The importance of alkyl nitrates and sea ice emissions to atmospheric NO_x sources and cycling in the summertime Southern Ocean marine

3 boundary layer

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11 Abstract. Atmospheric nitrate originates from the oxidation of nitrogen oxides ($NO_x = NO + NO_2$) and impacts both tropospheric 12 chemistry and climate. NO_x sources, cycling, and NO_x to nitrate formation pathways are poorly constrained in remote marine 13 regions, especially the Southern Ocean where pristine conditions serve as a useful proxy for the preindustrial atmosphere. Here, we measured the isotopic composition (δ^{15} N and δ^{18} O) of atmospheric nitrate in coarse-mode (> 1 µm) aerosols collected 14 15 in the summertime marine boundary layer of the Atlantic Southern Ocean from 34.5°S to 70°S, and across the northern edge 16 of the Weddell Sea. The δ^{15} N-NO₃⁻ decreased with latitude from -2.7‰ to -42.9‰. The decline in δ^{15} N with latitude is 17 attributed to changes in the dominant NO_x sources: lightning at the low latitudes, oceanic alkyl nitrates at the mid latitudes, 18 and photolysis of nitrate in snow at the high latitudes. There is no evidence of any influence from anthropogenic NO_x sources 19 or equilibrium isotopic fractionation. Using air mass back trajectories and an isotope mixing model, we calculate that oceanic 20 alkyl nitrate emissions have a δ^{15} N signature of -21.8‰ ± 7.6‰. Given that measurements of alkyl nitrate contributions to 21 remote nitrogen budgets are scarce, this may be a useful tracer for detecting their contribution in other oceanic regions. The 22 δ^{18} O-NO₃⁻ was always less than 70%, indicating that daytime processes involving OH are the dominant NO₃ oxidation pathway 23 during summer. Unusually low δ^{18} O-NO₃⁻ values (less than 31‰) were observed at the western edge of the Weddell Sea. The 24 air mass history of these samples indicates extensive interaction with sea ice covered ocean, which is known to enhance peroxy 25 radical production. The observed low δ^{18} O-NO₃⁻ is therefore attributed to increased exchange of NO with peroxy radicals, 26 which have a low δ^{18} O, relative to ozone, which has a high δ^{18} O. This study reveals that the mid- and high-latitude surface 27 ocean may serve as a more important NO_x source than previously thought, and that the ice-covered surface ocean impacts the 28 reactive nitrogen budget as well as the oxidative capacity of the marine boundary layer.

29 1 Introduction

30 Atmospheric nitrate (NO₃⁻), hereafter defined as gas phase nitric acid (HNO₃) and particulate NO₃⁻ (p-NO₃⁻), impacts air quality 31 and climate by contributing to atmospheric particulate matter (Park and Kim, 2005), and influencing the Earth's radiative heat 32 budget (IPCC, 2013). It also plays a major role in the biogeochemical cycling of reactive nitrogen (Altieri et al., 2021). NO₃⁻ 33 aerosols originate from the oxidation of nitrogen oxides, collectively referred to as NO_x ($NO_x = NO + NO_2$). NO_x cycling 34 controls the chemical production of tropospheric ozone (O_3) , a greenhouse gas and pollutant (Finlayson-Pitts and Pitts, 2000), 35 which in turn contributes to the oxidising capacity of the atmosphere (Alexander and Mickley, 2015). Globally, fossil fuel 36 combustion is the primary NO_x source (van der A et al., 2008), which far exceeds natural emissions such as biomass burning 37 (Finlayson-Pitts and Pitts, 2000), soil processes (Davidson and Kingerlee, 1997) and lightning (Schumann and Huntrieser, 38 2007).

39 Due to its remoteness, the summertime Southern Ocean (SO) marine boundary layer (MBL) can be representative of 40 preindustrial-like atmospheric conditions (Hamilton et al., 2014). The chemical composition of the Southern Ocean MBL is 41 characterised by low NO₃⁻ concentrations (Virkkula et al., 2006), representative of a background aerosol environment (i.e., 42 minimal anthropogenic influence). Furthermore, the South Atlantic sector of the Southern Ocean is primarily influenced by 43 natural NO_x sources. During summer, high lightning activity over South America and southern Africa results in NO_x production 44 between approximately 40° S and the intertropical convergence zone (ITCZ) (Nesbitt et al., 2000). As such, lightning is 45 expected to be the dominant NO_x source in the low latitude MBL (Schumann and Huntrieser 2007, van der A et al., 2008). 46 Because of its pristine nature, the summertime Southern Ocean serves as a unique region in which to study atmospheric 47 chemistry and is a useful preindustrial reference point for comparing the magnitude of anthropogenic aerosol impacts on 48 climate (Haywood and Boucher 2000; Hamilton et al., 2014).

49 The atmospheric chemistry of the polar MBL at the high southern latitudes differs from that of the mid- and low-latitude 50 MBL. During summer, high levels of photochemistry result in the emission of reactive gases from sea ice and snow cover in 51 the Antarctic. As a result, highly elevated concentrations of hydrogen oxide radicals ($HO_x = OH + peroxy radicals$), halogens, 52 nitrous acid (HONO), and NO_x have been observed during spring and summer in the polar regions (Brough et al., 2019). 53 Furthermore, photochemical production of NO_x within the surface snow of Antarctica and subsequent oxidation in the 54 overlying atmosphere represents a significant NO_3^- source to the Antarctic troposphere (Jones et al., 2000, 2001). NO_3^- 55 photolysis near the surface-air interface of ice crystals produces NO₂ (Grannas et al. 2007; Jones et al., 2000), which can be 56 released to the firm (i.e., the intermediate stage of ice between snow and glacial ice) air and escape the snowpack to the 57 overlying atmosphere (Erbland et al., 2013; Shi et al., 2015; Shi et al., 2018). During winter, additional NO_x sources to the 58 Antarctic atmosphere may include long-range transported peroxyacetyl nitrates (PAN) and stratospheric inputs (Savarino et 59 al., 2007; Lee et al., 2014; Walters et al., 2019).

Emission of alkyl nitrates (a group of nitrogen gases collectively referred to as RONO₂) from the surface ocean have been
 recently proposed as a potential NO_x source to the MBL in remote regions (Williams et al., 2014; Fisher et al., 2018).

62 Observations of elevated MBL alkyl nitrate concentrations suggest that a direct oceanic source exists in both the tropics (Atlas 63 et al., 1993; Blake et al., 2003), and the high-latitude Southern Ocean (Blake et al., 1999; Jones et al., 1999). Although the 64 exact mechanism remains unclear, experimental evidence suggests that oceanic RONO₂ production occurs via photochemical 65 processes involving the aqueous phase reaction of RO₂, derived from the photolysis of oceanic dissolved organic matter and 66 NO, derived from seawater nitrite photolysis (Dahl et al., 2003; Dahl and Saltzman, 2008). Supersaturated RONO₂ conditions in the surface ultimately drive a net flux from the ocean to the atmosphere (Chuck et al., 2002; Dahl et al., 2005). The photolysis 67 68 of emitted RONO₂ and subsequent OH oxidation in the overlying atmosphere leads to NO₃ formation (Fisher et al., 2018), and 69 /or RONO₂ can form aerosol NO₃⁻ directly by hydrolysis (Rindelaub et al., 2015).

70 Current global atmospheric models suggest that oceanic RONO₂ represents a significant source of nitrogen (N) to the 71 Southern Ocean MBL, accounting for 20% to 60% of the reactive N pool at the high-latitudes (60° S to 90° S) (Fisher et al., 72 2018). However, only one shipborne dataset with coincident ocean-atmosphere RONO₂ concentration measurements exists to 73 substantiate this notion (Hughes et al., 2008). Additionally, the NO_x source from RONO₂ degradation dominates relative to 74 model defined primary NO_x emission sources over the SO, which include shipping, aircraft and lightning (Fisher et al., 2018). 75 However, the lack of seawater observations available to constrain Southern Ocean RONO₂ distributions hamper the validation 76 of model fluxes. Better understanding of the Southern Ocean RONO₂ source is required to improve simulations and accurately 77 evaluate its contribution to the Southern Ocean MBL NO_x budget.

78 **1.1 Natural abundance isotopes of atmospheric nitrate**

Measurements of the oxygen (O) and N stable isotope ratios of atmospheric NO_3^- can be used to constrain NO_x sources, NO-NO₂ cycling, and NO_x to NO_3^- oxidation pathways, which are critical for understanding the reactive N budget in the atmosphere. This technique has been applied in polluted (Elliot et al., 2007; Zong et al., 2017), open ocean (Hastings et al., 2003; Morin et al., 2009; Kamezaki et al., 2019; Gobel et al., 2013; Altieri et al., 2013), and polar environments (Morin et al., 2009; Walters et al., 2019). Stable isotope ratios are reported as a ratio of the heavy to light isotopologues of a sample relative to the constant isotopic ratio of a reference standard, using delta (δ) notation in units of "per mil" (‰) following Eq. (1):

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$$\delta = \left(\left(R_{sample} / R_{standard} \right) - 1 \right) \times 1000$$

where *R* represents the ratio of ${}^{15}N/{}^{14}N$ or ${}^{18}O/{}^{16}O$ in the sample and in the reference standard, respectively. The reference for O is Vienna Standard Mean Ocean Water (VSMOW) and for N is atmospheric N₂ (Bölhke et al., 2003).

(1)

88 When NO_x is converted to NO₃⁻, the N atom is conserved. As such, it is generally expected that the N stable isotope ratio 89 of atmospheric NO₃⁻ (δ^{15} N-NO₃⁻) reflects the δ^{15} N of the source NO_x, (Kendall et al., 2007) plus any isotopic fractionation 90 associated with NO/NO₂ cycling or NO_x to NO₃⁻ conversion. For example, the δ^{15} N of lightning generated NO_x is close to 0‰ 91 (Hoering, 1957) and is distinct from stratospheric and snowpack NO_x. Savarino et al., (2007) used the degree of N₂O 92 destruction in the stratosphere and the associated isotopic fractionation to derive an Antarctic stratospheric δ^{15} N-NO_x source 93 signature of 19‰ ± 3‰ (Savarino et al., 2007). In contrast, snow emitted NO_x typically has a very low δ^{15} N signature due to

- 94 the large fractionation ($^{15}\varepsilon$) of ~ - 48‰ (Berhanu et al., 2014, 2015) associated with NO₃⁻ photolysis in the snowpack, where 95 $^{15}\varepsilon = (\text{KIE} - 1) \times 1000\%$ and the kinetic isotope effect (KIE) is the ratio of the rates with which the two isotopes of N are 96 converted from reactant to product. If equilibrium isotope fractionation during NO/NO₂ cycling occurs, it results in the ¹⁵N 97 enrichment of NO₂ such that the NO₃⁻ formed from this NO₂ will have a higher δ^{15} N-NO₃⁻ than the initial NO_x source (Frever 98 et al., 1993; Walters et al., 2016). Equilibrium isotope fractionation during the transformation of NO_x to NO₃⁻ also results in 99 higher δ^{15} N-NO₃⁻ compared to the original NO_x source (Walters and Michalski, 2015). 100 In contrast to N, the O stable isotope ratio of atmospheric NO₃⁻ (δ^{18} O-NO₃⁻) is reflective of the oxidants involved in NO₃ 101 cycling prior to NO_3^- formation as well as the dominant NO_3^- formation pathway (Hastings et al., 2003; Michalski et al., 2003; 102 Alexander et al., 2020). The O atoms of NO_x are rapidly exchanged with oxidising agents in the atmosphere to produce NO₃⁻. 103 Tropospheric NO_x recycles rapidly with O_3 following the equations below: 104 $NO + O_3 \rightarrow NO_2 + O_2$ (R1) 105 $NO_2 + O_2 + h\nu \rightarrow NO + O_3$ (R2) 106 The oxidation of NO to NO₂ requires an atmospheric oxidant, typically O_3 throughout most of the troposphere (R1), while the 107 breakdown of NO₂ back to NO is photolytic and requires light (R2). Therefore, under nighttime/dark conditions (R2) shuts 108 down and NO_x is comprised almost entirely of NO₂. 109 The dominant daytime sink for NO_x is the oxidation of NO_2 by OH, which produces nitric acid (HNO₃) via (R3), where M is 110 a non-reacting molecule. 111 $NO_2 + OH + M \rightarrow HNO_3 + M$ (R3) 112 Under nighttime/dark conditions, the photolytic production of OH cannot occur and NO₂ is oxidised by O_3 (R4). HNO₃ is 113 ultimately formed via the hydrolysis of dinitrogen pentoxide (N_2O_5), following the reactions (R5) and (R6) below: 114 $NO_2 + O_3 \rightarrow NO_3 + O_2$ (R4) 115 $NO_3 + NO_2 + M \rightleftharpoons N_2O_5(g) + M$ (R5) 116 $N_2O_5(g) + H_2O(l) + surface \rightarrow 2HNO_3(aq)$ (R6) 117 NO₃ can also react with hydrocarbons (HC) (e.g., dimethylsulphide (DMS)) to form HNO₃ following reaction (R7) below: 118 $NO_3 + HC/DMS \rightarrow HNO_3 + products$ (R7) 119 Lastly, in regions with elevated halogen concentrations, NO₂ can be oxidised by reactive halogens, for example bromine oxide 120 (BrO), to form HNO₃ following (R8) and (R9) below: 121 $NO_2 + BrO \rightarrow BrONO_2$ (R8) 122 $BrONO_2 + H_2O + surface \rightarrow HNO_3 + HOBr$ (R9) 123 Typically, aerosol δ^{18} O-NO₃⁻ is interpreted as being determined by the dominant NO_x oxidation pathways, (R3) versus 124 (R4) to (R9). If some combination of R4-R9 occurs, then O₃ is the main oxidant, whereas during (R3), one of the O atoms
- originates from OH. The OH radical exchanges with H_2O vapor in the troposphere, therefore the $\delta^{18}O$ of OH is a function of
- 126 the δ^{18} O of H₂O vapour, which generally ranges from -27.5% to 0% in the subtropics and over the Southern Ocean (Michalski
- 127 et al., 2012; Guilpart et al., 2017; Dar et al., 2020), and equilibrium isotope exchange between OH and H₂O (Walters and

128 Michalski, 2016). In contrast the δ^{18} O of tropospheric O₃ is much higher, the most recent estimate being 114.8±10.4‰ (Vicars

and Savarino, 2014). Therefore, a higher δ^{18} O for atmospheric NO₃⁻ reflects the increased influence of O₃ on NO_x to NO₃⁻

130 conversion (R4-R9), and the δ^{18} O-NO₃⁻ is lower when (R3) is favoured, due to the lack of exchange of O atoms with O₃ 131 (Hastings et al., 2003; Fang et al., 2011; Altieri et al., 2013).

Here, we present the concentration and isotopic composition of coarse mode (> 1 μ m) atmospheric NO₃⁻ collected in the MBL of the Southern Ocean between Cape Town, South Africa and coastal Antarctica, as well as across the Weddell Sea gyre, during summer. Using air mass back trajectories, surface ocean nitrite measurements, and the aerosol δ^{15} N- and δ^{18} O-NO₃⁻, we address 1) the major NO_x sources as well as the main oxidants in NO/NO₂ cycling and NO_x to NO₃⁻ conversion across a large latitudinal transect of the Atlantic Southern Ocean and within the Weddell Sea gyre, and 2) the influence of sea-ice and snowpack emissions on NO_x/NO₃⁻ chemistry in the high-latitude MBL.

138 **2) Methods**

139 **2.1**) Sample collection

140 Samples were collected on board the Research Vessel (R/V) SA Agulhas II during one cruise subdivided into three legs. Leg 141 one refers to the voyage south from Cape Town (33.9° S, 18.4° E) to Penguin Bukta (71.4° S, 2.5° W) in early summer (7 to 142 19 December 2018) as part of the South African National Antarctic Expedition's annual relief voyage (SANAE 58). Leg two 143 is the Weddell Sea Expedition (WSE) from 4 January to 21 February 2019. All data were recorded in GMT. The WSE refers 144 to the voyage west from Penguin Bukta to the northern edge of the Weddell Sea gyre to Larsen C ice shelf, followed by a 145 detour to King George Island before returning to the Weddell Sea and sailing back to Penguin Bukta. Leg three refers to the 146 SANAE 58 return voyage north from Penguin Bukta to Cape Town in late summer (27 February to 15 March 2019). From 147 here on, legs one, two and three will be referred to as early summer, the Weddell Sea, and late summer, respectively.

Size-segregated atmospheric 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 0.82 m³ min⁻¹ though a five-stage cascade impactor (TE-235; Tisch Environmental), loaded with combusted (400°C for 4 hours) glass fibre filters (TE-230-GF; Tisch Environmental) that have a surface area of approximately 119 cm². Aerosol nitrate in the MBL is predominantly present in the coarse mode (> 1 μ m), therefore only filter stages 1 through 4 were analysed, where the aerodynamical diameter of particles collected are as follows: stage 1 (> 7 μ m); stage 2 (3 to 7 μ m); stage 3 (1.5 to 3 μ m) and stage 4 (1 to 1.5 μ m).

A sector collector was used to restrict HV-AS activity to avoid contamination from ship stack emissions (Campbell Scientific Africa). The HV-AS only began operating if the wind was blowing at an angle less than 75° or greater than 180° from the bow of the ship for a minimum of ten minutes at a speed of at least 1 m s⁻¹. Filters were removed from the cascade impactor inside a laminar flow cabinet (Air Science), placed in individual zip-sealed plastic bags and stored at -20°C until analysis. Given that the MBL of the Southern Ocean is characterised by low atmospheric NO_3^- concentrations, an attempt was made to ensure that at least 24 hours of in-sector sampling had passed before filters were removed from the cascade impactor. However, this was not always possible as on occasion the filters had to be removed early to avoid contamination due to unusual ship manoeuvres or stagnant conditions. Therefore, sampling times ranged between 13 and 88 hours across the three legs. The details of each cruise leg can be found in the supplemental information (Table S1).

During the research voyage, a field blank was collected by fitting the cascade impactor with a set of filters and walking the cascade impactor from the laboratory to the HV-AS in the same way that atmospheric samples were deployed. The cascade impactor was placed into the HV-AS and then immediately removed without the HV-AS turning on, after which the filters were removed from the cascade impactor and stored in the same manner as the atmospheric samples. All chemical analyses performed on samples were also performed on the field blank filters to assess possible contamination during filter deployment or sample handling.

171 **2.2**) Sample analysis

Filter stages 1 to 4 were extracted using ultra-clean deionised water (DI; 18.2 M Ω) under a laminar flow cabinet (Air Science). The extraction ratio was approximately 30 cm² of filter in 25 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 a clean 30 mL HDPE bottle and stored at -20°C until analysis (Baker et al., 2010).

176 Aerosol nitrate concentrations ([NO₃⁻]) were determined using a Thermo Scientific Dionex Aquion Ion Chromatography (IC) system (precision of ± 0.3 µmol L⁻¹). The anion IC contained an AG22 RFIC 4 x 50 mm guard column and AG22 RFIC 177 178 4×250 mm analytical column. A six-point standard curve that encompassed the range of sample concentrations (extract [NO₃-179]: 1.3 to 27.7 μ mol L⁻¹) was run on each day of analysis (Dionex Seven Anion-II Standard) and an R² value > 0.999 was 180 required for sample analysis to proceed. Final aerosol [NO₃⁻] were corrected by subtracting the field blanks, which represented 181 35% of the total $[NO_3^-]$ on average. Aerosol samples were also analysed for $[NO_3^-]$ using a Lachat OuikChem® flow injection 182 autoanalyzer (precision of $\pm 0.8 \text{ }\mu\text{mol} \text{ }L^{-1}$). The average [NO₃⁻] measured using the Lachat OuikChem® flow injection 183 autoanalyzer and the IC system is reported (Table S3).

184 Nitrogen and oxygen isotopic ratios were measured using the denitrifier method (Sigman et al., 2001 and Casciotti et al., 2002). To determine the ${}^{15}N/{}^{14}N$ and ${}^{18}O/{}^{16}O$ of NO₃⁻, a natural strain of denitrifying bacteria, *Pseudomonas aureofaciens*, that 185 lack the terminal nitrous oxide (N₂O) reductase enzyme were used to convert aqueous NO_3^- quantitatively to N₂O gas. The 186 187 product N₂O was analysed by continuous flow isotope ratio mass spectrometry (IRMS) using a Delta V Advantage IRMS 188 interfaced with an online N₂O extraction and purification system. Individual analyses were referenced to injections of N₂O 189 from a pure gas cylinder and then standardized through comparison to the international reference materials of IAEA-N3 and USGS34 for δ^{15} N-NO₃⁻, and IAEA-N3, USGS34 and USGS35 for δ^{18} O-NO₃⁻ (Table S2) (Böhlke et al., 2003). The 15 N/¹⁴N of 190 samples was corrected for the contribution of ¹⁷O to the peak at mass 45 using an average reported Δ^{17} O value of 26‰ from 191 192 atmospheric nitrate collected in the Weddell Sea (Morin et al., 2009). The pooled standard deviation for all measurements of 193 IAEA-N3 and USGS34 for δ^{15} N-NO₃⁻, and IAEA-N3, USGS34 and USGS35 for δ^{18} O-NO₃⁻ are reported (Table S2). All 194 samples were measured in triplicate in separate batch analyses. The pooled standard deviation from all replicate analyses of 195 samples was 0.25‰ for δ^{15} N-NO₃⁻ and 0.64‰ for δ^{18} O-NO₃⁻. The average δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻ computed for each filter 196 deployment was weighted by the [NO₃⁻] observed for each stage and error was propagated according to standard statistical 197 practises (Table S3).

Seawater samples were collected in triplicate every two hours from the ship's underway system (position at depth approximately 5 m) for the analysis of surface ocean nitrite concentrations ($[NO_2^{-}]$). $[NO_2^{-}]$ 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) (Table S4).

202 2.3) Air mass back trajectory analysis

To determine the air mass source region for each aerosol sample, air mass back trajectories (AMBTs) were computed for each hour in which the HV-AS was operational for at least 45 minutes of that hour. Given that the ship was moving, a different date, time and starting location was used to compute each AMBT. An altitude of 20 m was chosen to match the height of the HV-AS above sea level and 72-hour AMBTs were computed to account for the lifetime of NO₃⁻ in the atmosphere. All AMBTs were computed with NOAA's Hybrid Single-Particle Lagrangian Integrated Trajectory model (HYSPLIT v 4), using NCEP Global Data Assimilation System (GDAS) output, which can be accessed at https://www.arl.noaa.gov/ready/hysplit4.html (NOAA Air Resources Laboratory, Silver Spring, Maryland) (Stein et al., 2015; Rolph 2016).

210 3) Results

211 The coarse mode (> 1 μ m in diameter) aerosol [NO₃⁻] computed by summing the [NO₃⁻] of stages 1 through 4, ranged from 212 15.1 to 235.0 ng m⁻³ (Fig. 1a and Table 1). The mass-weighted δ^{15} N of coarse mode aerosol NO₃⁻ ranged from -43.1‰ to -213 2.7‰ (Figs. 1b, 2 and Table 1). There were no clear trends in atmospheric $[NO_3^-]$ or $\delta^{15}N$ -NO₃⁻ with aerosol size (Table S5). 214 The highest nitrate concentrations occurred between 34° S and 45° S, and then decreased with increasing latitude. Similarly, higher values characterized δ^{15} N-NO₃⁻ between 34° S and 45° S (-4.9 ± 1.3‰), and then decreased with increasing latitude 215 216 (Fig. S2). At the high latitudes (south of 60° S), median values of 26.21 ng m⁻³ and -22.2‰ were observed for nitrate 217 concentration and δ^{15} N, respectively. Coincident mass-weighted δ^{18} O-NO₃⁻ values ranged from 16.5% to 70% (Figs. 1c, 3 218 and Table 1). No latitudinal trend in δ^{18} O-NO₃⁻ was apparent, although distinctly low δ^{18} O-NO₃⁻ values were observed in the 219 Weddell Sea, as discussed in section 4.3 below. The difference between δ^{18} O-NO₃⁻ observed in the Weddell Sea (during January to February) and δ^{18} O-NO₃⁻ observed at corresponding latitudes (56° S to 70° S) during the early and late summer 220 221 transects is statistically significant (p-value = 0.009). The early and late summer cruise transects were similar spatially in that 222 both took place along the same hydrographic line (i.e., the Good Hope line), apart from the deviation to South Georgia during 223 late summer (Fig. 2a & b). Even though the early and late summer cruise transects occurred in December and March, respectively, there is no statistically significant difference in [NO₃⁻] (p-value = 0.43), δ^{15} N-NO₃⁻ (p-value = 0.53) or δ^{18} O-NO₃⁻

(p-value = 0.67) between them. Therefore, the early and late summer legs are discussed together and collectively referred to as the latitudinal transect.

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Table 1: The average (Avg), standard deviation (SD) and range of total coarse-mode (> 1 μ m) atmospheric nitrate concentration ([NO₃⁻]; ng m⁻³) and the mass weighted average N and O isotopic composition of coarse mode nitrate (δ^{15} N-NO₃⁻ and δ^{18} O-NO₃⁻; ‰) are shown. Cruise legs are denoted as follows: early summer (ES), Weddell Sea (WS) and late summer (LS).

Leg	$[NO_3^-]$ (ng m ⁻³)		δ^{15} N-NO ₃ ⁻ (‰ vs. N ₂)		δ^{18} O-NO ₃ ⁻ (‰ vs. VSMOW)	
	Avg (SD)	Range	Avg (SD)	Range	Avg (SD)	Range
ES	88.1 (70.2)	20.0 to 235.0	-19.5 (16.4)	-42.9 to -2.7	47.1 (17.8)	16.5 to 70.0
WS	29.4 (12.1)	15.1 to 59.5	-22.7 (7.2)	-38.1 to -11.6	38.4 (12.9)	18.8 to 60.3
LS	59.7 (59.4)	16.9 to 177.4	-15.0 (8.1)	-25.6 to -4.6	50.3 (6.3)	43.1 to 58.9



Figure 1. (a) The average (\pm 1 SD) coarse mode (> 1 µm) nitrate concentration [NO₃⁻] (ng m⁻³), and the weighted average (\pm 1 SD) δ^{15} N (b) and δ^{18} O (c) of atmospheric nitrate (δ^{15} N-NO₃⁻ (‰ vs. N₂) and δ^{18} O-NO₃⁻ (‰ vs. VSMOW), respectively), as a function of latitude (°S). Early and late summer latitudinal transects are denoted by the red triangles and green squares, respectively. Weddell Sea samples are denoted by blue circles. Where error bars (\pm 1 SD) are not visible, the standard deviation is smaller than the size of the marker.

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Figure 2. 72-hour AMBTs (grey lines) computed for each hour of the voyage when the HV-AS was operational for more than 45 minutes of the hour during early summer (a), late summer (b), and in the Weddell Sea (c). The colour bar represents the weighted average $\delta^{15}N$ of coarse mode (> 1 µm) atmospheric nitrate ($\delta^{15}N-NO_3^{-}$). Individual AMBTs for each aerosol sample from the Weddell Sea are shown in Fig. S1. The white represents the location of the sea ice determined using satellite derived sea-ice concentration data, obtained from passive microwave sensors AMSR2 (Advanced Microwave Scanning Radiometer 2, Spreen et al., 2008).

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Figure. 3. 72-hour AMBTs (grey lines) computed for each hour of the voyage when the HV-AS was operational for more than 45 minutes of the hour during early summer (a), late summer (b), and in the Weddell Sea (c). The colour bar represents the weighted average δ^{18} O of coarse mode (> 1 µm) atmospheric nitrate (δ^{18} O-NO₃⁻). Individual AMBTs for each aerosol sample from the Weddell Sea are shown in Fig. S1 The white represents the location of the sea ice (see Fig. 2 caption).

296 4) Discussion

297 Our observations reveal a latitudinal gradient in atmospheric NO_3^- concentration and $\delta^{15}N-NO_3^-$, which we hypothesize 298 may be attributed to the varying contribution of the dominant NO_x sources present between Cape Town and coastal Antarctica. 299 In contrast, $\delta^{18}O-NO_3^-$ depicts no latitudinal trend; however, relatively low $\delta^{18}O-NO_3^-$ values are observed in the Weddell Sea,

- 300 which we hypothesize may be attributed to the influence of sea ice emissions on NO_x cycling. Below, we first discuss the
- 301 extent to which anthropogenic NO_x sources may influence the observed atmospheric NO₃⁻ concentrations and δ^{15} N signatures.
- 302 Then we discuss the dominant NO_x sources to low, mid and high latitude Southern Ocean MBL NO_3^- , determined in part from
- 303 72-hour AMBTs, as well as the role of various oxidants in NO/NO₂ cycling and NO₂ oxidation.

304 4.1) Minimal influence of anthropogenic NO_x sources

305 Aerosol NO_3^{-1} concentrations were low (< 100 ng m⁻³; Fig. 1a) for most air masses sampled along the latitudinal transect 306 and in the Weddell Sea, consistent with the expectation of minimal influence from anthropogenic NO_x sources. For 307 comparison, $[NO_3^-]$ in a polluted urban airshed over South Africa can be > 500 ng m⁻³ (Collett et al., 2010). Interestingly, NO_3^- 308 concentrations were higher (± 200 ng m⁻³; Fig. 1a) in samples collected near the South African coast at the beginning of the 309 latitudinal transect (i.e., above 43° S). However, 72-hour AMBTs computed for all latitudinal transect samples indicate that 310 sampled air masses originated from over the South Atlantic sector of the Southern Ocean (Fig. 2a and 2b), with no continental 311 influence and limited opportunity for direct anthropogenic NO_x emissions to contribute to aerosol NO₃⁻, assuming NO₃⁻ has a 312 lifetime of 72 hours (Alexander et al., 2020). Furthermore, contamination from ship stack emissions was avoided by using a 313 sector collector to restrict HV-AS activity to certain wind directions (Sect. 2.1). As such, the higher atmospheric NO_3^{-1} 314 concentrations observed near South Africa are best explained by greater lightning NO_x production, which generally occurs 315 between 40° S and the ITCZ during summer (Nesbitt et al., 2000; van der A et al., 2008).

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317 4.2) Interpretation of natural NO_x sources using the N isotopic composition of atmospheric NO₃⁻

318 Aerosol δ^{15} N-NO₃⁻ ranged from -2.7‰ for low-latitude air masses to -42.9‰ for high-latitude air masses (including those 319 sampled in the Weddell Sea; Fig. 1b). As discussed in section 1.1, the δ^{15} N-NO₃⁻ reflects the δ^{15} N of the source NO_x plus any 320 isotopic fractionation imparted from NO/NO₂ cycling or NO_x to NO₃⁻ conversion. Similar to previous studies, we surmise that 321 NO_x equilibrium fractionation is unlikely to be relevant in our system, as NO_x concentrations are significantly lower than O_3 322 concentrations (Elliott et al., 2007; Morin et al., 2009; Walters et al., 2016; Park et al., 2018). Typical O₃ concentrations 323 observed at coastal sites in Antarctica are on the order of 20 ppbv (parts per billion by volume) (Nadzir et al., 2018), whereas 324 the sum of NO and NO₂ rarely exceeds 0.04 ppbv (Jones et al., 2000; Weller et al., 2002; Bauguitte et al., 2012). Under these 325 conditions NO_x isotopic exchange occurs at a much slower rate than (R1) and (R2), such that little to no equilibrium isotope 326 fractionation is expressed and the $\delta^{15}N$ of the NO₃⁻ should reflect the $\delta^{15}N$ of the NO_x source (Walters et al., 2016). 327 Additionally, equilibrium isotope effects are temperature dependent (increasing with decreasing temperature) and here ambient 328 temperatures decline with increasing latitude. Therefore, if equilibrium isotope fractionation were occurring during NO-NO₂ 329 cycling and/or NO_x to NO₃⁻ conversion, one would expect δ^{15} N-NO₃⁻ to increase with latitude, as both fractionation processes 330 produce NO₃⁻ with a δ^{15} N signature higher than the source NO_x. However, the opposite trend is observed here whereby δ^{15} N- NO₃⁻ decreases with increasing latitude (Fig. 1b). Therefore, we discount the hypothesis that equilibrium isotope effects can explain the latitudinal gradient in δ^{15} N-NO₃⁻.

333 NO_3 in the Antarctic troposphere may also derive from stratospheric denitrification, whereby HNO₃ is injected into the 334 troposphere from the stratosphere via the subsidence and penetration of polar stratospheric clouds (PSC). However, this 335 phenomenon typically occurs in winter when the tropospheric barrier is weak and the lower stratosphere is cold enough for 336 PSC formation (Savarino et al., 2007; Walters et al., 2019). Furthermore, δ^{15} N-NO₃⁻ originating from stratospheric inputs is 337 estimated to be 19‰ \pm 3‰ (Savarino et al., 2007), a value substantially greater than the atmospheric δ^{15} N-NO₃⁻ observed here 338 for high-latitude air masses; thus, we discount a direct influence from stratospheric NO_x. We propose that the observed 339 variation in atmospheric δ^{15} N-NO₃⁻ across the Southern Ocean is best explained by the changing contribution of three 340 dominant NO_x sources: lightning, surface ocean alkyl nitrate emissions, and photochemical production on snow and ice. 341 determined using AMBT analyses and typical NO_x source signatures where possible, as discussed below.

342 **4.2.1**) High-latitudes: Photochemical NO_x source

343 Aerosol δ^{15} N-NO₃ was relatively low in air masses from the southern high-latitudes, including in the Weddell Sea (average 344 of -24.3‰; Figs. 1b & 2). The latitudinal gradient in lightning NO_x production suggests that lightning NO_x is greatly reduced 345 at high latitudes(Nesbitt et al., 2000). Similar to other studies in the region (Savarino et al., 2007; Morin et al., 2009), we 346 suggest that photochemical NO_x production on snow or ice accounts for the low aerosol δ^{15} N-NO₃⁻ in high-latitude air masses, 347 where high-latitude air mass samples are defined as those exposed to the Antarctic continent or the surrounding sea ice (with 348 sea ice concentration being at least 50%). Antarctic estimates for isotopic fractionation associated with snow NO_3^- photolysis 349 during summer range from -47.9‰ to -55.8‰ for laboratory and field experiments, respectively (Berhanu et al., 2014, 2015), resulting in the emission of low δ^{15} N NO_x to the overlying atmosphere (Savarino et al., 2007; Morin et al., 2009; Shi et al., 350 2018; Walters et al., 2019). Therefore, NO₃⁻ photolysis explains the very low δ^{15} N-NO₃⁻ observed in high-latitude air masses 351 352 in early and late summer that crossed snow-covered continental ice or sea ice before being sampled (Figs. 2a & b). During 353 early summer, air masses spent significantly more time over the snow-covered continent compared to late summer and the sea 354 ice extent was greater in early summer compared to late summer (Figs. 2a & b). Combined, these dynamics resulted in a much 355 lower δ^{15} N-NO₃⁻ for high-latitude air masses during early summer compared to late summer (minimum value of -42.9% vs -25.6‰). Similarly low MBL δ^{15} N-NO₃⁻ values (< -30‰) were recently observed for the southern high latitudes of the Indian 356 357 ocean (Shi et al., 2021). Our data are also consistent with vear-round studies of atmospheric NO_3^- at coastal Antarctica 358 (Savarino et al., 2007) and the South Pole (Walters et al., 2019), where δ^{15} N-NO₃⁻ was reported to range from -46.9% to 10.8% 359 and from -60.8% to 10.5%, respectively. Both studies observed a seasonal cycle in δ^{15} N-NO₃⁻ whereby the lowest values 360 occurred during sunlit periods (i.e., summer) due to snowpack NO_x emissions and the highest values occurred during dark 361 periods (i.e., winter) due to stratospheric inputs (Savarino et al., 2007; Walters et al., 2019).

362 **4.2.2) Low- to Mid-latitudes: Oceanic NO_x source**

At the northern extent of our transects, the low-latitude aerosol samples, defined as those with air mass back trajectories originating from anywhere north of 43° S in early summer and 41° S in late summer (Fig. 2), had the highest average δ^{15} N-NO₃⁻ signature (-4.9 ± 1.3‰; n = 5). These values can be attributed to lightning-generated NO_x, which has a δ^{15} N signature close to 0‰ (Hoering 1957). Lightning activity at the low latitudes is also consistent with the higher atmospheric [NO₃⁻] observed (Fig. 1a) and is further supported by co-occurring high [NO₃⁻] and relatively high δ^{15} N-NO₃⁻ (Fig. S2). An average atmospheric δ^{15} N-NO₃⁻ signature of -4‰ was previously reported for the low latitude Atlantic Ocean, between 45° S and 45° N, and similarly attributed to a combination of natural NO_x sources including lightning (Morin et al., 2009).

370 Aerosol samples across the mid-latitudes had an average δ^{15} N-NO₃⁻ of -13.2‰ (Figs. 1b & 2). Mid-latitude air masses are 371 defined as those originating from anywhere south of 43° S in early summer and south of 41° S in late summer that made no 372 contact with Antarctica or any surrounding sea ice (Fig. 2a & b), therefore these samples were unlikely to be influenced by 373 snow emitted NO_x with its light isotopic signature. The beginning of the mid-latitude zone (i.e., 43° S and 41° S in early and 374 late summer, respectively) was defined by the presence of non-zero sea surface nitrite concentrations in early and late summer 375 (Fig. 4). However, the observed aerosol δ^{15} N-NO₃⁻ was too low (-14.5‰ to -11.2‰) to be explained solely by lightning 376 generated NO_x. In the absence of any signature of anthropogenic NO_x emissions (Sect. 4.1), we argue that the dominant NO_x 377 source for the mid-latitude samples originates from seawater.



Figure 4. 72-hour AMBTs computed for each hour of the voyage during early (a) and late (b) summer, when the HV-AS was operational for more than 45 minutes of the hour. AMBTs are colour coded by the weighted average $\delta^{15}N$ of atmospheric nitrate ($\delta^{15}N-NO_3^{-}$), represented by the horizontal colour bar. Over-layered are the surface ocean nitrite concentrations (circles; [NO₂⁻]; µmol L⁻¹), measured along each transect and represented by the vertical colour bar.

382

383 As mentioned in section 1, the most likely mechanism for an oceanic NO_x source is via the photolysis of surface ocean 384 derived RONO₂ in the MBL. NO derived from seawater nitrite is thought to limit RONO₂ production (Dahl and Saltzman 385 2008; Dahl et al., 2012), such that non-zero nitrite concentrations are required for RONO₂ production to occur. Here, surface ocean nitrite concentrations were relatively high, in particular from ~ 41° S to 50° S (Fig. 4). Furthermore, the latitudinal 386 387 extent of mid-latitude air masses with low δ^{15} N-NO₃⁻ signatures corresponds well with the same latitudinal extent in which 388 non-zero surface ocean nitrite concentrations occurred (Fig. 4). As such, we suggest that in this region oceanic RONO₂ emission is the main source to the Southern Ocean MBL, ultimately resulting in the low δ^{15} N-NO₃⁻ values observed for mid-389 390 latitude air masses.

391 No estimates exist for the δ^{15} N of oceanic RONO₂, however RONO₂ photolysis may yield isotopically light NO_x given that 392 NO₃⁻ photolysis produces low δ^{15} N products (e.g., Frey et al., 2009). Therefore, once oxidised in the overlying atmosphere, 393 NO_x derived from oceanic RONO₂ photolysis may form atmospheric NO₃⁻ with a low δ^{15} N signature. Aerosol δ^{15} N-NO₃⁻

394 values have been observed to range from -14.1% to -7.3% in the eastern equatorial Pacific (Kamezaki et al., 2019) and from 395 -6‰ to ~0‰ (average = -3.4‰) in the western equatorial Pacific (Shi et al., 2021). Observed δ^{15} N-NO₃⁻ is higher in the 396 western compared the eastern equatorial Pacific, which could be attributed to the proximity of the western equatorial Pacific 397 to continental/anthropogenic NO_x sources, resulting in NO₃⁻ having a higher δ^{15} N signature. The low average δ^{15} N-NO₃⁻ 398 observed for the mid-latitude air masses of the Southern Ocean MBL sampled in the present study (-14.5% to -11.2%). are 399 remarkably similar to those observed in the eastern equatorial Pacific (Kamazaki et al., 2019). Kamezaki et al., (2019) also concluded that such low δ^{15} N-NO₃⁻ values cannot be explained solely by lightning NO_x and given the lack of considerable 400 401 influence from any continental NO_x sources, they invoked the contribution of oceanic N emissions in the form of ammonia 402 (NH₃) and/or RONO₂. However, NH₃ flux data for the summertime Atlantic Southern Ocean derived from in situ 403 ocean/atmosphere observations suggest that the ocean in this region is a net sink of NH₃ (Altieri et al., 2021).

The latitudinal extent of our sampling campaign enabled us to estimate a range of likely values for the N isotopic composition of NO_3^- derived from oceanic RONO₂. We split the latitudinal transect into three regions, each characterised by the dominance of a different natural source of NO_3^- , i.e., lightning NO_x at the low-latitudes (Fig. 5 light orange), oceanic RONO₂ emissions at the mid-latitudes (Fig. 5 dark orange) and snowpack emissions at the high-latitudes (Fig. 5 red).



Figure 5. 72-hour AMBTs computed for each hour of the voyage during early (a) and late (b) summer, when the HV-AS was operational for more than 45 minutes of the hour. Light orange, dark orange and red AMBTs represent time spent over the low, mid- and high latitude SO, respectively. The white represents the location of the sea ice (see Fig. 2 caption).

411

412 Assuming that the dominant natural source of NO_3^- is the only source relevant in each latitudinal zone, we estimate the 413 contribution of each source to total NO_3^- formation by ascertaining the amount of time air masses spent in each zone. We 414 further assume that atmospheric $\delta^{15}N-NO_3^-$ reflects at most a combination of two sources based on the AMBTs of each sample, 415 either lightning NO_x and oceanic RONO₂ emissions near South Africa, or oceanic RONO₂ emissions and snowpack NO_x

emissions near Antarctica (Fig. 5 and Table S6). Using a two-end member mixing model the δ^{15} N signature of the source NO₃⁻ 416 417 derived from mid-latitude Southern Ocean RONO₂ emissions was calculated for all samples where air masses from the mid-418 latitude region contributed at least 10% (Table S6). This 10% threshold was chosen as the isotopic endmember of oceanic 419 RONO₂ is harder to determine with confidence when its contribution to total NO_3^- is less than 10%. As an example, the 420 AMBTs for sample ES 4 spent 3% of the time in the low-latitude zone and 97% in the mid-latitude zone. Using the measured 421 δ^{15} N-NO₃⁻ for ES 4 of -14.5‰ and assuming lightning NO_x has a δ^{15} N signature of 0‰, we calculate the δ^{15} N signature of the RONO₂-derived NO₃⁻ to be -14.9%. It is important to note that using this approach to estimate the δ^{15} N-NO₃⁻ from oceanic 422 423 RONO₂ emissions relies heavily on AMBTs generated using HYSPLIT. While HYSPLIT is a frequently used tool for assessing 424 air mass origin in the Southern Hemisphere and over Antarctica (Morin et al., 2008; Walters et al., 2019; Shi et al., 2021), it is 425 important to note that a spatial uncertainty of 15% to 30% of the trajectory path distance can be expected (Scarchilli et al., 426 2011). AMBTs also become increasingly uncertain the further back in time they are used (Sinclair et al., 2013). Some of this 427 uncertainty is alleviated by the fact that the AMBTs generated here are relatively short (< 5 days). Additionally, the spatial 428 scale of the low-, mid- and high-latitude zones is large, such that some variation in sample AMBTs will not significantly alter 429 the expected dominant NO₃⁻ source.

Using this approach for each filter deployment along the latitudinal transect, an average δ^{15} N-NO₃⁻ from oceanic RONO₂ emissions of -21.8 ± 7.6‰ was estimated. Furthermore, the contribution of RONO₂ emissions can explain the lowering of δ^{15} N from 0‰ for the low-latitude air mass samples. For example, the highest δ^{15} N observed in the study was -2.7‰, and this sample has a < 5% contribution from the mid-latitude zone. The other two low-latitude samples have 30% to 40% contribution from the mid-latitude zone and their δ^{15} N is lower (Table S3), as expected due to the influence of RONO₂ emissions.

The influence of low δ^{15} N-NO₃⁻ from RONO₂ emissions is not limited to the Southern Ocean, and this estimate of the N isotopic composition for the RONO₂ derived NO₃⁻ source may be useful to constrain the contribution of RONO₂ emissions to NO₃⁻ formation in other ocean regions with elevated surface ocean nitrite concentrations, such as the tropical Pacific.

438 **4.3**) The O isotopes of atmospheric nitrate

439 The corresponding δ^{18} O values allow us to determine the pathways of NO₃⁻ formation from NO_x. However, an assumption

must first be made regarding the oxidation of NO to NO₂. While the dominant oxidant of NO to NO₂ is O₃ (R1) in most of the

- 441 troposphere, over the open ocean there can be a significant contribution via the reaction of NO with peroxy radicals (HO₂ and
- 442 its organic homologues RO_2) (Alexander et al., 2020). Peroxy radicals compete with O_3 to convert NO into NO_2 via R10:

443 NO + HO₂ (or RO₂)
$$\rightarrow$$
 NO₂ + OH (or RO)

440

- 444 The δ^{18} O of peroxy radicals is much lower than that of O₃ because the O atoms derive from atmospheric O₂, which has a well-
- 445 defined δ^{18} O of 23.9‰ (Kroopnick and Craig, 1972). The δ^{18} O-NO₂ can then be calculated using Eq. (2),
- 446 δ^{18} O-NO₂ = (δ^{18} O-O₂)(1-f) + (δ^{18} O-O₃*)(f)

(2)

(R10)

where f is the fraction of NO₂ formed from R1, (1-f) is the fraction formed from R10, and the terminal δ^{18} O-O₃ value (δ^{18} O-O₃*) is 130.4 ± 12.9‰ (Vicars and Savarino, 2014).

- 449 The δ^{18} O-NO₃⁻ is then determined using Eq. (3) in which two thirds of the O atoms in NO₃⁻ come from NO₂ and one third
- 450 comes from OH i.e., R3, or using Eq. (4) in which three sixths of the O atoms in NO_3^- come from O_3 , two sixths come from 451 NO₂ and one sixth comes from H₂O i.e., R4-R6 (Hastings et al., 2003; Alexander et al., 2020).
- 452 δ^{18} O-NO₃-(R3) = (2/3)(δ^{18} O-NO₂) + (1/3)(δ^{18} O-OH)

(3)

(4)

- 453 δ^{18} O-NO₃ (R4-R6) = (1/2)(δ^{18} O-O₃*) + (1/3)(δ^{18} O-NO₂) + (1/6)(δ^{18} O-H₂O)
- 454 We assume that 15% of NO to NO₂ conversion occurs via HO₂/RO₂ oxidation and 85% by O₃ oxidation as is suggested by 455 global models (Alexander et al., 2020), and use the minimum and maximum δ^{18} O-H₂O range of -27.5% to 0%, the 456 temperature-dependent equilibrium isotope exchange between OH and H_2O (Walters and Michalski, 2016), and the resulting 457 minimum and maximum estimates for δ^{18} O-OH of -67.4‰ to -41.0‰. Using these assumptions and Eq. (3) and (4), the 458 expected δ^{18} O-NO₃⁻ for the daytime OH oxidation pathway (R3) is 46.5% to 71.4%, and for the dark reactions (R4-R6) is 459 88.7% to 113.5%. The observed δ^{18} O-NO₃ values were all less than 70% (Figs. 1c and 3), suggesting that NO_x oxidation by 460 OH (R3) was indeed the dominant pathway for atmospheric NO₃⁻ formation during summer. The low δ^{18} O-NO₃⁻ values 461 observed suggest a minimal influence of O_3 in the oxidation chemistry, ruling out both the halogen (R8 to R9) and DMS (R7) 462 related NO_3^- formation pathways in addition to N_2O_5 hydrolysis (R4-6). This is consistent with previous year-round studies of 463 atmospheric NO₃⁻ at coastal Antarctica (Savarino et al., 2007) and the South Pole (Walters et al., 2019) where δ^{18} O-NO₃⁻ was 464 at a minimum in summer (59.6‰ and 47.0‰, respectively). Both studies confirm the importance of HO_x oxidation chemistry 465 in summer when solar radiation enhances the production of these oxidants, followed by a switch to O_3 dominated oxidation 466 chemistry in winter (Savarino et al., 2007; Ishino et al., 2017; Walters et al., 2019).
- 467 Interestingly, most aerosol samples have a δ^{18} O-NO₃⁻ less than 46.5‰ (n=19), the lower limit estimated above for the OH 468 pathway. This suggests that there is more NO to NO₂ conversion via HO_2/RO_2 oxidation occurring than the global average. A 469 maximum HO₂/RO₂ contribution to NO oxidation of ~63% is required to explain the lowest δ^{18} O-NO₃⁻ value, which was 470 observed over the mid-latitudes during early summer. Increased RO₂ production over the mid-latitudes could derive from 471 RONO₂ photolysis in the MBL, which we hypothesise is happening in this region based on the δ^{15} N-NO₃⁻ (Sect. 4.2.2). 472 Although the lowest δ^{18} O observation occurred in the mid-latitudes, the majority of low δ^{18} O-NO₃⁻ values were observed in 473 the Weddell Sea, away from the region of maximum RONO₂ emissions. Approximately half of the Weddell Sea samples have 474 a δ^{18} O-NO₃⁻ < 31‰, which would require a HO₂/RO₂ contribution to NO oxidation upwards of 40% (more than double the 475 contribution estimated by global models (Alexander et al., 2020)). These δ^{18} O-NO₃⁻ observations are unusually low compared 476 to previous observations for the same region in spring (Morin et al., 2009). We hypothesize that the large contribution of 477 HO₂/RO₂ to NO/NO₂ oxidation (i.e., a decrease in f in Eq. (2)) resulting in these low δ^{18} O-NO₃ values is due to the influence 478 of sea ice emissions. The 72-hour AMBTs for these low δ^{18} O-NO₃⁻ Weddell Sea samples indicate that all the air masses either 479 originated from, or spent a significant amount of time recirculating, over the sea ice covered region of the western Weddell 480 Sea (Fig. 6b). By contrast, aerosol samples from the Weddell Sea with δ^{18} O-NO₃ values greater than 31% have air masses 481 that experienced significantly more oceanic influence (Fig. 6a). There is evidence that sea ice can lead to enhanced peroxy 482 radical production (Brough et al., 2019). In that work, increased HO₂ + RO₂ concentrations were observed during spring at a

483 coastal Antarctic site when air masses arrived from across a sea ice covered zone. This was attributed to the oxidation of 484 hydrocarbons by chlorine atoms, which leads to increased RO₂ concentrations via R11 and R12:





Figure. 6. 72-hour AMBTs (light blue lines) computed for each hour of the voyage in the Weddell Sea, when the HV-AS was operational for more than 45 minutes of the hour. The vertical colour bar represents the weighted average δ^{18} O of atmospheric nitrate (δ^{18} O-NO₃⁻), where δ^{18} O-NO₃⁻ was > 31‰ (a) and < 31‰ (b). The white represents the location of the sea ice (see Fig. 2 caption).

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514 Cl atoms are much more reactive with hydrocarbons than OH (Monks, 2005) and can enhance hydrocarbon oxidation even 515 when present at low concentrations. Brough et al. (2019) suggest that air masses that traversed the sea ice zone contained 516 photolabile chlorine compounds that built up at night until photolysis occurred during the next day (Brough et al., 2019). Although our study was conducted in summer (the season of minimum sea ice extent), the sampling locations were uniquely positioned at the western edge of the Weddell Sea gyre where significant sea ice remained (Fig. 6). Therefore, we suggest that chlorine chemistry over the sea ice increased RO₂ concentrations at the time of our sampling, allowing the NO + RO₂ pathway to play a more significant role in the Weddell Sea and resulting in low δ^{18} O-NO₃⁻ values. We note that the only other estimates of δ^{18} O-NO₃⁻ from the Weddell Sea ranged from ~ 50‰ to 110‰ during springtime, and these samples were associated with air masses that spent almost no time over the sea ice and therefore had limited potential for this peroxy radical chemistry to drive down the δ^{18} O-NO₃⁻ to the low values we observe (Morin et al., 2009).

524 **5**) Conclusions

525 Our observations across a large latitudinal gradient of the summertime Southern Ocean MBL suggest it is dominated by natural 526 NO_x sources with distinct isotopic signatures. Aerosol NO_3^- was predominantly formed from lightning generated NO_x with a 527 $\delta^{15}N$ of ~ 0% at the lower latitudes, whereas snowpack NO_x emissions with a $\delta^{15}N \sim -48\%$ dominated the MBL inventory at 528 higher latitudes. Over the mid-latitudes, NO₃⁻ derived primarily from oceanic RONO₂ emissions, with an estimated δ^{15} N 529 signature of $\sim -22.0\%$. Additional research is needed to improve our mechanistic and isotopic understanding of surface ocean 530 RONO₂ formation, flux, and conversion to aerosol nitrate in order to constrain the contribution of oceanic RONO₂ emissions 531 to NO₃⁻ formation in other ocean regions where this source has been invoked, such as the tropical Pacific (Kamezaki et al., 532 2019). The isotopic composition of NO_3^- observed here can further inform interpretations of Antarctic ice core NO_3^- isotope 533 records to understand aerosol climate forcing and controls on the atmospheric oxidation budget over millennia (Freyer et al., 534 1996; Jiang et al., 2019) – the interpretation of which relies on knowledge of the NO_x isotopic source signatures in the polar 535 atmosphere.

536 The δ^{18} O-NO₃⁻ values were consistently lower than 70‰, which confirms NO_x oxidation by OH (R3) to be the dominant pathway for atmospheric NO₃⁻ formation during summer. However, unusually low δ^{18} O-NO₃⁻ values observed at the mid-537 538 latitudes and in the Weddell Sea indicate the increased importance of peroxy radicals (and decreased importance of O_3) in NO 539 oxidation to NO_2 in the MBL At the mid latitudes peroxy radicals (RO_2) may derive from RONO₂ photolysis, while in the 540 Weddell Sea, sea ice appears to play an important role in the formation of this oxidant via its influence on chlorine chemistry 541 (Brough et al., 2019). This implies that snow covered sea ice is not only a source of NO_x but also other species that have the 542 potential to change the composition of the atmosphere above the ice and impact NO_x oxidation chemistry. These results also 543 highlight the utility of δ^{18} O-NO₃⁻ to identify the major oxidants in NO oxidation, as well as NO_x to NO₃⁻ conversion. In 544 particular, δ^{18} O-NO₃⁻ can serve as a useful tool for testing our understanding of the relative importance of HO₂/RO₂ in NO/NO₂ 545 cycling, which can be difficult to constrain in some environments.

546 Our study challenges the traditional paradigm that considers the ocean as a passive recipient of N deposition, as the 547 Southern Ocean mid-latitude NO₃⁻ source may derive almost entirely from oceanic RONO₂ emissions. In the tropical equatorial 548 Pacific atmosphere, Kamezaki et al. (2019) also suggested evidence for a low δ^{15} N-NO₃⁻ source derived from the ocean. In the

- subtropical Atlantic Ocean MBL, Altieri et al. (2016) found that biogeochemical cycling in the surface ocean can directly influence the lower atmosphere serving as a source of aerosol organic N and ammonium. This study suggests that the surface waters of the Southern Ocean may also serve as a NO_x source, ultimately resulting in NO_3^- aerosol formation. As such, the surface ocean may play a bigger role in atmospheric oxidative capacity over remote marine regions than previously thought.
- 553

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- 558
- 559 *Competing interests.* The authors declare that they have no conflict of interest.
- 560

561 *Data availability statement*. Data sets for this research are available at https://doi.org/10.5281/zenodo.5740618.

562

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