Biogeochemical controls on ammonium accumulation in the surface layer of the Southern Ocean

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1. Abstract

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18 The production and removal of ammonium (NH4+) are essential upper-ocean nitrogen cycle 19 pathways, yet in the Southern Ocean where NH₄⁺ has been observed to accumulate in surface 20 waters, its mixed-layer cycling remains poorly understood. For surface seawater samples 21 collected between Cape Town and the marginal ice zone in winter 2017, we found that NH4+ 22 concentrations were five-fold higher than is typical for summer, and lower north than south of 23 the Subantarctic Front (0.01-0.26 µM versus 0.19-0.70 µM). Our observations confirm that 24 NH₄⁺ accumulates in the Southern Ocean's winter mixed layer, particularly in polar waters. NH₄⁺ 25 assimilation rates were highest near the Polar Front (12.9 ± 0.4 nM day⁻¹) and in the Subantarctic 26 Zone $(10.0 \pm 1.5 \text{ nM day}^{-1})$, decreasing towards the marginal ice zone $(3.0 \pm 0.8 \text{ nM day}^{-1})$ despite 27 the high ambient NH₄⁺ concentrations in these southernmost waters, likely due to the low 28 temperatures and limited light availability. By contrast, rates of NH₄+ oxidation were higher south 29 than north of the Polar Front (16.0 ± 0.8 versus 11.1 ± 0.5 nM day⁻¹), perhaps due to the lower 30 light and higher iron conditions characteristic of polar waters. NH₄⁺ concentrations were also 31 measured on five transects of the Southern Ocean (Subtropical- to marginal ice zone) spanning 32 the 2018/2019 annual cycle. These measurements reveal that mixed-layer NH₄+ accumulation 33 south of the Subantarctic Front derives from sustained heterotrophic NH₄⁺ production in late 34 summer through winter that in net, outpaces NH₄⁺ removal by temperature-, light-, and iron-35 limited microorganisms. Our observations thus imply that the Southern Ocean becomes a 36 biological source of CO2 to the atmosphere in autumn and winter not only because nitrate

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2. Introduction

accumulation.

40 The Southern Ocean impacts the Earth system through its role in global thermohaline circulation,

drawdown is weak, but also because the ambient conditions favour net heterotrophy and NH4+

41 which drives the exchange of heat and nutrients among ocean basins (Frölicher et al., 2015;

- 43 Sarmiento et al., 2004). The Southern Ocean also plays an integral role in mediating climate, by
- 44 transferring carbon to the deep ocean via its biological and solubility pumps (Sarmiento & Orr,
- 45 1991; Volk & Hoffert, 1985) and through the release of deep-ocean CO₂ to the atmosphere during
- 46 deep-water ventilation (i.e., CO₂ leak; Broecker & Peng, 1992; Lauderdale et al., 2013; Sarmiento
- 47 & Toggweiler, 1984). Upper Southern Ocean circulation is dominated by the eastward-flowing
- 48 Antarctic Circumpolar Current (ACC) that consists of a series of broad circumpolar bands
- 49 ("zones") separated by oceanic fronts. These fronts can drive water mass formation (Ito et al.,
- 50 2010) and nutrient upwelling that supports elevated productivity (Sokolov & Rintoul, 2007).
- 51 Concentrations of the essential macronutrients, nitrate (NO₃) and phosphate (PO₄³-), are
- 52 perennially high in Southern Ocean surface waters, in contrast to most of the global ocean.
- 53 Assimilation of these nutrients, and thus primary productivity, is limited in the Southern Ocean
- 54 by numerous overlapping factors, including temperature, light, micronutrient concentrations, and
- 55 grazing pressure (e.g., Boyd et al., 2001; Martin et al., 1990; Reay et al., 2001; Smith Jr &
- 56 Lancelot, 2004). The strength of these limitations varies with sector (i.e., longitude), zone (i.e.,
- 57 latitude), and season, resulting in spatial and temporal variability in chlorophyll-a, primary
- 58 production, plankton community composition, and nutrient uptake regime (Mdutyana et al.,
- 59 2020; Mengesha et al., 1998; Shadwick et al., 2015; Thomalla et al., 2011). In addition to the
- 60 seasonality of temperature and light, Southern Ocean ecosystems are influenced by seasonal
- changes in nutrient availability. In winter, deep mixing replenishes the nutrients required for 61
- 62 phytoplankton growth but the low temperatures and light levels impede biological activity
- 63 (Rintoul & Trull, 2001). Once the mixed layer shoals in spring and summer, phytoplankton
- 64 consume the available nutrients until some form of limitation (usually iron; Nelson et al., 2001;
- 65 Nicholson et al., 2019) sets in. This balance between wintertime nutrient recharge and
- 66 summertime nutrient drawdown is central to the Southern Ocean's role in setting atmospheric
- 67 CO₂ (Sarmiento & Toggweiler, 1984).
- 68 The onset of iron limitation following the spring/early summer bloom in the Southern Ocean
- 69 drives phytoplankton to increased reliance on recycled ammonium (NH₄⁺; Timmermans et al.,
- 70 1998), the assimilation of which has a far lower iron requirement than that of NO₃ (Price et al.,
- 71 1994). The extent to which phytoplankton rely on NO₃ versus NH₄ as their primary N source
- 72 has implications for Southern Ocean CO2 removal since phytoplankton growth fuelled by
- 73 subsurface NO₃ ("new production") must be balanced on an annual basis by the export of sinking
- 74 organic matter ("export production"; Dugdale & Goering, 1967), which drives CO2 sequestration
- 75 (i.e., the biological pump; Volk & Hoffert, 1985). By contrast, phytoplankton growth on NH₄+ or
- other recycled N forms ("regenerated production") yields no net removal of CO2 to the deep 76
- 77 ocean (Dugdale & Goering, 1967). Considerable research has focused on NO₃ cycling in the
- 78 Southern Ocean mixed layer because of the importance of this nutrient for the biological pump
- 79 (e.g., Francois et al., 1992; Johnson et al., 2017; Mdutyana et al., 2020; Primeau et al., 2013;
- 80 Sarmiento & Toggweiler, 1984) and global ocean fertility (Fripiat et al., 2021; Sarmiento et al.,
- 81 2004). By contrast, the cycling of regenerated N within the seasonally-varying mixed layer -
- 82 including the production of NH₄⁺ and its removal by phytoplankton and nitrifiers – remains
- 83 poorly understood.
- 84 NH₄⁺ is produced in the euphotic zone as a by-product of heterotrophic metabolism (Herbert,
- 85 1999) and as a consequence of zooplankton grazing (Lehette et al., 2012; Steinberg & Saba,

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87 2008), and is removed by phytoplankton uptake (in euphotic waters) and nitrification (mainly in 88 aphotic waters). Heterotrophic bacteria can also consume NH₄⁺ (Kirchman, 1994) and have been 89 hypothesized to do so at significant rates in the Southern Ocean mixed layer in winter (Cochlan, 90 2008; Mdutyana et al., 2020). The assimilation of NH₄⁺ by phytoplankton requires relatively little 91 energy (Dortch, 1990) such that NH₄⁺ is usually consumed in the euphotic zone as rapidly as it 92 is produced (Glibert, 1982; La Roche, 1983), resulting in very low surface NH₄⁺ concentrations 93 in the open ocean (<0.2 μM; Paulot et al., 2015). Additionally, NH₄⁺ is often the preferred N 94 source to small phytoplankton (Dortch 1990), which typically dominate when iron and/or light 95 are limiting (Deppeler & Davidson, 2017; Pearce et al., 2010; Tagliabue et al., 2014) since their 96 higher cell surface area-to-volume ratio renders them less vulnerable to diffusion- and/or light limitation (Finkel et al., 2004; Fujiki & Taguchi, 2002; Hudson & Morel, 1993; Mei et al., 2009). 97

98 In addition to the implications for size distribution, the dominant N source to phytoplankton is 99 indicative of their potential for CO₂ removal, as per the new production paradigm (Dugdale & Goering, 1967). The N isotopic composition (δ^{15} N, in % vs. N₂ in air, = (15 N/ 14 N_{sample}/ 15 N/ 14 N_{air} 100 101 -1) x 1000) of particulate organic N (PON; a proxy for phytoplankton biomass) can be used to 102 infer the dominant N source to phytoplankton (Altabet, 1988; Fawcett et al., 2011; 2014; Lourey 103 et al., 2003; Van Oostende et al., 2017) since the assimilation of subsurface NO₃ yields PON 104 that is higher in δ^{15} N than that fuelled by recycled NH₄⁺ uptake (Treibergs et al., 2014). As such, measurements of bulk δ^{15} N-PON can be used to infer the net N uptake regime. 105

106 Nitrification, the oxidation of NH₄⁺ to nitrite (NO₂⁻) and then NO₃⁻ by chemoautotrophic bacteria 107 and archaea, was historically considered unimportant in euphotic zone waters due to the evidence for light inhibition of nitrifiers (Hooper & Terry, 1974; Horrigan & Springer, 1990; Olson, 1981) 108 109 and the fact that they are outcompeted by phytoplankton for NH₄⁺ (Smith et al., 2014; Ward, 110 1985; 2005; Zakem et al., 2018). However, this view has been challenged in numerous ocean 111 regions (Yool et al., 2007), including the Southern Ocean (Smart et al., 2015; Cavagna et al., 112 2015; Fripiat et al., 2015; Mdutyana et al., 2020). Wintertime upper-ocean NH₄⁺ dynamics thus 113 have implications for annual estimates of carbon export potential, insofar as NO₃ produced by 114 nitrification in the winter mixed layer that is subsequently supplied to spring and summer

phytoplankton communities constitutes a regenerated rather than a new N source on an annualbasis (Mdutyana et al., 2020).

117 Surface concentrations of NH₄⁺ are typically near-zero in spring and early- to mid-summer in the 118 open Southern Ocean (Daly et al., 2001; Henley et al., 2020; Sambrotto & Mace, 2000; Savoye 119 et al., 2004) due to assimilation by phytoplankton. In late summer, a peak in NH₄+ concentration 120 has been observed and attributed to enhanced bacterial and zooplankton activity following 121 elevated phytoplankton growth (Becquevort et al., 2000; Dennett et al., 2001; Mengesha et al., 122 1998). The limited available observations suggest that wintertime surface NH₄⁺ concentrations 123 are high (often >1 μM), particularly south of the Subantarctic Front (SAF) (Bianchi et al., 1997; 124 Henley et al., 2020; Philibert et al., 2015; Mdutyana et al., 2020; Weir et al., 2020). It thus appears 125 that NH₄⁺ is not depleted following the late summer peak in its concentration, which indicates 126 enhanced NH₄⁺ regeneration, either coincident with (but in excess of) NH₄⁺ assimilation in winter 127 and/or prior to this in late summer and/or autumn. Under these conditions, the Southern Ocean 128 mixed layer may become net heterotrophic and thus a biological source of CO2 to the 129 atmosphere.

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133 Here, we focus on NH₄⁺ cycling in the Southern Ocean mixed layer, mainly in winter, which is a

- 134 season assumed to be largely biologically dormant (Arrigo et al., 2008; Schaafsma et al., 2018)
- 135 and for which NH₄⁺ cycle data are scarce. We confirm that NH₄⁺ accumulates throughout the
- 136 winter mixed layer south of the SAF, and examine the potential drivers thereof. Using NH₄⁺
- 137 concentration data collected over a full annual cycle, we propose that these drivers include a
- 138 contribution from the residual late-summer NH₄+ pool, sustained NH₄+ production in the autumn
- 139 and winter, and limited wintertime NH4+ uptake and oxidation that nonetheless exceed the rate of
- in situ NH4+ production. Finally, from our temporally-resolved NH4+ concentration data, we 140
- propose for the first time a measurement-based seasonal cycle for the mixed-layer NH₄⁺ pool 141
- 142 south of the SAF.

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3. Methods

- 3.1 Cruise tracks and sample collection
- 145 Samples were collected for a series of analyses on the southward (S) and northward (N) legs of
- 146 a winter cruise between Cape Town, South Africa, and the marginal ice zone (MIZ) onboard the
- 147 R/V SA Agulhas II (VOY 025; 28 June to 13 July 2017) (Fig. 1). Samples were also collected for
- 148 NH₄⁺ concentration analysis on three cruises onboard the R/V SA Agulhas II during 2018/19:
- 149 early- and late summer samples were collected during the SANAE 58 Relief Voyage (6
- 150 December 2018 to 15 March 2019; VOY035); winter samples were collected during the SCALE
- 151 2019 (www.scale.org.za) winter cruise to the MIZ (18 July to 12 August 2019; VOY039); and
- 152 spring samples were collected during the SCALE 2019 spring cruise to the MIZ (12 October to
- 153 20 November 2019; VOY040) (Fig. S1).
- 154 Leg S of VOY025 in winter 2017 crossed the Atlantic sector and due to logistical constraints,
- 155 involved only surface underway collections, while leg N bordered the Atlantic and Indian sectors
- 156 (30°E; WOCE IO6 line) and included eight conductivity-temperature-depth (CTD) hydrocast
- 157 stations. Frontal positions were determined using the ship's hull-mounted thermosalinograph,
- 158 supported by temperature, salinity, and oxygen concentration data from CTD measurements
- 159 made during leg N. The salinity and oxygen sensors were calibrated against seawater samples
- 160 that were analyzed for salinity using a Portasal 8410A salinometer and for dissolved oxygen by
- 161 Winkler titration (Strickland & Parsons, 1972). Frontal positions were determined from sharp
- 162 gradients in potential temperature, salinity, potential density, and oxygen concentrations (Belkin
- & Gordon, 1996; Lutjeharms & Valentine, 1984; Orsi et al., 1995). For leg N, the mixed layer 163
- depth (MLD) was determined for each Niskin (up)cast as the depth between 10 m and 400 m at 164
- which the Brunt Väisälä Frequency squared, N^2 , reached a maximum (Carvalho et al., 2017). 165
- During leg S, samples were collected every four hours from the ship's underway system (~7 m 166
- 167 intake; "underway stations") while samples on leg N were collected from surface Niskin bottles
- 168 (~10 m, approximately 55% light depth) mounted on the CTD rosette ("CTD stations"). NH₄⁺
- 169 samples were also taken at 13 depths over the upper 500 m at the CTD stations. During the
- 170 2018/19 cruises, NH₄ samples were collected every two hours from the ship's underway system.
- 171 At all stations, 40 mL of unfiltered seawater was collected for the analysis of NH₄⁺ concentrations
- 172 in duplicate 50 mL high density polyethylene (HDPE) bottles that had been stored ("aged") with
- 173 orthophthaldialdehyde (OPA) working reagent. Unfiltered seawater was collected in duplicate

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- 175 50 mL polypropylene centrifuge tubes for the analysis of NO₃-, NO₂-, and PO₄³-, and in a single
- tube for urea. Immediately following collection, NH₄⁺ and nutrient samples were frozen at -20°C. 176
- 177 Duplicate size-fractionated chlorophyll-a samples were collected by filtering seawater (500 mL)
- 178 through 25 mm-diameter glass fibre filters (0.3 µm and 2.7 µm; Sterlitech GF-75 and Grade D,
- 179 respectively). Acetone (90%) was added to foil-wrapped borosilicate tubes containing the filters
- 180 and incubated at -20 °C for 24 hours. Duplicate seawater samples (4 L) were also gently vacuum-
- 181 filtered through combusted 47 mm-diameter, 0.3 µm GF-75 and 2.7 µm Grade-D filters for POC
- 182 and PON concentrations and δ^{15} N-PON. Filters were stored in combusted foil envelopes at -
- 183 80°C.
- 184 For microscopy, unfiltered seawater samples (250 mL) were collected during leg S in amber
- glass bottles and immediately fixed by the addition of 2.5 mL of Lugol's iodine solution (2% 185
- 186 final concentration), then stored at low room temperature in the dark until analysis. For flow
- 187 cytometry, seawater samples were collected in triplicate 2 mL microcentrifuge tubes, fixed with
- 188 glutaraldehyde (1% final concentration), and stored at -80°C until analysis (Marie et al., 2005).
- 189 Ten incubation experiments were conducted during leg S to measure net primary production
- 190 (NPP). In addition, four NPP experiments were conducted during leg N using seawater collected
- 191 from Niskin bottles fired at ~10 m. In all cases, pre-screened (200-µm mesh; to remove large
- 192 grazers) seawater was collected in three 2-L polycarbonate bottles to which NaH13CO3 was added
- 193 at ~5% of the estimated ambient DIC concentration. ¹³C enrichment was re-calculated post-cruise
- 194 using measured DIC concentrations, and these enrichments were used in all NPP rate
- 195 calculations. Bottles were incubated for 5 to 6.5 hours in custom-built deck-board incubators
- 196 shaded with neutral-density screens to mimic the 55% light level and supplied with running
- 197 surface seawater. Following incubation, each sample was divided (1 L per size fraction) and
- 198 gently vacuum filtered through combusted 0.3 µm and 2.7 µm glass fibre filters that were stored
- 199 in combusted foil at -80°C until analysis.
- 200 N uptake (as NO₃, NH₄⁺ and urea) and NH₄⁺ oxidation experiments were conducted at five
- 201 stations during leg S, with NH₄+ oxidation measured at two additional stations at the ice edge
- 202 (Fig. 1). On leg N, experiments were also conducted using seawater collected from ~10 m at the
- 203 same four CTD stations as the NPP experiments. Duplicate 1 L polycarbonate bottles were
- 204 amended with ¹⁵N-labeled NO₃, NH₄ or urea at ~10% of the ambient N concentration, estimated
- 205 based on past wintertime measurements (Mdutyana et al., 2020) and, in the case of NH₄+,
- coincident shipboard analyses. ¹⁵N enrichment was re-calculated post-cruise using the measured 206
- 207 nutrient concentrations, and these enrichments were used in all rate calculations. Incubations and 208
- filtration were carried out as for NPP, although 500 mL was used per size fraction. For NH₄+
- 209 oxidation, duplicate black 250 mL HDPE bottles were amended with $0.1~\mu M$ $^{15}NH_4^+$ and $0.1~\mu M$
- ¹⁴NO₂ (the latter as a "trap" for the ¹⁵NO₂ produced by NH₄ oxidation; Ward 2011). NH₄ * 210
- oxidation bottles were incubated for 24 hours under the same temperature conditions as the N 211
- 212 uptake and NPP experiments. Subsamples (50 mL) were collected from each bottle immediately
- 213 following tracer addition (T₀) and at the end of the experiments (T_f), and frozen at -20°C until
- 214 analysis.
- 215 3.2 Sample processing

3.2.1. Ammonium concentrations

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- 217 On all cruises, NH₄⁺ concentrations were measured shipboard using the fluorometric method of
- 218 Holmes et al. (1999) and a Turner Designs Trilogy fluorometer 7500-000 equipped with a UV
- 219 module. The detection limit, calculated as twice the pooled standard deviation of all standards,
- 220 was 0.06 µM. To prevent possible in/efflux of ammonia (NH₃) due to the temperature difference
- 221 between surface waters and the shipboard laboratory, samples were frozen immediately upon
- 222 collection, for a maximum of 24 hours. OPA working reagent was added to the frozen samples
- 223
- prior to defrosting them for analysis. Samples were slowly warmed to room temperature in a
- 224 water bath after OPA addition, incubated in the dark for four hours once defrosted, and then each
- 225 replicate was measured in triplicate. Standards and blanks were made daily using Type-1 Milli-
- 226 Q water. Precision was \pm 0.03 μM for replicate samples and standards.
- 227 During VOY040 (spring 2019), we investigated the possibility that the ship's underway system
- alters the seawater NH₄+ concentrations (e.g., due to contamination or cell breakage). We 228
- 229 collected surface samples from the underway and Niskin bottles concurrently and measured an
- 230 average NH4+ concentration difference of 0.07 \pm 0.15 μM (n=17), with no noticeable trend of
- 231 one method consistently yielding higher/lower concentrations. We thus have no reason to doubt
- 232 NH₄⁺ concentrations measured for seawater samples collected from the ship's underway system.

233 3.2.2. Macronutrient concentrations

- 234 Following the winter 2017 cruise, duplicate seawater samples were analysed manually for NO₂
- 235 and PO₄³⁻ (Bendschneider & Robinson, 1952; Murphy & Riley, 1962) using a Thermo Scientific
- 236 Genesys 30 Visible spectrophotometer. Precision and detection limit was $\pm~0.05~\mu M$ and 0.05
- 237 μM for NO_2^- and \pm 0.06 μM and 0.05 μM for PO_4^{3-} . The concentrations of $NO_3^- + NO_2^-$ and
- 238 Si(OH)₄ were measured using a Lachat QuickChem 8500 Series 2 flow injection autoanalyzer.
- 239 Aliquots of a certified reference material (JAMSTEC) were measured during each run to ensure
- 240 measurement accuracy (SD \leq 2%). The precision of the NO₃ + NO₂ and Si(OH)₄ measurements
- 241 was $\pm 0.4 \mu M$ and $\pm 0.2 \mu M$, respectively, and the detection limit was $0.1 \mu M$ and $0.2 \mu M$. NO_3
- 242 concentrations were calculated by subtraction (i.e., NO₃ + NO₂ - NO₂), with error propagated
- 243 according to standard statistical practices. Urea-N (hereafter, urea) concentrations were
- 244 determined via the room-temperature, single-reagent colorimetric method (Revilla et al., 2005)
- 245 using a Thermo Scientific Genesys 30 Visible spectrophotometer; precision was \pm 0.04 μ M and
- 246 the detection limit was $0.04~\mu M$.

247 3.2.3. Chlorophyll-a concentrations

- 248 Chlorophyll-a concentrations ([chl-a]) were determined shipboard using the nonacidified
- 249 fluorometric method (Welschmeyer, 1994). The Turner Designs Trilogy fluorometer was
- 250 calibrated with an analytical standard (Anacystis nidulans, Sigma-Aldrich®) prior to and
- 251 following the cruise. The [chl-a] of the 0.3-2.7 µm size class (hereafter, "pico" size class) was
- 252 calculated by subtracting the measured [chl-a] of the >2.7 µm size class (hereafter, "nano+" size
- 253 class) from the >0.3 µm size class (hereafter, "bulk"). Given previous work showing that the
- 254 winter Southern Ocean phytoplankton community is composed primarily of small cells (i.e.,
- 255 typically <15 µm; e.g., Hewes et al., 1985; 1990; Weber & El-Sayed, 1987), we did not separate
- 256 micro- from nanophytoplankton.

257 3.2.4. Bulk POC, PON and δ^{15} N-PON

- 258 The NPP and N uptake filters were fumed with hydrochloric acid in a desiccator for 24 hours to
- 259 remove inorganic carbon, then dried for 24 hours at 40°C and packaged into tin cups. Filters for
- δ^{15} N-PON were dried in the same way, but not acidified. Samples were analysed using a Delta
- V Plus isotope ratio mass spectrometer (IRMS) coupled to a Flash 260 elemental analyser, with
- 262 a detection limit of 0.17 μ mol C and 0.07 μ mol N and precision of ± 0.005 At% for C and
- N. Unused pre-combusted filters (blanks) were included in each batch run. POC and PON content
- 264 was determined from daily standard curves of IRMS area versus known C and N masses. For the
- 265 isotope ratios, sample measurements were referenced to internal laboratory standards calibrated
- against IAEA reference materials that were measured after every 5-7 samples.

3.2.5. Size-fractionated rates of NPP and N uptake

- 268 Carbon and N uptake rates (NPP, ρNH_4^+ , ρNO_3^- , $\rho Urea$) were calculated according to Dugdale
- 269 & Wilkerson (1986) as:

$$\rho M = \frac{[PM] \times (At\%_{meas} - At\%_{amb})}{T \times (At\%_{init} - At\%_{amb})}$$
(Eqn 1)

where,
$$At\%_{init} = \frac{([M] \times At\%_{amb}) + ([M_{tracer}] \times At\%_{tracer})}{[M] + [M_{tracer}]}$$
 (Eqn 2)

- Here, M is the species of interest (C, NH₄+, NO₃-, or urea); ρM is the uptake rate of that species
- 273 (nM hour⁻¹, i.e., nmol C or N L⁻¹ hour⁻¹); [PM] is the concentration of POC or PON (μ M) on the
- filters; [M] is the ambient concentration of DIC, NH₄⁺, NO₃⁻, or urea at the time of sample
- collection; [M_{tracer}] is the concentration of NaH¹³CO₃, ¹⁵NH₄+, ¹⁵NO₃-, or ¹⁵N-urea added to the
- 276 incubation bottles; and T is the incubation period (days). DIC concentrations were measured
- 277 shipboard using a VINDTA 3C instrument and ranged from 2017 to 2130 μM (Bakker et al.,
- 278 2016). The PM and ρM of the picoplankton size class was calculated by subtracting the
- 279 nanoplankton from the bulk measurements. Daily rates were computed by multiplying the hourly
- 280 rates by the number of daylight hours, the latter calculated using the sampling latitude and day
- of the year (Forsythe et al., 1995).
- 282 The f-ratio (Eppley & Peterson, 1979), used to estimate the fraction of NPP potentially available
- 283 for export, was calculated as:

$$f - ratio = \frac{\rho N o_3^-}{\rho N_{tot}}$$
 (Eqn 3)

- where $\rho N_{tot} = \rho N H_4^+ + \rho N O_3^- + \rho U rea$. Urea uptake was not measured at underway stations
- 286 50.7°S and 55.5°S (both in the Antarctic Zone); here, the f-ratio was calculated omitting ρUrea.
- 287 For the two Antarctic Zone stations at which urea uptake was measured, including pUrea
- 288 decreased the f-ratio by 8-25% compared to that calculated using only ρNO₃⁻ and ρNH₄⁺.

289 3.2.6. Ammonia oxidation rates

- 290 The azide method (McIlvin and Altabet 2005) was used to convert NO₂ produced by NH₄+
- 291 oxidation to N₂O gas that was measured using a Delta V Plus IRMS with a custom-built purge-

and-trap front end (McIlvin & Casciotti, 2011). This configuration yields a detection limit of 0.2 nmol N with a δ^{15} N precision of \pm 0.1‰. The δ^{15} N of NO₂⁻ was derived from ⁴⁵N₂O/⁴⁴N₂O and the rate of NH₄⁺ oxidation (NH₄⁺ox. nM day⁻¹) was calculated following Peng et al. (2015) as:

$$NH_{4 \text{ ox}}^{+} = \frac{A[^{15}NO_{2}^{-}]}{f_{NH_{2}^{+}}^{15} \times T}$$
 (Eqn 4)

Here, Δ [$^{15}NO_2^-$] is the change in the concentration of $^{15}NO_2^-$ (nM) between the start and end of the incubation, calculated as the difference in the measured $\delta^{15}N$ of NO_2^- between the T_f and T_0 samples, $f_{NH_4^+}^{15}$ is the fraction of the NH₄+ substrate labelled with ^{15}N at the start of the incubation, and T is the incubation length (days). All $^{15}NO_2^-$ produced during the incubations was assumed to derive from $^{15}NH_4^+$ oxidation. The detection limit ranged from 0.02 to 0.11 nM day⁻¹, calculated according to Santoro et al. (2013).

We note that isotope dilution (i.e., the dilution of ¹⁵NH₄⁺ by co-occurring ¹⁴NH₄⁺ regeneration) during the NH₄⁺ uptake and oxidation experiments could potentially lead to an underestimation of the rates (Glibert et al., 1982; Mdutyana, 2021). For the NH₄⁺ uptake experiments, their short duration (3 to 7.5 hours) would have rendered the effect of regeneration minor (Mdutyana et al., 2020). Moreover, the ¹⁵NH₄⁺ additions were high (100 nM) relative to both the ambient NH₄⁺ concentrations north of the SAF and the K_m values derived for NH₄⁺ uptake and oxidation in the winter Southern Ocean (150-405 nM and 28-137 nM, respectively; Mdutyana, 2021), making a significant dilution effect unlikely (Lipschultz, 2008). Finally, at the stations south of the SAF, the ambient NH₄⁺ concentrations were so high that even if the regeneration of ¹⁴NH₄⁺ occurred at an elevated rate (e.g., 50 nM day⁻¹; as has been measured in the late-summer Southern Ocean when remineralization is expected to be high; Goeyens et al., 1991), the ¹⁵N/¹⁴N of the NH₄⁺ pool

A further consideration is possible stimulation of the NH₄⁺ uptake and oxidation rates by ¹⁵NH₄⁺ addition (Lipschultz, 2008). Given the K_m values listed above and the high ambient NH₄⁺ concentrations measured in the PFZ and AZ, a stimulation effect could only be significant at the stations north of the SAF where the NH₄⁺ concentrations were 10-100 nM, and even then, to a lesser extent for NH₄⁺ oxidation than NH₄⁺ uptake given that ammonia oxidizers in the winter Southern Ocean become saturated at NH₄⁺ concentrations of 100-200 nM (Mdutyana, 2021). The rates reported for the stations north of the SAF should therefore be considered "potential rates." However, since our focus is mainly on explaining the accumulation of NH₄⁺ south of the SAF, having "potential" rather than "true" rates for the STZ and SAZ does not affect our conclusions.

would decrease by <1-2%. We thus consider the potential effect of isotope dilution to be minor.

3.2.7 Plankton community composition

Microplankton groups (>15 μm) were identified and counted in a subsample (20 mL) from each amber bottle using the Utermöhl technique (Utermöhl, 1958) and following the recommendations of Hasle (1978). Plankton groups and individual species were counted and identified using an inverted light microscope (Olympus CKX41) at 200x magnification. This level of magnification limited the cell sizes that could be reliably distinguished to >15 μm. For each sample, at least 100 cells were enumerated to ensure a statistically valid count.

- 330 Pico- and nanoplankton cells (<15 μm) were enumerated using an LSR II flow cytometer (BD
- Biosciences) equipped with blue, red, violet, and green lasers. Prior to analysis, 1 mL of sample
- was incubated with 1% (v/v) SYBR Green-I (a DNA stain) at room temperature in the dark for
- 333 10 minutes (Marie et al., 1997). From light scatter and autofluorescence, the DNA-containing
- particles were identified as nano- and picoeukaryotes, and *Synechococcus*. Additionally, small
- heterotrophic prokaryotes (i.e., bacteria and possibly archaea; hereafter "bacteria") were
- identified as DNA-containing particles with the lowest detectable autofluorescence (Marie et al.,
- definite as Divisioning particles with the lowest detectable automatics energy (wante et al.,
- 337 1997; Gasol & del Giorgio, 2000) (see also Text S2). All particles lacking DNA were considered
- 338 detritus. The populations of interest were gated using FlowJo 10.3 software (TreeStar, Inc.;
- 339 www.flowjo.com).
- 340 In this study, we did not directly measure NH₄⁺ regeneration (i.e., heterotrophy). Instead, we use
- 341 the abundance of heterotrophic bacteria as a qualitative indicator of NH₄⁺ regeneration potential,
- 342 recognizing that cell abundance does not imply activity. Additionally, we estimate the rate of
- 343 NH₄⁺ production from our concentration and rate data (see section 3.3). The availability of
- 344 organic matter to heterotrophs is inferred from the abundance of detritus.
- 3.3 Mixed-layer NH₄+ residence time and NH₄+ production rate estimates
- 346 The residence time of the mixed-layer NH₄+ pool can be estimated using the measured ambient
- 347 NH₄⁺ concentrations and corresponding NH₄⁺ removal rates as

$$NH_{4 \ residence \ time}^{+} = \frac{[NH_{4}^{+}]}{NH_{4 \ removal \ rate}^{+}}$$
 (Eqn 5)

- Here, NH₄⁺residence time is the time period (days) over which a given NH₄⁺ concentration will be
- depleted assuming a constant NH₄⁺removal rate. We set NH₄⁺removal rate = ρ NH₄⁺ + NH₄⁺ox in winter
- and = ρNH_4^+ in late summer given the evidence for negligible mixed-layer NH_4^+ oxidation rates
- in this latter season (Bianchi et al., 1997; Mdutyana et al., 2020).
- 353
- 354 To determine the contribution of late summer NH₄+ production to the wintertime NH₄+ pool (see
- section 5.2), we define a rate of NH₄+concentration decline:

$$NH_{4 \ rate \ of \ decline}^{+} = NH_{4 \ production \ rate}^{+} - NH_{4 \ removal \ rate}^{+}$$
 (Eqn 6)

- Here, NH₄+_{production rate} is the NH₄+ flux required to compensate for NH₄+ removal over the late-
- 358 summer-to-winter period, in order to yield the observed seasonal change in the ambient NH₄+
- 359 concentration.
- The rate of NH₄+concentration decline can also be defined as:

$$NH_{4 \text{ rate of decline}}^{+} = \frac{[NH_{4}^{+}]_{decline}}{t}$$
 (Eqn 7)

- Where [NH₄+]_{decline} is the difference between the late summer and winter NH₄+ concentrations
- 363 and t is the time period (days) over which the NH_4^+ concentration declines. Setting Eqn 6 and 7
- 364 equal yields:

$$NH_{4\ production\ rate}^{+} = \frac{[NH_{4}^{+}]_{decline}}{t} + NH_{4\ removal\ rate}^{+}$$
 (Eqn 8)

- Where, $NH_{4}^{+}_{vemoval rate} = \rho NH_{4}^{+} + NH_{4}^{+}_{ox}$. Eqns 7 and 8 assume that the elevated wintertime NH_{4}^{+}
- 367 concentrations result from continuous NH₄+ production in excess of removal rather than from
- sporadic events of removal and/or production occurring between late summer and winter.

369 3.4 Statistical analyses

- 370 The correlations among latitude, N concentrations, NPP, N assimilation rates, and NH₄+ oxidation
- 371 rates were investigated at the 5% significance level using the Pearson correlation coefficient and
- 372 the R packages, stats (R Core Team, 2020) and corrplot (Wei & Simko, 2017). Standard
- 373 deviations were propagated using standard statistical practices.

374 **4. Results**

375

4.1 Hydrography

- 376 Sea surface temperature (SST) decreased by ~17 °C between Cape Town (~34°S) and the edge
- 377 of the MIZ (61.7°S), with similar gradients measured for legs S and N. During leg N, fairly deep
- 378 MLDs were observed (124-212 m), similar to June and July climatological MLDs compiled from
- 379 Argo float data for this region (Dong et al., 2008). While the focus of this study is the surface
- 380 (i.e., upper ~10 m), we report the MLDs here to show that sampling took place under typical
- winter conditions, with the deep MLDs evincing ongoing winter mixing and associated nutrient
- recharge. Where not specified, the trends discussed below refer to the surface data only.
- see rectain the first specified, the french discussed below refer to the surface data only.
- 383 Latitudinal variations in each parameter are assessed by comparing the various Southern Ocean
- zones the Subtropical Zone (STZ) north of the Subtropical Front (STF), the Subantarctic Zone
 (SAZ) between the STF and the Subantarctic Front (SAF), the Polar Frontal Zone (PFZ) between
- the SAF and the Polar Front (PF), and south of the PF, the Open and Polar Antarctic Zones (OAZ
- and PAZ, which are divided by the Southern Antarctic Circumpolar Current Front (SACCF) and
- 388 collectively termed the Antarctic Zone (AZ); see Text S1 for detailed definitions of the fronts
- and zones, and Fig. 1 and S1 for their positions at the time of sampling). For each parameter, the
- average \pm 1 standard deviation (SD) for each Southern Ocean zone is reported in Table 1.

391 4.2 Macronutrient concentrations

- 392 In winter 2017, the surface and mixed-layer concentrations of NH₄⁺ ranged from below detection
- 393 to 0.70 μM (Fig. 2a and b). Surface concentrations were higher in the PFZ, OAZ, and PAZ (0.42
- 394 \pm 0.01 μ M, 0.52 \pm 0.01 μ M, and 0.58 \pm 0.01 μ M, respectively) than in the STZ and SAZ (0.08 \pm
- 395 $0.03~\mu\text{M}$ and $0.06\pm0.01~\mu\text{M}$, respectively), with a sharp gradient observed at the SAF. South of
- 396 the SAF, high NH₄⁺ concentrations persisted near-homogeneously throughout the mixed layer,
- with mixed layer averages ranging from $0.65 \pm 0.01~\mu M$ at station $58.5^{\circ}S$ to $0.27 \pm 0.01~\mu M$ at
- 398 station 48.0°S and averaging $0.47 \pm 0.02 \mu M$, with concentrations that were below detection
- 399 north of the SAF (Fig. 2b). Below the mixed layer, NH₄+ concentrations decreased rapidly at all
- 400 stations to values below detection by 200 m.
- The concentrations of NO_3^- and PO_4^{3-} increased southwards from $\leq 10~\mu M$ and $\leq 1~\mu M$ in the STZ
- 402 to >20 μM and >1.5 μM in the PFZ, OAZ, and PAZ (Fig. 2c and S3a), with the sharpest gradients
- occurring near the SAF. The concentrations of Si(OH)₄ increased rapidly across the PF, from an
- 404 average of $3.2 \pm 1.1 \,\mu\text{M}$ between 35.0°S and 48.0°S to $45.6 \pm 0.6 \,\mu\text{M}$ between 52.1°S and 58.9°S

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- 406 (Fig. S3b). The NO_2 concentrations were consistently low across the transect (0.16 ± 0.02 μ M;
- 407 Fig. S3c), as were the concentrations of urea $(0.20 \pm 0.04 \,\mu\text{M}; \text{Table 1})$, with slightly lower urea
- 408 concentrations observed in the SAZ than in the other zones.

409 <u>4.3 Chlorophyll-a, POC and PON</u>

- 410 The highest bulk [chl-a] was observed near the South African continental shelf, decreasing across
- 411 the STF and remaining low thereafter (Fig. 3a). The proportion of chl-a in the nano+ size class
- varied across the region but was >50% at all stations, with higher (>80%) contributions near the
- 413 fronts and at many OAZ and PAZ stations (Fig. 3b). The nano+ contribution was ≤60% at only
- 414 five stations (three in the SAZ, two in the OAZ).
- 415 The concentrations of bulk POC and PON were highest north of the STF and slightly higher in
- 416 the OAZ than in the SAZ and PFZ (Fig. S4a and b). The contribution of the nano+ size fraction
- 417 to POC and PON across the transect was $77.1 \pm 22.6\%$ and $66.9 \pm 24.2\%$, respectively (Fig. S4c
- 418 and d). The δ^{15} N-PON decreased southwards from the STZ and SAZ (1.7 ± 1.0‰) to the PFZ
- 419 and OAZ (0.5 \pm 0.5%; Fig. 4). Despite considerable differences among zones, the δ^{15} N-PON
- 420 was relatively homogenous within each zone.

421 4.4 Rates of net primary production, nitrogen uptake, and ammonium oxidation

- 422 Rates of bulk NPP were two- to six-fold higher in the SAZ and PFZ than has been reported
- previously for the Atlantic sector in winter (Mdutyana et al., 2020; Froneman et al., 1999) (Fig.
- 424 5a). By contrast, NPP was low in the OAZ, consistent with previous measurements (Kottmeier
- 425 & Sullivan, 1987; Mdutyana et al., 2020). The relative contribution of the nano+ size class
- 426 generally decreased southwards, from 85.4% at 37.0°S to 24.4% at 53.5°S, before increasing to
- 427 >80% near the SACCF.
- 428 The bulk NH₄⁺ uptake rates (ρNH₄⁺) generally increased southwards from the STZ to the SAZ
- 429 and PFZ, and then decreased across the OAZ to reach a minimum at the southernmost station
- 430 (Fig. 5b). In the nano+ size fraction, ρNH₄+ changed little latitudinally, although it was slightly
- 431 lower in the PFZ than in the other zones. The contribution of nanoplankton to ρNH₄⁺ ranged from
- 432 32.8% in the PFZ to 71.9% in the STZ. The bulk NO_3 uptake rates (ρNO_3) were also low in the
- 433 STZ, while the highest ρNO₃ was measured in the SAZ, with the rate then decreasing
- 434 southwards. ρNO₃ in the nano+ size class followed the same trend as total community ρNO₃,
- with the nanoplankton accounting for $71.5 \pm 0.3\%$ of bulk ρNO_3 on average. The rates of bulk
- 436 urea uptake (ρUrea) were highest in the STZ, with the SAZ and the PFZ hosting similar rates,
- and the lowest rates were measured in the OAZ. pUrea for the nano+ size class followed a similar
- 438 trend to bulk ρUrea, and nanoplankton accounted for 51.8% of ρUrea in the SAZ, increasing to
- 439 100% in the PAZ. The uptake rates of the different N forms were not significantly correlated
- with one another or with the ambient N concentrations (Table S1).
- 441 Ammonium oxidation rates (NH₄⁺_{ox}) increased southwards, with higher NH₄⁺_{ox} in the OAZ and
- PAZ than in the STZ, SAZ, and PFZ (Fig. 5c). NH₄+_{ox} was generally comparable to previous
- 443 wintertime measurements from the surface of the open Southern Ocean (Mdutyana et al., 2020).
- NH₄⁺ox was not correlated with the ambient NH₄⁺ concentration (Table S1).

4.5 Plankton community composition

445

- 446 Microplankton abundance was low, with the highest cell counts recorded at stations 37.2°S and
- 447 41.3°S in the STZ and no cells counted at 38.1°S (STZ) and 55.5°S (OAZ) (Fig. 6a). On average,
- 448 microplankton abundance was higher in the STZ than in the SAZ, PFZ, and OAZ. The greatest
- 449 diversity of microplankton groups was observed at 41.3°S in the STZ and at 50.0°S near the PF.
- 450 Centric diatoms (including Planktoniella, Coscinodiscus, and Thalassiosira species) were
- 451 detected only at the southernmost station 58.9°S (3 cells mL-1). Pennate diatoms (including
- 452 Pseudo-nitzschia, Pleurosigma, and Navicula species) were more abundant in the STZ, PFZ, and
- 453 OAZ, with negligible abundances in the SAZ. Higher pennate diatom abundances occurred near
- 454 the PF (7 cells mL⁻¹), as has been observed in summer (e.g., Bracher et al., 1999). Dinoflagellates
- 455 were identified at every station except 38.1°S and were most abundant in the STZ and PFZ. At
- 456 all but three stations, small (\sim 15 μ m) dinoflagellates were the most abundant group, although the
- 457 larger Protoperidinium dinoflagellate species (mainly heterotrophic; Jeong & Latz, 1994) were
- 458 almost as abundant in the PFZ and at 54.0°S. Microzooplankton (i.e., ciliates, 20-200 μm) were
- 459 most abundant in the STZ, and were also present in the PFZ at $46.1^{\circ}S$ (3 cells mL⁻¹) and $48.9^{\circ}S$
- (3 cells mL⁻¹) and in the OAZ at 50.0°S (1 cells mL⁻¹) and 54.0°S (4 cells mL⁻¹). All other 460
- 461 stations were characterized by negligible (<1 cells mL⁻¹) microzooplankton abundances.
- 462 Nano- and picoeukaryotes, Synechococcus, and heterotrophic bacteria (collectively, "small
- 463 cells") were roughly 10³-times more abundant than the microplankton (Fig. 6b). Notwithstanding
- 464 a lack of data from the STZ, the highest small cell abundances occurred in the SAZ near the SAF.
- 465 Across the transect, picoeukaryotes were generally more abundant than all other phytoplankton
- groups (average picoeukaryote contribution to total small cells of 12-54%; nanoeukaryotes of 7-466
- 39%; Synechococcus of 15-42%). A similar trend has been observed for the Southern Ocean in 467
- 468 spring (Detmer & Bathmann, 1997) and late summer (Fiala et al., 1998), in contrast to mid-
- 469 summer observations showing nanoplankton dominance (e.g., Ishikawa et al., 2002; Weber &
- 470 El-Sayed, 1987). Additionally, picoeukaryotes were two- to three orders of magnitude more
- 471 abundant in the SAZ and PFZ than in the OAZ. Nanoeukaryotes dominated near the PF at 50.0°S
- 472 (39%) and in the southern OAZ at 55.5°S (36%), while Synechococcus dominated at 42.7°S and
- 473 54.0°S (42% and 33%, respectively). In general, nanoeukaryote abundance was higher in the
- 474 SAZ than in the PFZ and OAZ, as was that of Synechococcus.
- 475 The contribution of heterotrophic bacteria to total small cells varied considerably (10-62%),
- 476 reaching a maximum south of the PF at 53.0°S and 57.8°S (62% and 50%), and with higher
- 477 abundances in the SAZ than in the PFZ and OAZ (Fig. 7). Additionally, heterotrophic bacterial
- 478 abundances were ten-fold lower to two-fold higher than the total pico- and nanophytoplankton
- 479 cell counts. Detrital particles were most abundant near the southern edge of the SAF, and were
- 480 generally more abundant in the PFZ than in the SAZ and OAZ (Fig. S5).

481 4.6 2018/19 cruises: ammonium concentrations

- 482 In early summer, surface NH₄⁺ concentrations were uniformly low across the transect (average
- of $0.11 \pm 0.09 \mu M$; Fig. 8a). South of the SAF, NH₄+ increased to an average concentration of 483
- 484 $0.81 \pm 0.92 \,\mu\text{M}$ by late summer (Fig. 8b). By winter 2019, the NH₄+ concentrations south of the
- 485 SAF were ~40% lower than they had been in late summer (Fig. 8c), and were similar to those

486 observed in winter 2017 (0.50 \pm 0.30 μ M and 0.52 \pm 0.11 μ M, respectively), confirming that our 2017 observations are generally representative of the wintertime Southern Ocean. By early 487 spring, the NH₄⁺ concentrations south of the SAF had declined to near or below detection (0.09 488 489 ± 0.08 μM; Fig. 8d) before rising again by late spring to an average value only slightly lower 490 than that measured in winter (0.40 \pm 0.74 μ M; Fig. 8e). However, the late-spring NH₄⁺ 491 concentrations were only elevated in the PFZ (range of 0.11 ± 0.01 to 4.39 ± 0.03 µM, average 492 of $0.77 \pm 1.11 \,\mu\text{M}$), as has been observed previously (Bathmann et al., 1997). Excluding the PFZ 493 data yields a far lower late-spring average of $0.17 \pm 0.11 \,\mu\text{M}$ south of the SAF, which we take 494 as more broadly representative of this season.

4.7 Mixed-layer NH₄⁺ residence time and NH₄⁺ production rate estimates

The NH₄+ $_{residence\ time}$ in winter 2017, computed using Eqn 5, ranged from 10 to 38 days (median of 21 days) south of the SAF and from 0 to 6 days (median of 2 days) north of the SAF. These values were estimated using wintertime measurements only and as such, may not be representative of the transition from summer to winter. To refine our estimates, we used average ρ NH₄+ and NH₄+ concentration measurements. South of the SAF in late summer, ρ NH₄+= 50.6 ± 24.0 nM day⁻¹ and the NH₄+ concentration = 0.81 ± 0.92 μ M (Deary, 2020), which together yield an NH₄+ $_{residence\ time}$ of 2 to 27 days (median of 5 days). The NH₄+ $_{residence\ time}$ north of the SAF, calculated using ρ NH₄+= 20.7 ± 8.6 nM day⁻¹ and NH₄+ concentration = 0.16 ± 0.45 μ M (Deary, 2020), was 1 to 17 days (median of 14 days).

The NH₄+ $^+$ production rate south of the SAF, calculated using Eqn 8 and an [NH₄+ $^+$]decline of 330 nM (i.e., the difference between late summer and winter 2019; 810 nM – 480 nM), t of 141 days, and NH₄+ $^+$ removal rate of 50.6 ± 24.0 nM day⁻¹ (here, the average late-summer pNH₄+ $^+$ south of the SAF is used to approximate NH₄+ $^+$ removal rate), was 52.9 ± 25.0 nM day⁻¹. Similarly, north of the SAF (using an [NH₄+ $^+$]decline of 20 nM, i.e., 160 nM – 140 nM, and NH₄+ $^+$ removal rate of 20.7 ± 8.6 nM day⁻¹), the NH₄+ $^+$ production rate was 50.7 ± 9.3 nM day⁻¹. If we instead use the average NH₄+ $^+$ removal rate and NH₄+ concentration measured in winter 2017 south (21.4 ± 0.6 nM day⁻¹ and 520 ± 110 nM) and north (18.4 ± 0.8 nM day⁻¹ and 80 ± 10 nM) of the SAF, the NH₄+ $^+$ production rate was 23.4 ± 6.6 nM day⁻¹ and 18.5 ± 6.6 nM day⁻¹, respectively. Using the range of NH₄+ $^+$ removal rate estimates and the average ambient NH₄+ concentration measured south of the SAF in winter 2017 (16.7 to 31.2 nM day⁻¹ and 520 nM) and late summer 2019 (22.6 to 98.6 nM day⁻¹ and 810 nM), we calculate that over the late-summer-to-winter transition, the NH₄+ $^+$ production rate ranged from 18.8 to 100.9 nM day⁻¹ (compared to 6.3 to 28.8 nM day⁻¹ north of the SAF).

5. Discussion

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5.1 <u>Drivers of NH₄⁺ cycling in the surface layer of the Southern Ocean</u>

Previous work has suggested that NH₄⁺ accumulates in the Southern Ocean mixed layer following the late summer increase in heterotrophy, then decreases into autumn as heterotrophic activity subsides, to be depleted by winter due to advective processes and biological removal (Koike et al., 1986; Serebrennikova & Fanning, 2004). However, our data show that NH₄⁺ concentrations are elevated in the mixed layer in winter, particularly south of the SAF (Fig. 2). Similarly elevated winter surface-layer NH₄⁺ has been observed previously in both the Atlantic and Indian sectors,

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The $NH_4^+_{production\ rate}$ calculated using Eqn 8 and an $[NH_4^+]_{decline}$ of 330 nM (i.e., 810 nM -480 nM), t of 141 days, and $NH_4^+_{removal\ rate}$ of 50.6 ± 24.0 nM day-1 (here, the average late-summer pNH_4^+ south of the SAF is used to approximate $NH_4^+_{removal\ rate}$), was 52.9 ± 25.0 nM day-1. If we instead use the average $NH_4^+_{removal\ rate}$ and $NH_4^+_{concentration}$ measured in winter 2017 (21.4 \pm 0.6 nM day-1 and 520 ± 110 nM), the $NH_4^+_{production\ rate}$ was $23.4 \pm$ 6.6 nM day-1. Using the range of $NH_4^+_{removal\ rate}$ values and the average ambient $NH_4^+_{concentration}$ measured south of the SAF in winter 2017 (16.7 to 31.2 nM day-1 and 520 nM) and late summer 2019 (22.6 to 98.6 nM day-1 and 810 nM), we calculate that over the late-summer-to-winter transition, the $NH_4^+_{production\ rate}$ ranged from 18.8 to 100.9 nM day-1.

572 with concentrations typically increasing towards the south (Philibert et al., 2015; Mdutyana et 573 al., 2020; Bianchi et al., 1997). Numerous overlapping processes are likely involved in setting 574 the ambient NH₄⁺ concentrations, as summarized in Fig. 9. In this study, we directly measured 575 the rates of NH₄⁺ uptake and oxidation, and estimated the rates of NH₄⁺ production, along with 576 qualitatively evaluating the role of heterotrophy from the relative abundance of heterotrophic 577 bacteria, phytoplankton, and detritus. For the NH₄⁺ cycle processes shown in Fig. 9 that are not 578 quantified or inferred from our dataset, we consider their potential role in Southern Ocean NH4+

579 cycling based on findings reported in the literature.

580 The high NH₄⁺ concentrations observed south of the SAF in winter may result from net NH₄⁺ 581 accumulation during late summer, autumn, and/or winter. The persistence of elevated NH₄⁺ 582 concentrations that are near-homogeneously distributed throughout the mixed layer is consistent 583 with a residence time for the winter NH₄⁺ reservoir in excess of the time-scale for upper-ocean 584 mixing. Indeed, we calculate a median residence time of 21 days south of the SAF, compared to 585 2 days north of the SAF. One implication of the long residence time computed for the polar zones 586 is that the wintertime NH4+ pool likely reflects both ongoing processes and those that occurred 587 earlier in the year. We posit that the elevated NH₄+ concentrations south of the SAF may result 588 from higher wintertime rates of NH₄₊ production than removal and/or from the gradual but incomplete depletion in winter of NH_4^+ produced mainly in late summer and autumn. We evaluate 589 590 both possibilities throughout the discussion below.

5.1.1 Ammonium removal

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592 Ammonium assimilation - Microbial growth is limited in the winter Southern Ocean (Arrigo et 593 al., 2008; Smith Jr et al., 2000, Takao et al., 2012), resulting in low cell abundances and nutrient 594 uptake rates (Church et al., 2003; Iida & Odate, 2014; Mdutyana et al., 2020). However, while 595 the concentrations of chl-a and rates of NPP were low across our transect, they were not 596 negligible (Fig. 3a and 5a), consistent with previous reports for this season (Mordy et al., 1995; 597 Pomeroy & Wiebe, 2001). Southern Ocean phytoplankton are adapted to survive suboptimal 598 conditions; for example, numerous species achieve their maximum growth rates at temperatures 599 that are considerably lower than the optimal growth temperatures of temperate and tropical 600 species (2-9 °C versus 10-30 °C and 15-35 °C, respectively), with sharp declines in growth rates 601 observed at temperatures outside this range (Boyd et al., 2013; Coello-Camba & Agusti, 2017; 602 Fiala & Oriol, 1990). In addition, ice-free Southern Ocean waters typically extend to <60°S in 603 the eastern Atlantic and western Indian sectors in winter, so that even though irradiance levels 604 may not be optimal for phytoplankton growth, there is always some light available for 605 photosynthesis. The hostile wintertime conditions of the open Southern Ocean do not, therefore, 606 prevent ecosystem functioning, although the microbial dynamics and associated biogeochemical 607 processes differ from those occurring in summer (Smart et al., 2015; Mdutyana et al., 2020).

608 We measured fairly low surface NH₄⁺ uptake rates (3.0-13.2 nM day⁻¹; Fig. 5b) compared to 609 previous wintertime observations (ranging from 32-66 nM day-1; Cota et al., 1992; Mdutyana et 610 al., 2020; Philibert et al., 2015). Such low rates, if generally representative of winter, would limit 611 mixed-layer NH₄⁺ drawdown, especially south of the PF where ρNH₄⁺ was particularly low. 612 Recycled N (NH₄⁺ + urea) nonetheless accounted for most of the N assimilated during winter,

613 including in the AZ (Fig. 5b). 614 The available δ^{15} N-PON data suggest that the preferential reliance of phytoplankton on recycled 615 N may have persisted from the late summer. In theory, PON generated in early-through midsummer from the assimilation of upwelled NO₃⁻ (δ^{15} N-NO₃⁻ of 5.2‰ in the AZ and 6.2‰ in the 616 SAZ; Smart et al., 2015; Fripiat et al., 2019; 2021) will have a δ^{15} N of ~0% in the AZ and 1-2% 617 618 in the SAZ given the isotope effect of NO₃⁻ assimilation and the degree of seasonal NO₃⁻ 619 drawdown (Sigman et al., 1999; Granger et al., 2004; 2010). Such δ^{15} N-PON values have indeed 620 been measured in the early- and mid-summer Southern Ocean (Lourey et al., 2003; Smart et al., 621 2020; Soares et al., 2015). By late summer, δ^{15} N-PON has been observed to decline to between 622 -5 and -1‰, with the lowest values occurring in the AZ (Lourey et al., 2003; Smart et al., 2020; 623 Trull et al., 2008). Since the δ^{15} N of recycled N is expected to be low (<0%; Checkley & Miller, 1989, Macko et al., 1986), the early-to-late summer decline in δ^{15} N-PON implicates a switch 624 625 from dominantly NO₃-- to dominantly recycled N-supported phytoplankton growth (Lourey et al., 2003). For the SAZ, the subsequent late summer-to-winter rise in δ^{15} N-PON (i.e., from \sim -626 627 1‰ to 1-2.5‰; Fig. 4) has previously been attributed to PON decomposition by heterotrophic bacteria (Smart et al., 2020), during which ¹⁴N-NH₄⁺ is preferentially remineralized, leaving the 628 629 remaining PON enriched in ¹⁵N (Möbius, 2013). That NH₄+ concentrations are not elevated in 630 the SAZ mixed layer in winter (Fig. 2b) indicates that the remineralized NH₄⁺ is rapidly re-631 assimilated by phytoplankton and/or oxidized to NO2 in this zone. In the AZ, the much lower 632 δ^{15} N-PON of -3 to -1‰ that we observe in winter surface waters requires the sustained 633 assimilation of low- δ^{15} N N (i.e., recycled N) to offset a remineralization-driven δ^{15} N rise akin to 634 that of the SAZ. We conclude that Southern Ocean phytoplankton preferentially consume 635 regenerated N from late summer until at least July (albeit at low rates in winter), particularly 636 south of the PF.

637 The fact that NH₄+ accumulated in the winter mixed layer despite being the preferred 638 phytoplankton N source in late summer through winter implies that low rates of NH₄⁺ uptake 639 contributed to its accumulation. Multiple factors may cause low rates of photoautotrophic NH₄⁺ 640 assimilation, including deplete NH4+ and micronutrient concentrations, light limitation, and low 641 temperatures. North of the SAF, NH₄⁺ concentrations below detection likely limited ρNH₄⁺, as 642 evidenced by the fact that in a series of experiments conducted on the same cruise, ρNH₄⁺ 643 increased with the addition of NH₄⁺ at these stations (Mdutyana, 2021). By contrast, south of the 644 SAF, NH₄⁺ concentrations were similar to or higher than the half-saturation constant (K_m) derived 645 for NH₄⁺ uptake in the winter Southern Ocean (0.2 to 0.4 μM; Mdutyana, 2021), suggesting that 646 something other than NH₄⁺ availability was limiting to phytoplankton at these latitudes.

647 Iron is not directly involved in NH₄⁺ assimilation but is required for electron transport during 648 photosynthesis and respiration, as well as for chlorophyll synthesis (Raven, 1988). While iron 649 limitation is widespread across the Southern Ocean (Janssen et al., 2020; Pausch et al., 2019; 650 Viljoen et al., 2019), iron availability appears to be higher in winter than during other seasons 651 (Mtshali et al., 2019; Tagliabue et al., 2014) due to enhanced mixing, storms, and increased 652 aeolian deposition (Coale et al., 2005; Honjo et al., 2000; Sedwick et al., 2008). The fact that 653 ρNO₃⁻ and ρNH₄⁺ were generally similar across the transect (Fig. 5b) argues against a dominant 654 role for iron in controlling ρNH₄⁺ since NO₃⁻ consumption has a far higher iron requirement than

655 NH₄⁺ assimilation (Morel et al., 1991).

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657 In contrast to NH4+ and iron availability, light limitation is exacerbated in winter due to low 658 insolation, increased cloud-cover, and mixed layers that can be hundreds of meters deeper than 659 the euphotic zone (Buongiorno Nardelli et al., 2017; Sallée et al., 2010). Light is thus often 660 considered the dominant constraint on Southern Ocean primary productivity in this season 661 (Thomalla et al., 2011; Llort et al., 2019; Wadley et al., 2014). However, since NH₄+ assimilation by phytoplankton is fairly energetically inexpensive (Dortch, 1990), it should occur even under 662 low light conditions (recognizing that light remains critical for coincident CO2 fixation). 663 Heterotrophic bacteria can also consume NH₄⁺ (Kirchman, 1994), including in the dark, as they 664 derive energy from organic carbon oxidation rather than light. At an ecosystem level, therefore, 665 666 NH₄⁺ assimilation may not be primarily limited by light, although this parameter clearly strongly 667 controls the rate and distribution of NPP (Fig. 5a).

668 Previous observations suggest that temperature can influence NH₄⁺ uptake, especially in winter 669 (Glibert, 1982; Reay et al., 2001). The negative effect of temperature appears to be enhanced 670 under high-nutrient and low-light conditions, at least in the case of phytoplankton growth rates 671 (Baird et al., 2001). Experiments conducted coincident with our sampling showed that the 672 maximum rate of NH4+ uptake (Vmax) achievable by the in situ community was strongly negatively correlated with temperature and latitude (Mdutyana, 2021), with the latter parameter 673 674 representing the combined role of light, temperature, and possibly iron, the average concentration of which appears to increase from the SAZ to the AZ (Tagliabue et al., 2012). We conclude that 675 these three drivers, along with NH4+ availability north of the SAF, may all play a role in 676 677 controlling photoautotrophic NH₄⁺ assimilation in the winter Southern Ocean, with complex 678 interactions among them that are difficult to disentangle.

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In addition to physical and chemical limitations, microbial preference for other N species may impact NH₄⁺ depletion. For example, the preferential uptake of urea and/or other dissolved organic N (DON) species by some organisms (e.g., picoeukaryotes, cyano- or heterotrophic bacteria) could cause a net decrease in the total NH₄⁺ uptake rates. While urea has been shown to constitute a large fraction of the total N assimilated by Southern Ocean phytoplankton in summer and autumn (albeit mainly in the SAZ; Joubert et al., 2011; Thomalla et al., 2011), we measured fairly low pUrea (Fig. 5b), which is perhaps unsurprising given the low ambient urea concentrations (Table 1). The exceptions were stations 37°S and 43.0°S where pUrea was higher than pNH₄⁺, coincident with very low ambient NH₄⁺ (0.10 μM and below detection) and relatively high urea concentrations (0.36 μM and 0.15 μM, respectively).

689 Community composition can also alter the N uptake regime. Small phytoplankton, such as the 690 numerically-dominant nano- and picoeukaryotes, are more likely to consume NH4+ and urea than 691 NO₃ (Koike et al., 1986; Lee et al., 2012; 2013), especially under conditions of iron and light 692 limitation (Sunda & Huntsman, 1997). Across our transect, reduced N (i.e., NH₄⁺ + urea) uptake 693 exceeded NO₃ uptake for both the total phytoplankton community (transect average of $12.0 \pm$ 694 0.9 nM day^{-1} for reduced N versus $5.8 \pm 1.0 \text{ nM day}^{-1}$ for NO₃; f-ratio of 0.36) and the pico size 695 fraction $(5.0 \pm 1.2 \text{ nM day}^{-1} \text{ versus } 1.9 \pm 1.2 \text{ nM day}^{-1}; \text{ f-ratio of } 0.27; \text{ Fig. 5b})$. That said, the 696 NO₃⁻ uptake rates were not negligible, including in the pico size fraction. In the PFZ and AZ, 697 NO₃ uptake by the picoplankton was far more strongly correlated with the abundance of 698 picoeukaryotes than Synechococcus (r = 0.75 and 0.03, respectively), consistent with 699 observations of dominant reliance on NO₃⁻ by picoeukaryotes and NH₄⁺ by Synechococcus in

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701 other ocean regions (Fawcett et al., 2011; 2014; Painter et al., 2014). Additionally, 702 Synechococcus abundance was strongly correlated with NH₄⁺ concentration south of the SAF (r 703 = 0.65). In the nano+ size class, NO₃ uptake was likely driven in the SAZ by dinoflagellates and 704 nanoeukaryotes, and in the PFZ and AZ by diatoms, which remain active in these zones in winter 705 (Weir et al., 2020). By contrast, nanoeukaryotes, which have a higher per-cell nutrient 706 requirement than the equally-abundant picoeukaryotes, may have dominated NH4+ uptake in the 707 PFZ and AZ given that higher nanoeukaryote abundances corresponded with lower NH4+ 708 concentrations at a number of stations (e.g., stations 50.0°S, 51.1°S, and 55.5°S; Fig. 6b).

709 The low abundances of diatoms and dinoflagellates and absence of coccolithophores across our 710 transect (Fig. 6a) is expected given the limitations imposed on nutrient uptake and CO₂ fixation 711 by winter Southern Ocean conditions. The lower surface area-to-volume ratio of large cells 712 means that they rapidly experience diffusion-limitation of NH₄⁺ and micronutrient uptake and are 713 more susceptible to light limitation (Finkel et al., 2004), resulting in their being outcompeted by 714 smaller species for essential resources (Franck et al., 2005; Cavender-Bares et al., 1999). The 715 near-absence of centric diatoms is also best explained thus, particularly given their low surface area-to-volume ratio compared to the more-abundant pennate species (Kobayashi & Takahashi, 716 2002) that are more likely to consume NH₄+ (Semeneh et al., 1998). Diatom success in winter 717 718 may also be limited by enhanced mixing, as this group generally prefers stratified waters 719 (Kopczynska et al., 2007).

In sum, NH₄⁺ uptake rates were low across our transect but not negligible, indicating that phytoplankton activity in winter, which is dominated by smaller species, is a sink for NH₄⁺. The hostile conditions of the winter Southern Ocean imposed limitations on NH₄⁺ uptake that varied with latitude, with NH₄⁺ concentrations controlling ρNH₄⁺ north of the SAF, while light and temperature were important south of the SAF. Additionally, *Synechococcus*, nanoeukaryotes, and pennate diatoms likely dominated NH₄⁺ assimilation, consistent with previous observations from

the Southern Ocean and elsewhere (Klawonn et al., 2019; Semeneh et al., 1998).

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731 732 Ammonium oxidation – Nitrification removes more mixed-layer NH₄⁺ in winter than phytoplankton assimilation south of the PF, with NH₄⁺ oxidation rates that were two- to five-times the co-occurring NH₄⁺ uptake rates (Fig. 5c). The comparative success of ammonia oxidisers may be due to decreased competition with phytoplankton for NH₄⁺, augmented by decreased photoinhibition (Wan et al., 2018; Lu et al., 2020), elevated NH₄⁺ availability (Baer et al., 2014; Mdutyana et al., 2020; Mdutyana, 2021), and the apparently minor effect of temperature

on NH₄+ oxidation (Bianchi et al., 1997; Baer et al., 2014; Horak et al., 2013; Mdutyana 2021).

One implication of the dominance of NH₄⁺ oxidation in winter is that in addition to the limitations on photoautotrophic NH₄⁺ assimilation discussed above, low phytoplankton success in the AZ

on photoautotrophic NH₄⁻ assimilation discussed above, low phytoplankton success in the AZ may result from nitrifiers outcompeting phytoplankton for scarce resources (e.g., trace elements

737 required for enzyme functioning, such as iron and copper; Amin et al., 2013; Maldonado et al.,

738 2006; Shafiee et al., 2019) under conditions of low incident light and enhanced mixing.

The K_m derived for NH_4^+ oxidation in the winter Southern Ocean has recently been reported to be low (0.03 to 0.14 μ M), with ammonia oxidizers observed to become saturated at ambient NH_4^+

741 concentrations of ~ 0.1 -0.2 μ M (Mdutyana, 2021). This means that south of the SAF in winter

742 2017, ammonia oxidizers were not substrate limited (as implied by the lack of correlation

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745 between NH₄+_{ox} and NH₄+ concentration; Table S1), which raises the question of why NH₄+

oxidation did not occur at higher rates. The answer may indirectly involve temperature, in that

- 747 psychrophilic organisms can be less responsive to high substrate concentrations at low
- 748 temperatures (Baer et al., 2014). Another possibility is that NH₄⁺ oxidation was iron-limited
- 749 (Shiozaki et al., 2016; Shafiee et al., 2019; Mdutyana, 2021). In any case, ammonia oxidisers
- 750 were moderately successful across the surface Southern Ocean in winter, with low light, reduced
- 751
- competition with phytoplankton, and substrate repletion likely explaining the elevated NH4+
- oxidation rates south of the PF compared to the stations to the north. 752

753 5.1.2 Ammonium production and other sources of ammonium

- 754 NH₄⁺ production must have been sustained during the winter to maintain a mixed-layer NH₄⁺
- 755 pool south of the SAF that was high in concentration relative to the early summer. Indeed, the
- 756 residence time estimated for NH₄⁺ in winter (10 to 38 days) is considerably shorter than the
- 757 transition from late summer to winter (approximately three months), indicating that heterotrophic
- 758 NH₄⁺ production, which would have occurred coincident with NH₄⁺ consumption, must have
- 759 been ongoing in winter. We estimate the rate of this wintertime NH_4^+ production to be 23.4 ± 6.6
- 760 nM day-1.

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- 761 Heterotrophic activity by bacteria - Heterotrophic bacteria contribute significantly to NH4+
- 762 production in the Southern Ocean (Hewes et al., 1985; Koike et al., 1986; Tréguer & Jacques,
- 763 1992), including in winter (Rembauville et al., 2017). In our dataset, lower ratios of
- 764 photosynthetic-to-heterotrophic cells were observed at stations with higher NH₄⁺ concentrations
- 765 (e.g., stations 48.9°S, 53.0°S, 54.0°S, and 57.8°S; Fig. S5a), consistent with a role for the
- 766 heterotrophic bacteria present at the time of sampling in generating the ambient NH₄⁺ pool. The
- 767 potential for ongoing heterotrophic activity can also be inferred from the high detrital particle
- 768 counts along the transect (Fig. 7). However, since heterotrophic bacteria are likely more active
- 769 in late summer and autumn when the temperature and the supply of labile PON are higher
- 770 (Becquevort et al., 2000; Dennett et al., 2001; Pomeroy & Wiebe, 2001; Smart et al., 2020), we
- 771 expect that the winter NH₄⁺ pool includes NH₄⁺ produced in late summer and autumn. A further
- 772 consideration is assimilation of NH₄⁺ by heterotrophic bacteria, reported to occur at elevated
- 773 rates in the Southern Ocean mixed layer in winter (Mdutyana et al. 2020; Text S3). If this process
- 774 is a persistent feature of the winter Southern Ocean, it will decrease the net contribution of
- 775 heterotrophic bacteria to NH4+ accumulation. We conclude that it is unlikely that the surface NH4+
- 776 pool measured in winter derived solely from wintertime bacterial NH₄⁺ production given that yet
- 777 higher NH₄⁺ concentrations have been observed in late summer and autumn (Becquevort et al.,
- 778 2000; Dennett et al., 2001), including in the present study (see section 5.2 below).
- 779 Heterotrophic activity by zooplankton - While the microzooplankton enumerated in this study
- 780 occurred at very low abundances, those that were present likely contributed to the NH₄⁺ flux. For
- 781 example, at stations 48.9°S and 54.0°S in the PFZ and AZ, respectively, both the ratios of
- 782 photosynthetic-to-heterotrophic cells and the absolute abundances of heterotrophic bacteria were
- 783 low, while the microzooplankton abundances and NH₄⁺ concentrations were elevated compared
- 784 to nearby stations. The implication of these observations is that elevated microzooplankton
- 785 abundances may help to explain high NH4+ concentrations in waters with low numbers of
- 786 heterotrophic bacteria, although we note that this scenario only occurred at two stations. On

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balance, we posit that microzooplankton are less important for wintertime NH₄⁺ production than heterotrophic bacteria given their low abundances in the surface layer (Fig. 6a; Atkinson et al.,

791 2012). That said, it is possible that the contribution of micro- (and/or macro-) zooplankton to the

NH₄ pool surpasses that of heterotrophic bacteria under certain conditions (Koike et al., 1986;

Priddle et al., 1998), such as in (late) summer and near regions of frontal upwelling in response

794 <u>to elevated rates of phytoplankton biomass accumulation.</u>

Above, we have assumed that NH₄⁺ production is the direct result of heterotrophy. However, there are other possible mechanisms of NH₄⁺ supply that should be considered. We briefly address some of these processes below, noting that for most, there are very few to no observations

798 available from the Southern Ocean.

799 DON cycling – NH₄⁺ can be released by heterotrophic bacteria that directly consume DON (e.g., 800 urea; Billen, 1983; Tupas & Koike, 1990), and possibly also by ammonia oxidisers that convert 801 DON to NH₄⁺ intracellularly, through the equilibration of the intra- and extracellular NH₄⁺ pools 802 (Kitzinger et al., 2019). DON can also be converted to NH₄⁺ through photodegradation by UV 803 radiation (e.g., Aarnos et al., 2012). Bacterial decomposition of DON (rather than PON) to NH₄+ 804 is implicit in most estimates of ammonification, however, and cellular NH₄⁺ efflux by ammonia 805 oxidisers is likely extremely low given that they require NH₄⁺ to fix CO₂. Additionally, the low 806 light flux to the surface Southern Ocean in winter means that photodegradation will not yield a 807 significant supply of NH₄⁺. Thus, DON conversion to NH₄⁺, through any mechanism, is probably 808 negligible.

809 External inputs of ammonium - High surface ocean NH₄+ concentrations may theoretically derive 810 from external inputs of NH₄+, such as N₂ fixation, NH₄+ aerosol deposition, or sea-ice melt. N₂ 811 fixation should be below detection in the winter Southern Ocean due to the cold temperatures, 812 low light and iron conditions, and high NO₃⁻ concentrations (Jiang et al., 2018; Knapp et al., 813 2012; Kustka et al., 2003). NH₄+ aerosols are unlikely to be abundant over regions of the Southern 814 Ocean remote from islands and coastal Antarctica, particularly in winter when NH₄⁺ aerosol 815 concentrations have been shown to reach a minimum (Legrand et al., 1998; Xu et al., 2019). 816 Moreover, the aerosols that are present over the open Southern Ocean will derive mainly from 817 surface-ocean NH₃ efflux; once re-deposited, this NH₄+ does not constitute a new input to surface 818 waters (Altieri et al., 2021). Finally, since our sampling took place before the sea-ice reached its 819 northernmost extent (Cavalieri & Parkinson, 2008), the dominant process would have been sea-820 ice formation rather than sea-ice melt, the latter an occasional source of NH4+ (Kattner et al., 821 2004; Zhou et al., 2014). In any case, we observed elevated NH₄+ concentrations as far north as

822 46°S, ~1700 km beyond the influence of sea-ice melt.

5.2 Seasonal cycling of NH₄⁺ in the Southern Ocean mixed layer south of the SAF

The NH₄⁺ concentration data collected over the 2018/19 annual cycle provide context for interpreting our winter 2017 dataset, allowing us to address our hypothesis that NH₄⁺ production

826 in late summer and autumn contributes to the elevated NH₄⁺ concentrations measured in winter.

The very low NH₄⁺ concentrations observed in early summer (Fig. 8a) are consistent with high

828 rates of phytoplankton NH₄⁺ assimilation during the spring and early-summer growing period

829 (Mdutyana et al., 2020; Savoye et al., 2004; Daly et al., 2001). By late summer, the NH₄+

830 concentrations increased (Fig. 8b) presumably due to elevated heterotrophic activity (i.e., 831 bacterial decomposition and zooplankton grazing) following the accumulation of algal biomass 832 (Mengesha et al., 1998; Le Moigne et al., 2013), coupled with iron- and/or silicate-limitation of 833 phytoplankton (Hiscock et al., 2003; Sosik & Olson, 2002) and enhanced grazing pressure 834 (Becquevort et al., 2000). Mixed-layer NH₄⁺ remained high between late summer and winter 835 (Fig. 8b-c), likely due to sustained heterotrophic NH₄⁺ production in excess of NH₄⁺ removal. 836 This notion is supported by estimates of the residence time of NH₄⁺. We calculate that in summer, 837 the in situ NH₄+ pool would be depleted in 2 to 27 days (median of 5 days) without coincident 838 NH_4^+ production. In addition, the net decline in NH_4^+ concentration of $0.31 \pm 0.97 \mu M$ between 839 late summer and winter requires an average NH₄⁺ production rate of 52.8 ± 25.0 nM/day given 840 the observed NH₄⁺ assimilation rates. This estimate is remarkably similar to the only 841 measurements of NH4+ regeneration available for the Southern Ocean, measured near the

Antarctic Peninsula in summer (average of 55 nM day⁻¹; Goeyens et al., 1991).

By early spring, the NH₄⁺ concentrations had declined (Fig. 8d), implicating increased photosynthetic activity, and thus nutrient assimilation, following the alleviation of light-limitation. We suggest that any NH₄⁺ remaining in late winter would have been consumed in early spring prior to significant NO₃⁻ drawdown because far less energy (i.e., light) is required for its assimilation (Dortch, 1990). The high NH₄⁺ concentrations subsequently observed in late spring (mainly in the PFZ; Fig. 8e) can be explained by elevated heterotrophic activity in response to high levels of regional phytoplankton growth driven by frontal upwelling of limiting nutrients (Becquevort et al., 2000; Mayzaud et al., 2002).

From our six transects of surface NH₄⁺ concentrations across the Southern Ocean, we propose a seasonal cycle for mixed-layer NH₄⁺ south of the SAF (Fig. 8f). Our proposal is consistent with previous characterizations of the early summer-to-autumn evolution of Southern Ocean NH4⁺ concentrations (i.e., from below detection due to phytoplankton assimilation to elevated due to net heterotrophy). However, it contradicts the hypothesis that NH₄+ will subsequently decline due to persistent but low rates of photosynthesis that yield insufficient biomass to support elevated heterotrophy in autumn, thus driving a coincident decrease in photosynthetic and heterotrophic activity (Koike et al., 1986; Serebrennikova & Fanning, 2004). Instead, our data evince a gradual decline in mixed-layer NH₄⁺ concentrations from late summer through winter. This decline can be explained by heterotrophic NH₄+ production outpacing NH₄+ removal in late summer/autumn, with NH₄⁺ regeneration then decreasing during winter to lower rates than the combined rate of NH₄⁺ assimilation and oxidation. By late spring, NH₄⁺ reaches concentrations similar to those observed in early summer as the improved growing conditions (i.e., elevated light and iron availability; Ellwood et al., 2008; Mtshali et al., 2019) allow phytoplankton to rapidly consume any NH₄⁺ remaining at the end of winter and subsequently produced in spring. An exception to this scenario is elevated, localized NH₄+ production near fronts, such as we observed in late spring 2019, which likely resulted from biological activity supported by frontal upwelling of silicateand iron-bearing Upper Circumpolar Deep Water (Prézelin et al., 2000).

6. Summary and implications

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Our study of the upper Southern Ocean, focused on the infrequently-sampled winter season, provides new insights into the internal cycling of N in the mixed layer of a globally-important

region. We attribute the elevated NH₄⁺ concentrations that persist in the winter mixed layer south of the SAF to sustained heterotrophic NH₄⁺ production in excess of NH₄⁺ removal, driven by temperature-, light-, and possibly iron-limitation of phytoplankton and nitrifiers. We further suggest that heterotrophic bacteria are the main NH₄⁺ producers in winter and that the contribution of external sources to the Southern Ocean's mixed-layer NH₄⁺ pool is negligible. From observations of surface NH₄⁺ concentrations made between December 2018 and November 2019, we deduce that the elevated mixed-layer NH₄⁺ concentrations measured in winter cannot be due solely to wintertime NH₄⁺ production. Instead, we propose that NH₄⁺ accumulates to its highest concentrations in late summer following the peak phytoplankton growing season, after which sustained heterotrophy throughout the autumn and winter prevents this NH₄⁺ from being fully depleted until the early spring, even though the rate of NH₄⁺ removal must exceed that of NH₄⁺ production over this period. Measurements of heterotrophic NH₄⁺ production rates are required to confirm the hypothesized seasonal cycle of NH₄⁺ in the Southern Ocean mixed layer, and higher spatial resolution sampling of plankton community composition and N removal rates may help to explain local variability in NH₄⁺ concentrations, particularly near the fronts.

In net, the Southern Ocean mixed layer is a biological source of CO₂ to the atmosphere in autumn and winter (Mongwe et al., 2018). The persistence of elevated NH₄⁺ concentrations across the polar Southern Ocean between late summer and winter implies that this biological CO₂ production occurs not only because NO₃⁻ drawdown is weak relative to NO₃⁻ supply at this time (e.g., Gibson & Trull, 1999; Gray et al., 2018; Hauck et al., 2015; Mongwe et al., 2018; Shadwick et al., 2015), but also because the ambient conditions allow for NH₄⁺ accumulation. There are additional implications of our observations. For example, NH₄⁺ concentrations >1 μM (and at times >0.5 μM) have been reported to inhibit NO₃⁻ assimilation, including in the Southern Ocean (Cochlan, 1986; Goeyens et al., 1995; Philibert et al., 2015; Reay et al., 2001). Inhibition of NO₃⁻ assimilation due to the seasonal accumulation of NH₄⁺ would constitute an inefficiency in the biological pump. However, we observed little evidence of this effect in winter 2017 – the southward decrease in ρNO₃⁻ was not stronger than that of ρNH₄⁺ despite the latitudinal increase in NH₄⁺ concentration, and we observed no relationship between NH₄⁺ concentration and the proportion of NO₃⁻ to NO₃⁻+NH₄⁺ uptake (i.e., the f-ratio; Table S1).

The implications of NH₄⁺ cycling extend beyond the upper ocean to the atmosphere, since ammonium aerosols that influence Earth's albedo (Tevlin & Murphy, 2019) are formed in the marine boundary layer from reactions of NH₃ gas with acidic species. In the remote Southern Ocean, marine NH₃ emissions, which are the largest natural contributors to NH₃ globally, are likely the dominant local source of NH₃ to the atmosphere (Paulot et al., 2015). Surface ocean NH₄⁺ concentrations play a central role in determining the sign and magnitude of the air-sea NH₃ flux, along with wind speed, surface ocean temperature, and pH. Therefore, the biogeochemical pathways that underpin seasonal changes in surface ocean NH₄⁺ concentrations represent an important control on the remote Southern Ocean air-sea NH₃ flux, with consequences for aerosol composition, cloud formation, and climate (Altieri et al., 2021).

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Figure and Table Captions

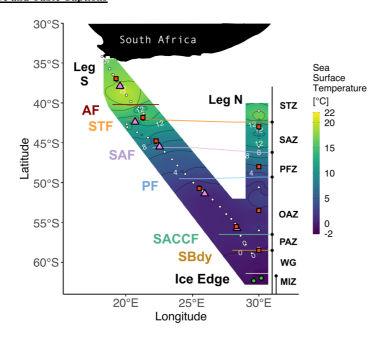


Figure 1: Winter 2017 cruise track overlaid on sea surface temperature (SST) measured by the hull-mounted thermosalinograph. The underway (Leg S) and CTD (Leg N) stations are indicated by white circles. Stations at which net primary production (NPP), nitrogen uptake, and ammonium oxidation experiments were conducted are denoted by red squares. The pink triangles indicate stations where only NPP experiments were conducted while the green circles show stations where only ammonium oxidation was measured. Solid lines indicate the positions of the fronts, identified from measurements of temperature and salinity. Abbreviations for fronts: AF – Agulhas Front (~40.2°S); STF – Subtropical Front (~42.1°S); SAF – Subantarctic Front (~45.6°S); PF – Polar Front (~49.5°S); SACCF – Southern Antarctic Circumpolar Current Front (~56.5°S); SBDY – Southern Boundary (~58.5°S). Abbreviations for zones: STZ – Subtropical Zone; SAZ – Subantarctic Zone; PFZ – Polar Frontal Zone; OAZ – Open Antarctic Zone; PAZ – Polar Antarctic Zone; WG – Weddell Gyre; MIZ – Marginal Ice Zone. Together, the OAZ and PAZ constitute the Antarctic Zone (AZ). See Text S1 for detailed definitions of the fronts and zones. Figure produced using the package ggplot2 (Wickham, 2016).

Table 1: Mean (\pm 1 SD) of surface ocean POC, PON, chl-a, and nutrient concentrations, cell abundances, and nutrient uptake rates measured in each zone of the Southern Ocean in winter 2017. Where no SD is given, only one sample was measured. The >0.3 μm and >2.7 μm size fractions are referred to as "bulk" and "nano+", respectively. "% of nano+" refers to the average relative contribution of the nano+ size fraction to total chl-a, POC, or PON, calculated for each station within a zone. The f-ratio including ρUrea

 is only shown for zones where $\rho Urea$ was measured at all stations. "ND" indicates no data available. Abbreviations as in Figure 1.

	STZ	SAZ	PFZ	OAZ	PAZ
NH ₄ ⁺ (μM)	0.08±0.03	0.06±0.01	0.42±0.01	0.52±0.01	0.58±0.01
PO ₄ 3- (μM)	0.44±0.07	0.90±0.06	1.59±0.1	2.00±0.13	1.99±0.09
NO ₃ - (μM)	3.6±0.2	10.5±0.5	21.5±0.2	26.7±0.4	27.5±0.4
Si(OH) ₄ (μM)	2.6±0.1	2.5±1.8	6.6±0.1	40.3±0.5	45.0±0.8
NO ₂ - (μM)	0.15±0.02	0.13±0.02	0.17±0.02	0.19±0.01	0.21±0.02
Urea (µM)	0.23±0.04	0.11±0.04	0.26±0.08	0.24	0.21±0.03
chl-a (bulk) (µg L ⁻¹)	0.65±0.08	0.43±0.05	0.35±0.03	0.25±0.02	0.21±0.00
chl-a (nano+) (µg L ⁻¹)	0.50±0.05	0.30±0.04	0.24±0.02	0.18±0.02	0.17±0.02
chl-a (pico) (µg L⁻¹)	0.15±0.1	0.13±0.07	0.11±0.04	0.06±0.03	0.04±0.02
chl-a (% of nano+)	77.5±13.9	73.1±10.9	69.8±8.7	76.7±11.3	80.1±8.5
POC (bulk) (μM)	4.4±6.7	3.4±0.4	3.2±0.3	3.4±0.5	3.5+0.2
POC (nano+) (μM)	2.6±0.5	2.6±0.4	1.9±1.2	1.9±0.4	4.6
PON (bulk) (μM)	0.6±0.2	0.5±0.1	0.4±0.1	0.5±0.1	0.5±0.1
PON (nano+) (μM)	0.3±0.1	0.3±0.1	0.2±0.3	0.2±0.1	0.4±0.0
POC (% of nano+)	79.7±24.6	79.6±19.0	50.9±33.2	77.2±21.8	ND
PON (% of nano+)	69.0±31.9	67.1±17.2	53.8±24.1	67.0±21.9	51.1±24.7
POC:chl-a (g g ⁻¹)	103.0±22.1	102.5±14.4	122.5±11	234.1±29.2	219.3±1.0
POC:PON (M/M)	7.81±6.49	6.90±1.25	7.13±0.71	6.72±1.62	5.80±3.75
δ ¹⁵ N-PON	1.4±0.9	1.2±1.0	0.3±0.5	-1.3±0.5	-1.3±0.4
NPP (bulk) (nM day ⁻¹)	497.1±42.4	277.5±21.3	289.7±19.2	85.3±26.1	27.7±0.2
NPP (nano+) (nM day ⁻¹)	384.7±29.7	178.2±23.4	193.5	49.6±5.0	ND
ρΝΗ ₄ + (bulk) (nM day-1)	5.7±0.8	8.9±1.1	12.9±0.4	4.8±0.1	3.0±0.8
ρNH ₄ + (nano+) (nM day ⁻¹)	4.0±1.1	4.1±1.2	4.2±4.7	3.1±0.4	ND
ρNO ₃ - (bulk) (nM day-1)	4.1±0.4	11.5±1.4	5.9±1	3.6±0.4	3.7±1.8
ρNO ₃ - (nano+) (nM day-1)	3.4±0.3	6.6±0.4	4.3±0.4	2.6±0.8	2.7±1.2
ρUrea (bulk) (nM day ⁻¹)	7.5±0.6	6.9±0.3	6.5±1.0	2.1±0.3	0.6±0.01
ρUrea (nano+) (nM day ⁻¹)	4.9±0.3	3.8±0.2	4.0±0.6	1.3±0.2	0.7±0.4
f-ratio (bulk) (including ρUrea)	0.21±0.31	0.43±0.11	0.23±0.18	ND	0.51±0.53
f-ratio (bulk) (excluding ρUrea)	0.43±0.32	0.57±0.12	0.31±0.18	0.43±0.16	0.55±0.54
NH ₄ ⁺ ox (nM day ⁻¹)	9.3±0.5	12.9±0.6	11.1	17.7±0.6	14.3±1.0
Total microplankton (cells mL ⁻¹)	13±11	5±3	9±3	6±6	4±2
Centric diatoms (cells mL ⁻¹)	<1	<1	<1	<1	1±2
Pennate diatoms (cells mL ⁻¹)	2±4	<1	2±1	2±3	<1
Dinoflagellates (cells mL ⁻¹)	7±6	4±0	6±2	3±2	2±0
Micro-zooplankton (cells mL ⁻¹)	4±3	<1	2±2	1±2	<1
Nanoeukaryotes (cells mL ⁻¹)	ND	2.2±1.4 E+03	1.5±0.7 E+03	1.6±0.7 E+03	1.4E+03
Picoeukaryotes (cells mL ⁻¹)	ND	4.5±2.9 E+03	4.9±3.7 E+03	1.5±0.5 E+03	8E+02
Synechococcus (cells mL ⁻¹)	ND	3.8±1.8 E+03	2.3±1.1 E+03	1.4±0.2 E+03	1E+03

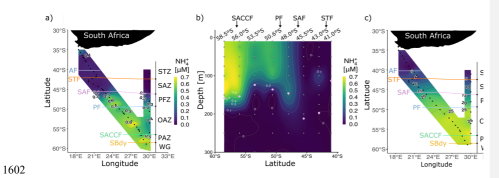


Figure 2: Concentrations of dissolved ammonium (NH $^+$) a) at the surface for Legs S and N and b) with depth (0-300 m) for Leg N, and c) concentrations of nitrate (NO $^-$) at the surface for Legs S and N. Pink circles in panel b show the mixed layer depth at the CTD stations. Abbreviations are as in Figure 1. Figure produced using the package ggplot2 (Wickham, 2016).

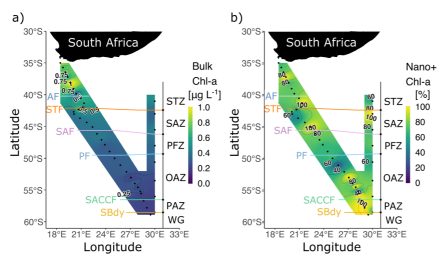
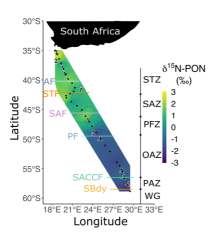


Figure 3: a) Bulk chlorophyll-a (chl-a) concentrations and b) the proportion of chlorophyll-a in the nano+size fraction at the surface for Legs S and N. Abbreviations are as in Figure 1. Figure produced using the package ggplot2 (Wickham, 2016).



1612 Figure 4: Bulk δ^{15} N-PON at the surface for Leg S in winter 2017. Two stations nearest South Africa at which biomass concentrations were extremely high have been excluded. Abbreviations are as in Figure 1614 1. Figure produced using the package ggplot2 (Wickham, 2016).

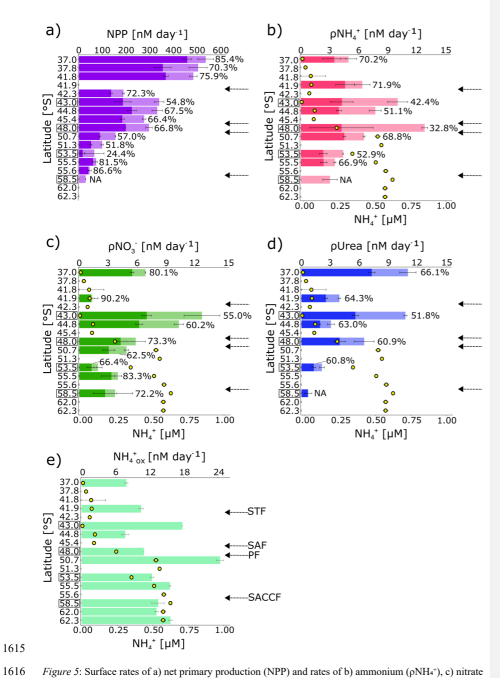


Figure 5: Surface rates of a) net primary production (NPP) and rates of b) ammonium (ρ NH₄⁺), c) nitrate (ρ NO₅⁻), and d) urea (ρ Urea) uptake by the pico (light colours) and nano+ (dark colours) size fractions, with the full length of the bars indicating the bulk rates, and e) NH₄⁺ oxidation. Error bars indicate ± 1

standard deviation of duplicate experiments. The percentage of total NPP and N uptake attributable to the nano+ size fraction is written next to each bar in panels a-d. NPP and NH₄+ uptake were not measured for the nano+ size fraction at 58.5°S, and urea uptake was not measured at 50.7°S and 55.5°S. Rates were not measured at the latitudes where no data are shown. In panels b-e, the surface NH₄+ concentration at each station is shown by the yellow circles. Leg N stations (at which samples were collected from Niskin bottles fired at 10 m) are indicated by black boxes surrounding the latitude. By contrast, samples were collected at the Leg S stations (no square surrounding the latitude) from the ship's underway system (~7 m). Fronts are indicated with arrows (labeled in panel e), and abbreviations are as in Figure 1. Figure produced using the package ggplot2 (Wickham, 2016).

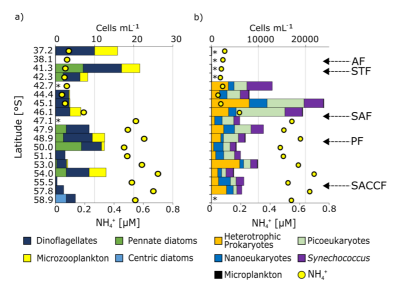


Figure 6: Surface community composition for a) plankton \geq 15 µm (enumerated by microscopy) and b) the total community <15 µm (enumerated by flow cytometry). For context, the surface NH₄+ concentration at each station is shown by the yellow circles. * indicates stations at which no measurements were made while the absence of a bar with no * indicates that no cells were detected. Note that the abundances shown on panel b (top x-axis) are >2 orders of magnitude greater than those shown in panel a. The "microplankton" shown in panel a are included on panel b (slim black bars) to illustrate the difference in abundance between the micro- and pico+nano populations. The frontal positions are indicated on panel b, with abbreviations as in Figure 1.

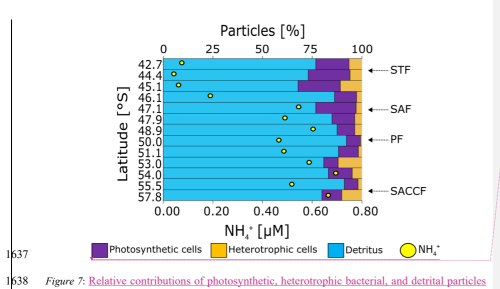
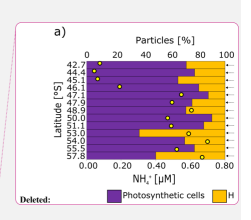
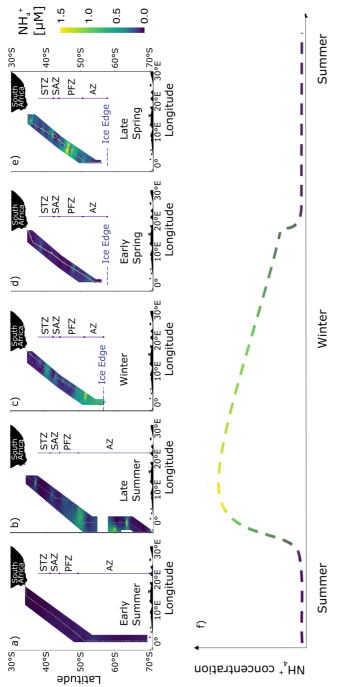


Figure 7: Relative contributions of photosynthetic, heterotrophic bacterial, and detrital particles to the total flow cytometry counts at the surface during leg S. The coincident NH_4^+ concentrations are shown as yellow dots. Abbreviations are as in Figure 1,



Deleted: Relative abundances of a) total photosynthetic versus heterotrophic bacteria and b) detritus versus heterotrophic bacteria at the surface for Leg S. The surface NH₄* concentration at each station is indicated by the yellow dots. The values in maroon text on the right side of panel a are the photosynthetic-to-heterotrophic cell ratios. The upper x-axis in panel b begins at 75% in order to highlight the (much smaller) heterotrophic bacterial contribution to the summed detrital + heterotrophic particles. Frontal abbreviations are as in Figure 1.



gradient in panel f shows the transition between late summer and late winter. Panels a and b cover a latitudinal extent of 30-70°S, while panels c-e cover 30-60°S due to the presence of sea-ice. Abbreviations are as in Figure 1, with AZ referring to the combined OAZ and PAZ. Figure produced using the package ggplot2 (Wickham, 2016). Figure 8: Surface concentrations of NH⁺ across the eastern Atlantic sector of the Southern Ocean measured between December 2018 and November 2019. Five unique transects (additional to the winter 2017 dataset presented in Fig. 2a) are shown: a) early summer 2018, b) late summer 2019, c) winter 2019, d) early spring 2019, and e) late spring 2019. f) The proposed seasonal cycle of NH4* concentrations in the mixed layer south of the Subantarctic Front. The colour

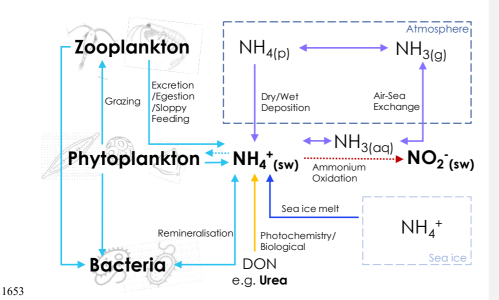


Figure 9: Schematic of the possible mixed-layer NH_4^+ assimilation and production pathways. Bold text indicates components of the NH_4^+ cycle that were directly measured in this study (seawater concentrations of NH_4^+ , NO_2^- , and urea; phytoplankton, bacterial, and microzooplankton cell abundances), and dotted lines indicate processes for which we have direct rate measurements (phytoplankton uptake of NH_4^+ ; oxidation of NH_4^+ to NO_2^-). Dashed-line boxes represent the atmosphere and sea-ice, with all other processes occurring in the ocean. DON – dissolved organic nitrogen; $NH_{3(aq)}$ – aqueous (seawater) ammonia; $NH_{4(p)}$ – ammonium aerosols (including ammonium sulphate, ammonium bisulphate, and ammonium nitrate); $NH_{3(g)}$ – ammonia gas.