBTs) were computed for each hour in which the HV-AS was operational for at least 237 45 minutes of that hour. Given that the ship was moving, a different date, time and starting location was used to 238 compute each AMBT. An altitude of 20 m was chosen to match the height of the HV-AS above sea level and 72-239 hour AMBTs were computed to account for the lifetime of NO3- in the atmosphere. Model estimates of the 240 atmospheric lifetime of NO3⁻ range from approximately three to five days (Lu et al., 2021). AMBTs become 241 increasingly uncertain the further back in time they are used (Sinclair et al., 2013), particularly in the remote 242 Southern Hemisphere. To minimize this uncertainty, the shortest possible AMBTs are generated while still 243 accounting for the lifetime of NO3⁻ (i.e., 72-hours). Daily 120-hour AMBTs computed for the duration of each 244 voyage were additionally computed (See supplemental), to confirm that even when utilising the maximum 245 estimate for NO3 atmospheric lifetime, no continental influence from southern Africa is expected. All AMBTs 246 were computed with NOAA's Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Stein 247 et al., 2015; Rolph et al., 2017), using NCEP Global Data Assimilation System (GDAS) output, which can be 248 accessed at http://www.arl.noaa.gov/ready/hysplit4.html (NOAA Air Resources Laboratory, Silver spring, 249 Maryland).

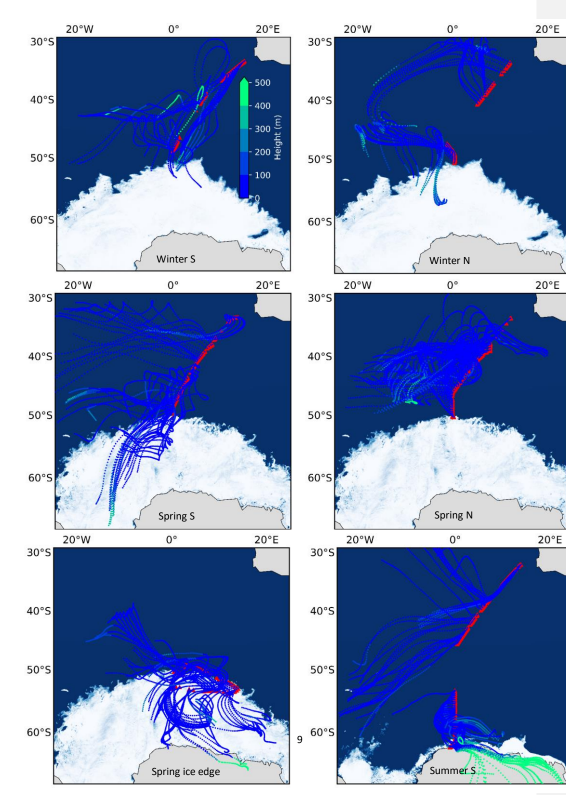
250 3 Results and Discussion

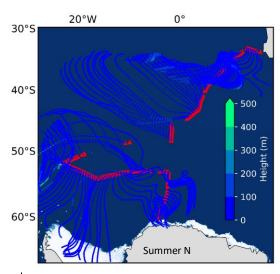
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Formatted: Font: (Default) Times New Roman, Bold Formatted: Font: Bold 251 During all three seasons sampling was conducted across the Atlantic Southern Ocean. AMBTs indicate that no 252 samples experienced any continental influence from South Africa (Fig 1), such that no direct anthropogenic 253 emission sources are considered. 72-hour AMBTs confirm that the Atlantic sector of the Southern Ocean was a 254 dominant source region for most samples collected throughout all seasons. Airmasses experienced very little 255 interaction with sea ice in winter (Fig 1a &b), while extensive interaction with sea ice was experienced by 256 airmasses sampled in spring, particularly at the high latitudes during the south bound leg (Fig 1c) and ice edge 257 transect (Fig 1e). In summer, some high latitudes air masses traversed coastal Antarctica before being sampled, 258 particularly on the south bound leg (Fig 1f), while some interaction with sea ice was also experienced by high 259 latitude air masses on both legs (Fig 1f & g). The potential for sea ice influence is supported by the relatively low 260 height (< 100 m) of AMBTs (Fig. 1). As a result, air masses originated from a mixture of source regions ranging 261 from the open ocean to sea ice to Antarctic continental ice. The remoteness of all the locations at which air masses 262 originated from motivates the investigation of natural NOx sources below.





264	Figure 1. 72-hour AMBTs computed for each hour of every filter deployment made in winter (19th July to 12th	
265	August 2019) on both the southbound (Winter S) and northbound (Winter N) voyages, in spring (13th October to	
266	19th November 2019) on the southbound voyage (Spring S), northbound voyage (Spring N) and ice edge	
267	transect (Spring ice edge) and in summer (7th to 21st December 2018 & 27th February to 15th March 2019) on the	
268	southbound (Summer S) and northbound (Summer N) voyages. Red triangles indicate the ships cruise track	
269	during each filter deployment. The AMBTs are coloured by height (m) (blue to green colour bar).	
270	A	
271	Figure 1.72-hour AMBTs computed for each hour of every filter deployment (grey lines) made in winter on both	
272	the southbound (a) and northbound (b) voyages, in spring on the southbound voyage (c), ice edge transect (d) and	
273	northbound voyage (e) and in summer on the southbound (f) and northbound (g) voyages. Red circles indicate the	
274	ships cruise track during each filter deployment.	
275	3.1 Seasonal variation in atmospheric NO ₃ ⁻ concentrations	
276	In winter, atmospheric $[NO_3]$ were very low across the Atlantic Southern Ocean ranging from below detection to	
277	22.3 ng m ⁻³ (Fig. $2a^2$; blue diamonds). A single outlier exists with a relatively high [NO ₃ ⁻] equivalent to 222.9 ng	

22.3 ng m⁻³ (Fig. 2a2; blue diamonds). A single outlier exists with a relatively high $[NO_3^-]$ equivalent to 222.9 ng m⁻³ in winter, although it is comparable to summertime $[NO_3^-]$ (Fig. 2a2; orange circles). In spring, atmospheric $[NO_3^-]$ ranged from 3.3 ng m⁻³ to 74.4 ng m⁻³, with one sample below detection limit. Higher $[NO_3^-]$ were observed at the lower latitudes and at the higher latitudes, while lower $[NO_3^-]$ were observed in the mid-latitude Atlantic Southern Ocean (Fig. 2a2; green squares). During summer atmospheric $[NO_3^-]$ were higher than winter and spring, ranging from 19.9 ng m⁻³ to 264.0 ng m⁻³. In contrast to winter and spring, a distinct latitudinal trend was observed in summer whereby the- $[NO_3^-]$ decreased with increasing latitude (Fig. 2a2; orange circles) (Burger et al., 2022).

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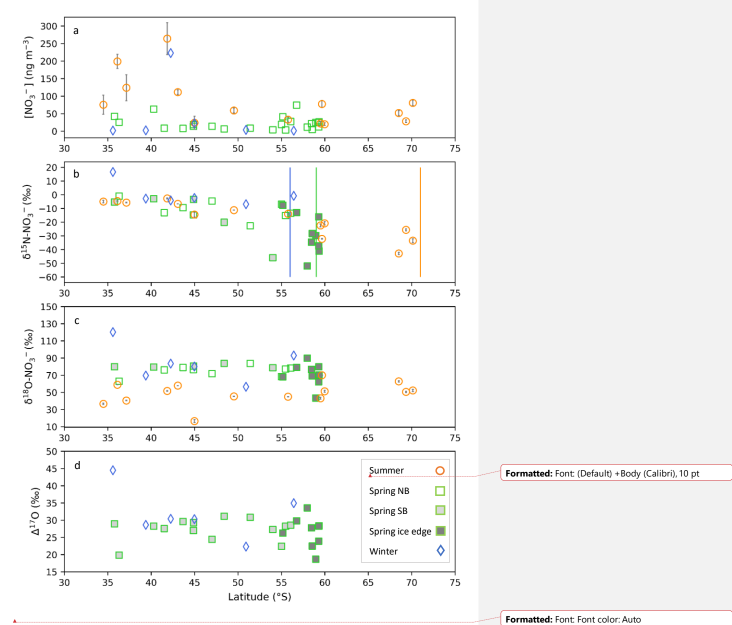




Figure 2. The average coarse mode (> 1 μm) atmospheric nitrate concentration [NO₃⁻] (ng m⁻³) (a), weighted
average δ¹⁵N of atmospheric nitrate (δ¹⁵N-NO₃⁻ (‰ vs. N₂)) (b), δ¹⁸O of atmospheric nitrate (δ¹⁸O-NO₃⁻ (‰ vs. N₂)) (b), δ¹⁸O of atmospheric nitrate (δ¹⁸O-NO₃⁻ (‰ vs. N₂)) (d) as a function of latitude (° S). Winter, spring

and summer are denoted by blue diamonds, green squares, and orange circles, respectively. For the summer data,

290 where error bars $(\pm 1 \text{ SD})$ are not visible, the standard deviation is smaller than the size of the marker. Spring data 291 are separated into northbound (NB), southbound (SB) and ice edge legs by clear, light grey and dark grey fills, 292 respectively for panels b-d. Vertical lines indicate the approximate location of the sea ice edge in summer (orange), 293 winter (blue) and spring (green), identified visually using satellite derived sea ice concentration obtained from 294 passive microwave sensors AMSR2 (Advanced Microwave Scanning Radiometer 2; Spreen et al., 2008). The 295 average coarse mode (> 1 μ m) atmospheric nitrate concentration [NO₃-] (ng m⁻³) as a function of latitude (° S). 296 Winter, spring and summer are denoted by blue diamonds, green squares, and orange circles, respectively. For the 297 summer data, where error bars (\pm 1 SD) are not visible, the standard deviation is smaller than the size of the 298 marker.

299 The seasonal cycle in atmospheric [NO₃⁻] that we observe, i.e., lowest concentrations in winter, higher in spring 300 and highest in summer, is similar to previous observations for the region. Atmospheric $[\mathrm{NO}_{3^-}]$ ranging from tens 301 of ng m⁻³ to approximately 100 ng m⁻³ have been observed for the Southern Ocean MBL during late spring (Morin 302 et al., 2009; Shi et al., 2021) and observations from a coastal Antarctic sites in the Atlantic sector at east Antarctica 303 showed elevated $[NO_3^-]$ ($\geq \sim 2040 \text{ to } 70 \text{ ng m}^{-3}$) inthroughout late spring and to early summer (Wagenbach et al., 304 1998; Wolff et al., 2008Shi et al., 2022). Seasonal studies at coastal and inland Antarctic sites observed the lowest 305 [NO3-] during winter (Wagenbach et al., 1998; Savarino et al., 2007; Wolff et al., 2008; Ishino et al., 2017; Walters 306 et al., 2019).

307 The seasonality in atmospheric [NO₃⁻] is largely driven by the seasonality in sunlight availability. Maximum 308 atmospheric [NO3-] observed in late spring/early summer in coastal Antarctica were attributed to reactive N 309 released from the post depositional processing/recycling of snow NO3- (Savarino et al., 2007). After NO3- is 310 deposited to the snowpack, it can be photochemically reduced to NO_x, and (re)emitted to the overlying atmosphere 311 (Jones et al., 2000; Jones et al., 2001). During winter, extended periods of darkness lead to reduced photochemical 312 activity above the snow, resulting in background level [NO₃-] (Lee et al., 2014). Over the open ocean, increased 313 UV radiation in spring and summer compared to winter may lead to greater NO3- production from the 314 photolytically derived oceanic RONO2 source (Fisher et al., 2018). Ground-based studies at various Antarctic 315 sites in Antarctica demonstrate that UV radiation is highest increases inthroughout spring and and is highest in 316 early summer, when stratospheric O3 concentrations are at a minimum and the noon solar zenith angle is low (Aun 317 et al., 2020; Lakkala et al., 2020; Li et al., 2020). This, in addition to greater lightning NO_x production during 318 spring and summer at the lower southern latitudes (< 40 °S) (Nesbitt et al., 2000) likely explain why higher [NO₃⁻] 319 are observed in spring and summer as compared to winter.

320 3.2 Seasonal variation in NO_x sources

 $\label{eq:second} 321 \qquad \text{While [NO_3^-] provides valuable information regarding the seasonal and spatial variation in the quantity of}$

 $\label{eq:sources} 322 \qquad \text{tropospheric NO}_{3^-} \text{ present, the N isotopic composition serves as a useful tool for identifying NO}_x \text{ sources that lead}$

- 323 to aerosol NO₃⁻ formation. Here, we present and interpret the mass weighted coarse-mode average δ^{15} N-NO₃⁻,
- computed for each filter deployment. In remote environments where O₃ concentrations largely exceed NO₄.
- 325 concentrations, as is the case for the remote Southern Ocean, NO_g isotopic exchange occurs at a much slower rate
- than the Leighton Cycle reactions. Therefore, little to no equilibrium isotope fractionation is expressed, and the
- 327 $\delta^{15}N \text{ of } NO_3^{-1}$ is assumed to reflect the $\delta^{15}N$ of the $NO_{\underline{s}}$ source (Walters et al., 2016).

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328 3.2.1 Evidence for stratospheric NO_{3-x}

329 There was one unusually high δ^{15} N-NO₃⁻ value equivalent to 16.6‰ for the first filter deployment of the 330 southbound leg in winter. Despite an elevated $\delta^{15}N$ signature, its [NO₃⁻] was consistent with that of most 331 wintertime samples. The δ^{15} N-NO₃⁻ of this wintertime sample is similar to the δ^{15} N of stratospherically sourced 332 NO_{3} , estimated to be $19 \pm 3\%$ (Savarino et al., 2007). Stratospheric input is additionally supported by the air 333 mass history of this sample, which indicates that air originated from as far south as the sea ice edge for the duration 334 of the sample deployment (Fig. 1a). Near the sea ice edge, some AMBTs originate from greater heights (300 to 335 400 m) and descend towards the sampling location (< 100 m) (Fig. S1).s. Coastal Antarctic studies suggest that 336 the deposition of PSCs during winter results in stratospheric NO3- inputs to the Antarctic troposphere (Wagenbach 337 et al., 1998; Savarino et al., 2007). Winter, when this sample was collected is the only time of year when Antarctic 338 temperatures are expected to be cold enough (<195 K) for polar stratospheric cloud (PSC) formation (von Savigny 339 et al., 2005; Wang et al., 2008).

340 Furthermore, this sample is unique in that it has a relatively high δ^{18} O-NO₃⁻ and Δ^{17} O-NO₃⁻, 120.2‰ and 44.5‰, 341 respectively. Tropospheric oxidation typically produces Δ^{17} O-NO₃⁻ values ranging from 17.3‰ to 42.7‰ (Morin et al., 2011; Ishino et al., 2017; Walters et al., 2019). Stratospheric sourced Δ^{17} O-NO₃⁻ is elevated in comparison 342 343 to tropospheric Δ^{17} O-NO₃⁻ because stratospheric O₃ has a greater isotope anomaly than tropospheric O₃, and/or 344 dominance of the N2O5 and ClONO2 pathways allow for greater transfer of the anomaly to NO3- via O3 (Savarino 345 et al., 2007; McCabe et al., 2007). High Δ^{17} O-NO₃⁻ (> ~40‰) observed in winter, are often attributed to 346 contributions by stratospheric denitrification (Savarino et al., 2007; McCabe et al., 2007; Frey et al., 2009; Walters et al., 2019) The combination of elevated Δ^{17} O-NO₃⁻ and δ^{15} N-NO₃⁻ is consistent with a stratospheric NO₃⁻ source 347 348 for this sample. Given the evidence that this sample likely does not reflect tropospheric oxidation chemistry, it is 349 left out of the below analysis.

350 3.2.2 Transported NO_x

365

351 $Previous \ \underline{modelling} \ studies \ \underline{suggesthave \ shown} \ that \ tropospheric \ transport \ of \ NO_x \ \underline{emitted} \ in \ the \ from \ the \ mid \ to$ 352 low latitudes (i.e., soil, lightning, thermal decomposition of peroxyacetylnitrate (PAN) and fossil fuel 353 combustion), contributes to the Antarctic NO3⁻ budget in winter (Lee et al., 2014; Shi et al., 2018). PAN 354 decomposition has previously been suggested as a NOx source to coastal Antarctica during winter and early spring 355 (Savarino et al., 2007; Jones et al., 2011). However, transported NO_x results in minimal NO₃-, regarded as 356 background level concentrations (Lee et al., 2014), consistent with most of our winter observations (Fig. 2a2; blue 357 diamonds). During winter, δ^{15} N-NO₃⁻ was relatively invariant across the Atlantic Southern Ocean (Fig. 2b³; blue 358 diamonds) with an average of $-3.4 \pm 2.1\%$ (n = 5). This is consistent with a lack of snowpack NO_x emissions at 359 the high latitudes during July/August due to weak or absent solar radiation (Shi et al., 2022). Furthermore, air 360 mass back trajectory analyses indicate that sea ice had a very minor influence on the winter samples (Fig. 1a &b). 361 Figure 3. The weighted average δ^{45} N of atmospheric nitrate (δ^{45} N NO₂⁻(% vs. N₂)) as a function of latitude (0 S). 362 Winter, spring and summer are denoted by blue diamonds, green squares, and orange circles, respectively. For the

summer data, where error bars (± 1 SD) are not visible, the standard deviation is smaller than the size of the
 marker. Spring data are separated into northbound (NB), southbound (SB) and ice edge legs by clear, light grey

and dark grey fills, respectively. Vertical lines indicate the approximate location of the sea ice edge in summer

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(orange), winter (blue) and spring (green), identified visually using satellite derived sea ice concentration obtained
 from passive microwave sensors AMSR2 (Advanced Microwave Scanning Radiometer 2; Spreen et al., 2008).

Albeit outside of the winter months, previous studies report an average δ^{15} N-NO₃⁻ for the low latitude Atlantic Ocean (between 45° S and 45° N) on the order of -3 to -4‰ (Baker et al., 2007; Morin et al., 2009), attributed to a combination of natural NO_x sources including lightning, biomass burning and soil emissions (Morin et al., 2009). This is also similar to the spring observations where higher values of δ^{15} N-NO₃⁻ were observed at the lower latitudes (-3.2 ± 1.8‰; n = 3). As such, the winter samples and low-latitude spring samples could be representative of a combination of natural NO_x sources emitted further north and transported to the mid-to-low latitude Atlantic Ocean.

375 Not all winter samples isotopically indicative of the transported background NOx source had low [NO3-]. One 376 unusually high $[NO_3^-]$ value (222.9 ng m⁻³) was observed in at the lower latitudes (Fig. 2a2; blue diamonds). Due 377 to the similarity in isotopic composition among winter samples, we can assume that despite a higher $[NO_3]$, this 378 sample also originated from a combination of natural NO_x sources transported from the lower latitudes. 379 Furthermore a [NO3-] on the order of 200 ng m-3 is consistent with summertime [NO3-] observations (Fig 2a2; 380 orange circles), when natural NOx sources dominated (see section 3.2.3). Our results thus confirm that like in 381 summer, natural NO_x sources can at times lead to relatively high [NO₃⁻], even in winter when background 382 conditions are typically experienced.

In addition, the winter dataset presented here clearly highlights the utility of the isotopes in distinguishing NO_x
 sources. The initial winter sample had a low concentration indicative of the background conditions; however, the
 triple stable isotopic composition of the sample confirms that it originated from the stratosphere (see sect. 3.2.1).
 In contrast, the anomalously high [NO₃⁻] sample observed in winter was not consistent with minimal background

 $\label{eq:source} 387 \qquad NO_x \mbox{ emissions, however its } \delta^{15}N \mbox{ confirmed that this was in fact the most likely source.}$

388 3.2.3 Snowpack photolysis and oceanic NOx sources

Springtime δ^{15} N-NO₃⁻ ranged from -52.0‰ to -1.1‰ and samples with the lowest δ^{15} N-NO₃⁻ were observed at the high latitudes (Fig. <u>2b</u>3; green squares). The range in δ^{15} N-NO₃⁻ observed for spring is consistent with late spring/early summer (November to December) observations from the Indian Ocean sector of the Southern Ocean (Shi et al., 2021) and summer (December and March) observations from the Atlantic sector (Burger et al., 2022). Springtime δ^{15} N-NO₃⁻ is also consistent with long-term records of δ^{15} N-NO₃⁻ measured at coastal Antarctica (Wagenbach et al., 1998) and on the east Antarctic Plateau (Winton et al., 2020), for the same season. Given the similarity in δ^{15} N-NO₃⁻ between spring and summer we expect the dominant NO_x sources to be the same.

396 During spring, air mass back trajectories indicate substantial sea ice influence at the high latitudes during the 397 southbound leg and during the ice edge transect (Fig. 34a & c). There is a large isotope effect associated with 398 snow NO3⁻ photolysis during summer in the Antarctic (Berhanu et al., 2014, 2015; Frey et al., 2009; Erbland et 399 <u>al., 2013</u>), resulting in the emission of low δ^{15} N-NO_x (~ -48‰) to the overlying atmosphere (Savarino et al., 2007; Morin et al., 2009; Shi et al., 2018; Walters et al., 2019). The low δ^{15} N-NO₃⁻ samples from the high latitudes 400 401 (minimum -52.0‰) are clearly influenced by sea ice (Fig. 2b3, Fig. 34a &c), but the air masses do not cross the 402 Antarctic continent. This suggests that the low δ^{15} N-NO_x likely comes from snow nitrate photolysis from the snow 403 on sea ice, before a net loss of NO32 from the snowpack leads to any large 15N enrichment in the snow and

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404 subsequently the atmosphere (Shi et al., 2018). We conclude that NO_x as a result of photolysis of snow nitrate on 405 sea ice can explain the relatively low $\delta^{15}N-NO_3^-$ observed in samples collected at the high latitudes on the spring 406 southbound leg and during the ice edge transect (Fig. 2b grey filled squares)."

407 This suggests that the low δ^{15} N NO_{*} likely comes from snow nitrate photolysis from the snow on sea ice. We

408 conclude that NO₃⁻ photolysis from snow on sea ice can explain the relatively low δ^{15} N NO₃⁻ observed in samples

409 collected at the high latitudes on the spring southbound leg and during the ice edge transect (Fig. 3 grey filled410 squares).

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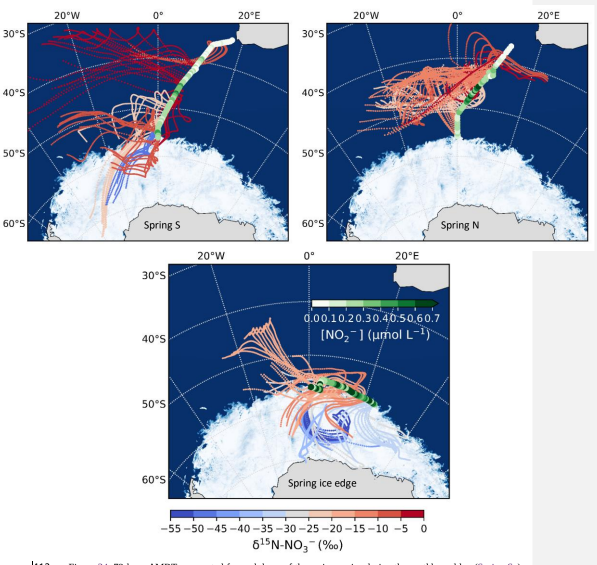


Figure <u>3</u>4. 72-hour AMBTs computed for each hour of the spring cruise during the southbound leg (Spring Sa),
the northbound leg (Spring Nb) and the ice edge transect (Spring ice edgee), when the HV-AS was running for
more than 45 min of the hour. AMBTs are colour coded by the weighted average δ¹⁵N-NO₃⁻, represented by the
blue to red colour bar. Overlaid are the surface ocean nitrite concentrations ([NO₂⁻]), represented by the green
colour bar. The white area represents the location of the sea ice determined using satellite -derived sea ice
concentration data obtained from the passive microwave sensors ASMR2 (Advanced Microwave Scanning
Radiometer2; Spreen et al., 2008).

421 Higher δ^{15} N-NO₃⁻ values (-22.7 to -1.0‰) were observed during spring for the northbound leg (Fig. 2b open 422 squares; Fig. 3b). The $\delta^{15}N$ of atmospheric NO₃⁻ that originates from snowpack emissions, depends on the $\delta^{15}N$ 423 of the local snowpack NO_x source. ¹⁵N enrichment in the snow due to preferential loss of ¹⁴N during photolysis₇ 424 NO3-loss, can eventually lead to increased 615N-NOx-via photolysis, and ultimately higher values of atmospheric 425 δ^{15} N-NO₃⁻ (Shi et al., 2018). However, the air mass histories of the samples with air mass histories that indicate 426 no contact with surrounding sea ice (i.e., the northbound leg; Fig. 2b3 open squares; Fig. 34b), suggesting that 427 limited-any influence from snowpack NOx emissions was limited. These samples originated from over the mid-428 latitude region of the Southern Ocean where detectable sea surface nitrite was present (Fig. 34b). The NO which 429 originates from derived from seawater nitrite in seawater is thought to limit sea surface RONO2 production, As a 430 result, such that non-zero elevated nitrite concentrations are required for RONO2 production to occur in seawater 431 (Dahl & Saltzman 2008; Dahl et al., 2012). Oceanic RONO2 has been long proposed as an important primary 432 NO: source to the Antarctic (Jones et al., 1991). Recent studies have used modelling and isotopic approaches to 433 investigate the relative importance of oceanic RONO2 compared to other sources of NO3- in the Southern Ocean 434 MBL, particularly in summer recently been proposed as an important NO2⁻ source to the region in summer (Fisher 435 et al., 2018; Burger et al., 2022)... Hhowever, limited co-occurring ocean atmosphere measurements are available 436 to constrain the seasonality of the RONO₂ source. While δ^{15} N-RONO₂ has yet to be directly quantified, it was 437 recently estimated to have an average δ^{15} N signature of ~ -22‰ in the summertime Southern Ocean (Burger et 438 al.,2022) and -27.8‰ in the eastern equatorial Pacific (Joyce et al., 2022). Consistent with this relatively 439 isotopically light oceanic RONO₂ source are observations of relatively low aerosol δ^{15} N-NO₃⁻ from the mid-440 latitude Southern Ocean (Burger et al., 2022) and eastern equatorial Pacific (Kamezaki et al., 2019; Joyce et al., 441 2022), on the order of -15 to -7‰.

Trends in δ^{15} N-NO₃⁻ by air mass origin were most evident in the ice edge transect during which lower (higher) δ^{15} N-NO₃⁻ values were observed for samples with greater sea ice (oceanic) influence (Fig. <u>3</u>4c). The photolysis imprint on the NO₃⁻ stable isotope signal in the marine boundary layer above the sea ice is clearly observed and speaks to the importance of snow-covered sea ice as a NO_x source in the region during spring as well as summer. The increased importance of oceanic RONO₂ emissions as air mass origin migrates from sea ice covered to open ocean zones is also evidenced by the decrease in δ^{15} N-NO₃⁻ observed for air mass originating predominantly from over the ocean (Fig. <u>3</u>4c).

Isotopically, there is little evidence of RONO₂ emissions contributing to aerosol NO₃ in the winter samples. Reduced levels of UV radiation and minimal daylight hours (Fig. S $_{2}$ +) in winter likely hinders the contribution of the oceanic NO_x source to NO₃⁻ loading compared to spring/summer, despite detectable sea surface nitrite concentrations in winter (Fig. S $_{2}$ -). Additionally, photolysis in spring/summer serves to produce OH which is the primary oxidant for conversion of RONO₂-derived NO_x in the MBL (Fisher et al., 2018).

Some studies suggest that the photolysis of particulate NO_3^- (p- NO_3^-) associated with sea-salt aerosols in the MBL can serve as an important NO_x source (Zhou et al., 2003; Ye et al., 2016; Reed et al., 2017). However, the importance of this NO_x formation pathway remains unclear, with large variability in reported rates between studies (Ye et al., 2016; Reed et al., 2017; Kasibhatla et al., 2018; Romer et al., 2018). To our knowledge, there are no observations of p- NO_3^- photolysis from the Southern Ocean MBL, and the implications of this process on the isotopic composition of NO_3^- in the MBL have yet to be assessed. We know that NO_3^- photolysis in snow is Formatted: Superscript
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460 associated with a large fractionation, leading to the emission of isotopically light NOx while the remaining NO3-461 pool becomes enriched in ¹⁵N (eg., Frey et al., 2009; Berhanu et al., 2014; 2015; Shi et al., 2018). Thus, if the p-462 NO3⁻ we measured was affected by photolysis we would have expected to observe much higher or even positive 463 values of δ^{15} N-NO₃⁻ during spring and summer. Another scenario is that the p-NO₃⁻ we measured resulted from 464 the oxidation of NOx released by prior p-NO3⁻ photolysis. In this case, we would have expected to observe much 465 lower δ^{15} N-NO₃⁻ values over the open ocean, on par with those observed over the ice. Since neither of the above 466 scenarios matches the observations, we discount the potential influence of aerosol NO3⁻ photolysis as a significant 467 NOx source to the region during our study is unlikely.

468 Additionally, a strong anti-correlation fr = -0.86 is observed between $\delta^{15}N-NO_3^{-}$ and $\Delta^{17}O-NO_3^{-}$ for samples

469 <u>collected in spring which experience a greater than 75% sea ice influence (Fig. 4), determined based on air mass</u>

470 history. A similar relationship was observed at Dome C during summer (Erbland et al., 2013; Savarino et al.,

471 <u>2016</u>). Previous studies found that the production of enhanced Δ^{17} O-NO₃⁻ in polar regions is linked to the intensity

472 <u>of NO_x emissions from the snowpack (Moring et al., 2012; Savarino et al., 2016). The correlation between Δ^{17} O-</u>

473 <u>NO₃⁻ and δ^{15} N-NO₃⁻ could arise from an increased contribution of HONO photolysis to total OH production,</u>

474 which is co-emitted with NO_x from the snowpack (Grannas et al., 2007), and induces a greater ¹⁷O excess in OH

476 of snowpack emissions with reactive halogen chemistry as suggested by Morin et al. (2012). The $\Delta^{17}O/\delta^{15}N$

477 <u>relationship presented here for the spring samples with air mass histories that indicate extensive influence from</u>

478 snow covered sea ice, suggests that snowpack emissions may lead to enhanced Δ^{17} O transfer to NO₃

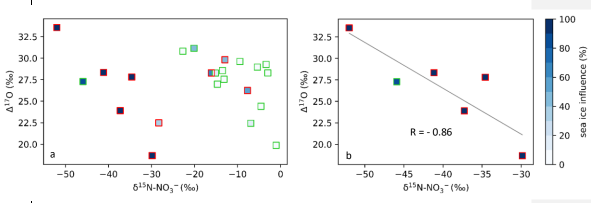
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Figure 4. The relationship between Δ¹⁷O-NO₃⁻ and δ¹⁵N-NO₃⁻ in spring (square symbols). In both panels, samples
collected along the ice edge are denoted by the red edge colour, with all other samples collect on the north and
southbound legs of the voyage denoted by the green edge colour. The colour bar (blues) indicates the percentage
sea influence experienced by each filter sample as determined using AMBTs. In panel a, all spring samples are
included. In panel b, only samples that experienced a sea ice influence > 75% are included. A straight line (grey)
is fitted to the data in panel b. Note the difference in x axis scale between panels.

- 485
- 486

487 3.3 Seasonal variation in oxidation

As mentioned in Sect. 1.1, the oxidation of NO and NO₂ can be determined using the oxygen isotopic composition of aerosol NO₃⁻. Here, we present and interpret the mass weighted coarse-mode average δ^{18} O-NO₃⁻ and Δ^{17} O-NO₃⁻, computed for each filter deployment.

491 During NO and NO₂ oxidation, the oxygen atoms of the responsible oxidants are incorporated into the NO₃ 492 product. The transferrable terminal oxygen atom of O_3 possesses an elevated $\Delta^{17}O$ -NO₃⁻(O_{3term}) and $\delta^{18}O$ (O_{3term}) 493 $(39.3 \pm 2\%)$ and $126.3 \pm 11.9\%$, respectively) (Vicars & Savarino, 2014), compared to other oxidants (e.g., OH 494 and peroxy radicals (RO₂/HO₂)) which possess a Δ^{17} O-NO₃⁻ \approx 0‰ (Michalski et al., 2011). The δ^{18} O of OH is 495 negative while the δ^{18} O of RO₂/HO₂ stems from that of atmospheric O₂ which is also low (23.9%; Barkan and 496 Luz 2005). These differences allow us to qualitatively multively assess NO and NO2 oxidation chemistry 497 involving contributions by various oxidants.7 Similarly to previous work conducted in the Southern Ocean MBL and in Antarctica (Walters et al., 2019; Shi et al., 2021), we make the assumptionas prior work suggests that 498 499 oxidant $\delta^{18}O$ values are known and directly represented in the NO₃-.

500 The relatively low δ^{18} O-NO₃⁻ values observed in summer (< 70%; Fig. <u>2c</u>5) are consistent with NO₂ oxidation via

501 OH (Burger et al., 2022). During summer, unusually low δ^{18} O-NO₃⁻ values were also observed equating to less 502 than the minimum expected for the OH oxidation pathway (< ~46‰; Burger et al., 2022). This was attributed to 503 an increased contribution by HO₂/RO₂ during NO oxidation to NO₂ (as opposed to O₃) which would decrease the 504 δ^{18} O of the product NO₃⁻. Increased abundance of RO₂ in the MBL was attributed to RONO₂ photolysis, 505 hypothesized to occur over the mid latitude Southern Ocean (Fisher et al., 2018; Burger et al., 2022), and/or the 506 presence of sea ice, which can lead to enhanced peroxy radical production (Brough et al., 2019).

Figure 5. The weighted average δ¹⁴O of atmospheric nitrate (δ¹⁴O NO₃⁻ (‰ vs. VSMOW)) as a function of latitude
(^o S). Winter, spring and summer are denoted by blue diamonds, green squares, and orange circles, respectively.
Spring data are separated into northbound (NB), southbound (SB) and ice edge legs by clear, light grey and dark
grey fills, respectively. For the summer data, where error bars (± 1 SD) are not visible, the standard deviation is
smaller than the size of the marker.

512 Interestingly, despite NO_x sources being the same in spring and summer (sect. 3.2), the δ^{18} O-NO₃⁻ data suggest 513 that the NO₃⁻ formation pathways differ (Fig. 2c5). Higher average δ^{18} O-NO₃⁻ values were observed in spring 514 compared to summer (Fig. $\underline{2c5}$). <u>Higher δ^{18} O-NO₃⁻ values in spring compared to summer may originate from NO_x</u> 515 oxidation by XO. In the Antarctic boundary layer, enhanced levels of BrO occur in spring; over sea ice covered 516 areas (Theys et al., 2011). The production of inorganic bromine has been proposed to be related to frost flowers 517 on thin sea ice (Kaleschke et al., 2004) and blowing of saline snow on sea ice (Yang et al., 2010). Significant 518 interaction with sea ice cover was experienced in spring, particularly at the ice edge transect, which could have 519 promoted NO3⁻ formation via the BrO pathway, resulting in increased values of $\delta^{18}O$ -NO3⁻.

520 <u>TIndeed</u>, the oxygen isotopic composition of NO₃⁻ in winter and spring were comparable as indicated by both

521 $\delta^{18}O$ (Fig. <u>2c</u>5) and $\Delta^{17}O$ (Fig. <u>2d</u>83). The $\delta^{18}O$ -NO₃⁻ ranged from 56.5‰ to 92.9‰ in winter (Fig. <u>2c</u>5; blue

diamonds) and from 62.3% to 89.8% in spring (Fig. $2c^5$; green squares). The Δ^{17} O-NO₃ ranged from 22.3% to

523 35‰ in winter (Fig. 2dS3; blue diamonds) and from 18.7‰ to 33.6‰ in spring (Fig. 2dS3; green squares).

524 Interestingly, there is more variability in the δ^{18} O and Δ^{17} O for the ice edge transect (Fig. <u>2c</u>5 & <u>2d</u>S3; dark shaded 525 squares) than the north and southbound transects. The overlap in $\delta^{18}O$ and $\Delta^{17}O$ in winter and spring suggest that 526 similar pathways lead to NO3- formation in both seasons, i.e., oxidation pathways that result in an increased 527 influence of O3 during oxidation (i.e., N2O5, DMS, XO) .-

528 A significant linear relationship was observed between δ^{18} O-NO₃⁻ and Δ^{17} O-NO₃⁻ in both winter and spring (Fig. 529 S4). This suggests isotopic mixing between two major oxidants (e.g., Fibiger et al., 2013; Shi et al., 2021). As 530 such, the highest end-member is representative of tropospheric O₃, and/or XO with a δ^{18} O of ~114 to 138‰ and 531 a $\Delta^{17}O$ of ~39‰. There are multiple options for the second oxidant with a $\Delta^{17}O = 0$ ‰, i.e., water vapour (H₂O_(v)), 532 OH, and O₂. Here, we use the δ^{18} O-H₂O_(y) from the average of observations along a similar cruise transect from 533 the Indian sector of the Southern Ocean (Dar et al., 2020). The average $\delta^{18}O-H_2O_{(v)}$ determined between ~33° S 534 and $\sim 60^{\circ}$ S (-13.9 \pm 1.4‰) was used for the winter samples given that AMBTs indicate that most air masses 535 originated within this latitudinal band, where there is minimal variation in $\delta^{18}O-H_2O_{(v)}$ (Dar et al., 2020). In 536 spring, the zone of air mass origin for our samples extends further south to ~70° S. As shown by Dar et al. (2020), 537 δ^{18} O-H₂O_(y) declines significantly between ~60° S and ~70° S. To account for this lowering in δ^{18} O-H₂O_(y) which 538 could influence higher latitude samples, an additional H₂O_(v) end member equivalent to the minimum observed 539 by Dar at al., 2020 (-27.5‰) was included for spring. The δ^{18} O-OH was calculated from the equilibrium 540 fractionation between OH and $H_2O_{(v)}$ (Walters & Michalskii et al., 20161) using the observed atmospheric 541 temperature range for winter and spring. The δ^{18} O-OH determined for winter ranges from -56.2 to -49.5‰ 542 (average = -52.8‰) and for spring ranges from -54.5 to -50.5‰ (average = -52.5‰). Therefore, a value of -5322%543 was used for both seasons. The atmospheric δ^{18} O-O₂ is well-constrained at 23.9‰ (Barkan and Luz 2005). The 544 δ^{18} O and Δ^{17} O values assumed for all oxidants or oxygen sources outlined above, are summarised in Table 1. 545 Mixing lines for the three oxidant pairs (OH/O₃, $H_2O_{(v)}/O_3$ and O_2/O_3) are indicated by the grey, orange and red 546 lines, respectively in Figure 6.

547 Table 1: A summary of the oxygen isotope ratios (δ^{18} O and Δ^{17} O) for the end member oxidants and/or oxidant 548 sources (O₃, OH, HO₂/RO₂ and H₂O) utilised in Sect. 3.3.

	a10 a (a)		. 17	
Oxidant/source	$\delta^{18}O(\%)$	References	$\Delta^{1}O(\%)$	References
Terminal O ₃	126.3 ± 11.9	Vicars & Savarino	<u>39.3 ± 2</u>	Vicars & Savarino
		<u>(2014)</u>		(2014)
OH	-52.7 ± 2.8^{a}	Walters &	~0	Michalski et al.
		Michalski (2016)		(2011)
HO_2/RO_2	23.88 ± 0.03	Barkan & Luz	~0	Michalski et al.
		(2005)		(2011)
H_2O	<u>-13.9 ± 1.4</u>	Dar et al. (2020)	~0	Michalski et al.
	_			(2011)

549 550 ^aThe average δ^{18} O-OH was calculated from the equilibrium fractionation between OH and H₂O_(v) (Walters & Michalski, 2016) using the observed atmospheric temperature range for winter and spring and the average $\delta^{18}O$ -551

H₂O (Dar et al., 2020).

552 To determine the lower endmember in each season, i.e., the second major oxidant in addition to ozone and/or XO, 553 a straight line was fitted to the data in $\delta^{18}O$ - $\Delta^{17}O$ space and the x-intercept at a $\Delta^{17}O = 0\%$ was determined. The 554 x-intercept in winter is ~-16‰. During winter, the linear relationship observed (Fig. 56a) is similar to what has 555 been seen in the Indian Ocean MBL and at coastal east Antarctica, where the x intercept was $-11 \pm 8\%$ (Shi et al., 556 2021) and $-15 \pm 6\%$ (Shi et al., 2022), respectively. The oxygen isotopic composition of the lower end member 557 in our winter data is most similar to that of $H_2O_{(v)}$. This is consistent with the average $\delta^{18}O-H_2O_{(v)}$ (= -13.96 ±

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1.45‰) observed between approximately 33° S and 60° S (Dar et al., 2020). Therefore, a mixing line between
H₂O_(v) and O₃ is the best fit to the winter observations (Fig. 56a; solid orange line). If we exclude an equilibrium
isotope fractionation between OH and H₂O_(v) (Michalski et al., 2011) such that δ¹⁸O-OH is similar to the δ¹⁸O of
H₂O_(v), then the lower end-member likely results from the OH oxidation pathway.

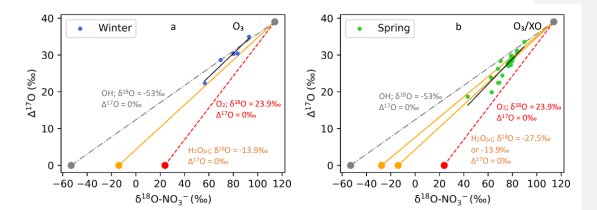


Figure 5. Winter and spring δ^{18} O-NO₃⁻ vs. Δ^{17} O-NO₃⁻ are plotted in panels (a) and (b), respectively. A straights line (black) is fitted to the data in each panel. In both panels the grey line represents the OH/O₃ mixing line, the orange line represents the H₂O_(v)/O₃ mixing line and the red line represents the O₂/O₃ mixing line. In panel b, an additional H₂O_(v)/O₃ mixing line is included (also in orange) to account for potentially lower values of δ^{18} O-H₂O_(v) (~ -27.5 ‰) at 60 ° to 70 °S.-

Figure 6. Winter and spring δ⁴⁸O-NO₃⁻-vs. Δ¹⁷O-NO₃⁻ are plotted in panels (a) and (b), respectively. A straight
line (black) is fitted to the data in each panel. In both panels the grey line represents the OH/O₃ mixing line, the
orange line represents the H₂O_{(v}/O₃ mixing line and the red line represents the O₂/O₂ mixing line.

570 By contrast, observations made in spring are best represented by mixing between 3 major oxidants: $H_2O_{(v)}$, O_3 and 571 O_2 . The x-intercept in spring is ~-4‰, making it more difficult to identify one low $\delta d^{18}O$ end member. The oxidant 572 source with the closest oxygen isotope composition is again H2O(v), indicating the prevalence of the OH pathway 573 (when δ^{18} O-OH ~ δ^{18} O-H₂O_(v)), however the x-intercept is greater in spring compared to winter, suggesting that 574 the lower end member has a higher δ^{18} O. H₂O_(v) data from the region suggest that we would not expect to see a $\delta^{18}O > -10\%$, therefore an increase in $H_2O_{(v)}\delta^{18}O$ from winter to spring can be ruled out. A more likely explanation 575 576 is that the springtime lower endmember consists of some combination of $H_2O_{(v)}$ and an additional higher $\delta^{18}O$ 577 oxidant that is less abundant in winter. The higher $\delta^{18}O$ oxidant is likely atmospheric O_2 ($\delta^{18}O = 23.9\%$, $\Delta^{17}O = 23.9\%$). 578 0% vs VSMOW; Barkan and Luz, 2005). This is consistent with the spread in the springtime observations, which 579 are bound by the decreased $H_2O_{(v)}/O_3$ mixing line to accommodate the influence of lower $\delta^{18}O-H_2O_{(v)}$ at the high 580 latitudes (Fig. 56b; orange line), and the atmospheric O2/O3 mixing line (Fig. 56b; red line).

The influence of atmospheric O₂ during spring likely results from the increased role for RO₂ (and/or HO₂) in NO_x cycling. This may be linked to increased RO₂ production over the mid-latitude Southern Ocean, derived from

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583 RONO₂ photolysis in the MBL (Burger et al., 2022). There is also evidence that sea ice can lead to enhanced 584 peroxy radical production (Brough et al., 2019), resulting in the potential for increased RO2 and HO2 585 concentrations to be observed in air masses that traverse the sea ice zone before being sampled. δ^{18} O-NO₃⁻ is 586 greater in winter and spring compared to summer (Fig. 5), highlighting the increased control of O_3 on the oxygen 587 isotopic composition of NO3- in winter and spring. Consistent with increased O3 influence are seasonally resolved 588 observations of O₃ concentration ([O₃]) at coastal Antarctica (Ishino et al., 2017; Shi et al, 2022) and Cape Grim, 589 Tasmania (Derwent et al., 2016), the latter being more representative of the MBL. In all cases, maximum [O₃] are 590 observed in winter, and minimum [O₃] are observed throughout summer. In spring, [O₃] concentrations are 591 noticeably reduced compared to the winter, but slightly elevated compared to summer. Higher & 80-80-30- values 592 in spring may also originate from NOx oxidation by XO, for example BrO as discussed in Section 3.3.

593

594 4 Conclusions

Seasonally resolved observations of atmospheric NO_3^- across the Atlantic Southern Ocean MBL suggest that natural NO_x sources dominate throughout the year. Similar NO_3^- sources are available to the MBL in both spring and summer, highlighting the importance of oceanic RONO₂ emissions in seasons other than the more frequently sampled summer months in the Southern Ocean. Although further research is required to improve our mechanistic and isotopic understanding of oceanic RONO₂ formation, fluxes and conversion to aerosol NO_3^- , this work contributes to our growing understanding of how the surface ocean influences the atmospheric reactive N cycle and oxidation chemistry of the MBL (Altieri et al., 2021; Burger et al., 2022; Joyce et al., 2022).

602 Furthermore, the large sea ice extent characteristic of spring highlights the importance of snow-covered sea ice as 603 a NOx source, in addition to the well documented summer source from snow covered continental ice (Jones et al., 604 2001; Walters et al., 2019; Winton et al., 2020). Currently no measurements of δ^{15} N-NO₃⁻ from snowpack on sea 605 ice exist for Antarctica, which is an important measurement gap that should be addressed in future studies. The 606 presence of sea ice may also play a role in the formation of peroxy radicals through its influence on chlorine 607 chemistry when sunlight is available (Brough et al., 2019). Peroxy radicals (RO₂), H₂O_(v) and O₃ serve as the 608 dominant atmospheric oxidants during spring, responsible for aerosol NO3- formation. In contrast, a lack of 609 sunlight and sea ice influence is experienced during winter, and mixing between two end-members, H2O(y) and 610 O₃, best explain the oxygen isotopic composition of the NO₃⁻ that is formed. Similar to coastal Antarctic sites, 611 reduced daylight hours and/or increased O3 abundance in the winter and spring MBL lead to greater O3 influence 612 on NO3⁻ formation, compared to the summer when OH oxidation chemistry dominates.

613 Winter is characterised by very low $[NO_3^-]$ concentrations with $\delta^{15}N$ signatures that reflect background conditions 614 similar to that of the low latitude Atlantic Ocean (Morin et al., 2009). Interestingly, despite being collected off the 615 coast of South Africa, the N and O isotopic composition of NO₃⁻ measured for the first wintertime sample reflects 616 a stratospheric NO_{3-*} source signal. This is also supported by AMBTs that originate near Antarctica, where 617 stratospheric denitrification is reported to occur (Savarino et al., 2007).

Our observations highlight the potential power of N and O isotopes of nitrate in distinguishing between the various
 natural NOx sources that result in NO₃⁻ formation, and constraining formation pathways of aerosol NO₃⁻. In order

620 to improve the utility of the N and O isotopes in the polar atmosphere, more measurements of the isotopic 621 composition of the regional sources, e.g., snow on sea ice, and regional processes, e.g., OH from HONO and sea 622 ice oxidant emissions, is needed. Even though it is complex, the utility of the N isotopes in distinguishing between 623 the various natural NO_x sources that result in NO_3^- formation in the MBL of the Atlantic Southern Ocean, 624 especially in the less frequently sampled seasons of winter and spring is evident. Furthermore, the O isotopes 625 serve as a useful were able to help-tool for constraining formation pathways of aerosol NO₃ seasonally. This is -626 especially important in the Atlantic Southern Ocean where oxidation chemistry is poorly constrained (Beygi et 627 al., 2011). The contribution of sea ice to oxidant production when sunlight returns in spring is also highlighted 628 by the O isotopes. As such, these data may be useful to modelling efforts attempting to characterise N cycling 629 between the surface ocean and lower atmosphere, and may help improve atmospheric oxidant budgets that are 630 less understood in unpolluted low-NOx environments.

631 Author contributions. KEA designed the study and sampling campaign, acquired funding and supervised the 632 research. KEA and MGH provided financial and laboratory resources and assisted in data validation. EJ performed 633 laboratory analysis of samples at Brown University. KAMS and JMB conducted the sampling at sea and JMB 634 performed laboratory analysis at the University of Cape Town. JMB analysed the data and prepared the manuscript 635 with contributions from all co-authors. KEA, MGH and EJ assisted with reviewing and editing the manuscript.

636 Competing interests. One author is a member of the editorial board of journal ACP. The authors declare no other637 conflicts of interest.

638 Data availability. Datasets for this research are available at https://doi.org/10.5281/zenodo.7142722

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