# The impact of snow nitrate photolysis on boundary layer chemistry and the recycling and redistribution of reactive nitrogen across Antarctica and Greenland in a global chemical transport model

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#### Abstract

The formation and recycling of reactive nitrogen (NO, NO<sub>2</sub>, HONO) at the air-snow interface has implications for air quality and the oxidation capacity of the atmosphere in snow-covered regions. Nitrate  $(NO_3^{-})$  photolysis in snow provides a source of oxidants (e.g., hydroxyl radical) and oxidant precursors (e.g., nitrogen oxides) to the overlying boundary layer, and alters the concentration and isotopic (e.g.,  $\delta^{15}N$ ) signature of NO<sub>3</sub><sup>-</sup> preserved in ice cores. We have incorporated an idealized snowpack with a NO3 photolysis parameterization into a global chemical transport model (GEOS-Chem) to examine the implications of snow NO<sub>3</sub><sup>-</sup> photolysis for boundary layer chemistry, the recycling and redistribution of reactive nitrogen, and the preservation of ice-core NO<sub>3</sub><sup>-</sup> in ice cores across Antarctica and Greenland, where observations of these parameters over large spatial scales are difficult to obtain. A major goal of this study is to examine the influence of meteorological parameters and chemical, optical, and physical snow properties on the magnitudes and spatial patterns of snow-sourced NO<sub>x</sub> fluxes and the recycling and redistribution of reactive nitrogen across Antarctica and Greenland. Snowsourced NO<sub>x</sub> fluxes are most influenced by temperature-dependent quantum yields of NO<sub>3</sub> photolysis, photolabile NO<sub>3</sub> concentrations in snow, and concentrations of lightabsorbing impurities (LAI) in snow. Despite very different assumptions about snowpack properties, the range of model-calculated snow-sourced NO<sub>x</sub> fluxes are similar in Greenland (0.5 to  $11 \times 10^8$  molec cm<sup>-2</sup> s<sup>-1</sup>) and Antarctica (0.01 to  $6.4 \times 10^8$  molec cm<sup>-2</sup> s<sup>-1</sup>) due to the opposing effects of higher concentrations of both photolabile  $NO_3^-$  and LAI in Greenland compared to Antarctica. Despite the similarity in snow-sourced NO<sub>x</sub> fluxes, these fluxes lead to smaller factor increases in mean austral summer boundary layer mixing ratios of total nitrate (HNO<sub>3</sub>+ NO<sub>3</sub><sup>-</sup>), NO<sub>x</sub>, OH, and O<sub>3</sub> in Greenland compared to Antarctica because of Greenland's proximity to pollution sources. The degree of nitrogen recycling in the snow is dependent on the relative magnitudes of snow-sourced NO<sub>x</sub> fluxes versus primary  $NO_3^-$  deposition. Recycling of snow nitrate in Greenland is much less than in Antarctica because the deposition of primary  $NO_3^-$  is up to 35 times larger than snow-sourced  $NO_x$  fluxes in Greenland. Photolysis-driven loss of snow  $NO_3^-$  is largely dependent on the time that NO<sub>3</sub><sup>-</sup> remains in the snow photic zone (up to 6.5 years in Antarctica and 7 months in Greenland), and wind patterns that redistribute snowsourced reactive nitrogen across Antarctica and Greenland. The loss of snow NO<sub>3</sub><sup>-</sup> is higher in Antarctica (up to 99%) than in Greenland (up to 83%) due to deeper snow photic zones and lower snow accumulation rates in Antarctica. Modeled enrichments in ice-core  $\delta^{15}N(NO_3)$  due to photolysis-driven loss of snow NO<sub>3</sub> ranges from 0 to 363‰ in Antarctica and 0 to 90% in Greenland, with the highest fraction of NO<sub>3</sub><sup>-</sup> loss and largest enrichments in ice-core  $\delta^{15}N(NO_3)$  at high elevations where snow accumulation rates are lowest. There is a strong relationship between the degree of photolysis-driven loss of snow  $NO_3^-$  and the degree of nitrogen recycling between the air and snow throughout all of Greenland and in Antarctica where snow accumulation rates are greater than 130 kg m<sup>-2</sup> a<sup>-1</sup> in the present day.

1 1. Introduction

2 Nitrogen oxides ( $NO_x = NO + NO_2$ ) emitted from fossil fuel combustion, biomass burning, 3 soil microbial activity, and lightning have adverse respiratory effects, contribute to the 4 formation of atmospheric acidity, and are a key ingredient in tropospheric oxidant cycling 5 leading to the formation of ground-level ozone (O<sub>3</sub>). O<sub>3</sub> also has adverse respiratory effects, is an effective greenhouse gas [UNEP, 2011], and its photolysis dominates 6 7 hydroxyl radical (OH) production in much of the troposphere [Thompson, 1992]. 8 Oxidation to form nitrate (HNO<sub>3</sub>/NO<sub>3</sub><sup>-</sup>) is the main sink for NO<sub>x</sub> in the troposphere 9 [Logan, 1983], and the lifetime of  $NO_x$  against oxidation to nitrate is expected to be 1-3 days in polar regions [Levy et al., 1999]. NO<sub>3</sub><sup>-</sup> is lost from the atmosphere through dry 10 and wet deposition to the Earth's surface, and has a global atmospheric lifetime of 11 12 roughly 5 days [Xu and Penner, 2012]. In Antarctica and Greenland, HNO<sub>3</sub> and NO<sub>3</sub><sup>-</sup> 13 deposited to the snowpack originates from both the troposphere (e.g., long-range 14 transport) [Geng et al., 2014a, Lee et al., 2014, Wespes et al., 2012] and stratosphere 15 [Davidson et al., 1989, Frev et al., 2009, Savarino et al., 2007]. In snow-covered regions, 16 the deposition of HNO<sub>3</sub> and NO<sub>3</sub><sup>-</sup> is not a permanent sink for NO<sub>x</sub>, as the photolysis of 17 snow  $NO_3^-$  returns reactive nitrogen (N<sub>r</sub>=NO<sub>x</sub>, HONO) back to the atmosphere, with 18 implications for other oxidants such as OH and O<sub>3</sub> [Domine and Shepson, 2002].

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20 Snow photochemistry significantly influences boundary layer chemistry and plays an 21 important role in oxidant production and cycling, especially in pristine regions, such as 22 Antarctica [Bloss et al., 2007, Chen et al., 2004, 2007, Grannas et al., 2007, Helmig et al., 23 2008, Sjostedt et al., 2007, Thomas et al., 2012]. Snow photochemistry may have more 24 widespread impacts since up to 40% of land on Earth is snow-covered at a given time 25 [*Grannas et al.*, 2007].  $NO_3^{-1}$  is not the only photochemically-active species in snow. The 26 photolysis of nitrite (NO<sub>2</sub>) in snow and the photolysis of snow-sourced formaldehyde 27  $(CH_2O)$ , nitrous acid (HONO), and hydrogen peroxide  $(H_2O_2)$  provide additional sources 28 of  $N_r$  and OH to the boundary layer. Bromine (Br<sub>2</sub>) is also produced in the snow via 29 reactions involving bromide (Br), photochemically-active species (e.g., NO<sub>3</sub>), and 30 photochemically-produced species (e.g., OH) within snow grains [Pratt et al., 2013].

31 32 In snow, NO<sub>3</sub><sup>-</sup> photolysis likely occurs in the liquid-like region (LLR) on the surface of 33 ice grains, in cracks between ice grains, or in brine pockets embedded within ice grains 34 [*Domine et al.*, 2013]. There are two channels for NO<sub>3</sub><sup>-</sup> photolysis at wavelengths 35 ( $\lambda$ )=290-345 nm. In the aqueous phase, NO<sub>3</sub><sup>-</sup> can photolyze to produce NO<sub>2</sub> and OH (E1), 36 or produce NO<sub>2</sub><sup>-</sup> and O(<sup>3</sup>P) (E2), but E1 is thought to be the dominant pathway [*Grannas* 37 *et al.*, 2007, *Mack and Bolton*, 1999, *Meusinger et al.*, 2014].

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The aqueous phase NO<sub>2</sub> produced in E1 can evaporate to the gas phase and be released into the interstitial air [*Boxe et al.*, 2005] and subsequently be transported to the overlying atmosphere via diffusion and windpumping [*Zatko et al.*, 2013]. The quantum yield ( $\phi$ ) in E1 is strongly influenced by the location of NO<sub>3</sub><sup>-</sup> in an ice grain. Quantum yields are 46 more weakly dependent on LLR temperature and pH;  $\phi$  values increase with increasing 47 temperature and pH [Chu and Anastasio, 2003]. Chu and Anastasio [2003] froze NO<sub>3</sub>-48 doped water in the lab and measured the quantum yield for E1 (0.003 molec photon<sup>-1</sup> at 49 T=253 K) as frozen ice grains were exposed to ultraviolet (UV) radiation. Zhu et al. 50 [2010] deposited HNO<sub>3</sub> on an ice film and measured  $\phi$  for E1 (0.6 molec photon<sup>-1</sup> at 51 T=253 K), as the frozen surface was irradiated with UV radiation. A recent study by 52 Meusinger et al. [2014] found  $\phi$ =0.003-0.44 molec photon<sup>-1</sup> at T=258 K for E1, which 53 nearly spans the full range of previously reported quantum yields. Results from 54 Meusinger et al. [2014] suggest that  $\phi$  is dependent on the length of time that snow is 55 exposed to UV radiation, as well as the location of  $NO_3^-$  in the ice grain. Meusinger et al. 56 [2014] suggest that two photochemical domains of  $NO_3^-$  exist: photolabile  $NO_3^-$  and  $NO_3^-$ 57 buried within the ice grain. The NO<sub>x</sub> produced from the photolysis of photolabile NO<sub>3</sub><sup>-</sup> 58 can escape the ice grain, while the  $NO_x$  produced from the photolysis of buried  $NO_3^-$  is 59 likely to undergo recombination chemistry within the snow grain, thus lowering the 60 quantum yield of  $NO_x$  for  $NO_3^-$  photolysis. Recombination chemistry involves  $NO_3^$ reformation from photo-products (e.g., NO<sub>x</sub> and OH) within the ice grain, which alters 61 oxygen isotopes (e.g.,  $\Delta^{17}$ O) of NO<sub>3</sub>, but does not impact bulk snow NO<sub>3</sub> concentrations 62 nor its nitrogen isotopes ( $\delta^{15}$ N) [Frev et al., 2009, Erbland et al., 2013, 2015]. 63 64

65 The NO<sub>2</sub><sup>-</sup> produced in E2 is quickly photolyzed at longer wavelengths ( $\lambda$ =290-390 nm) in 66 the LLR or can react with OH or H<sup>+</sup> in the LLR to produce N<sub>r</sub> [*Grannas et al.*, 2007]:

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 $\begin{array}{ll} 68 & \operatorname{NO}_2(\operatorname{aq}) + \operatorname{hv}(+\operatorname{H}^+, \operatorname{aq}) \xrightarrow{\bullet} \operatorname{NO}(\operatorname{aq}) + \operatorname{OH}(\operatorname{aq}), \\ 69 & \operatorname{NO}_2(\operatorname{aq}) + \operatorname{OH}(\operatorname{aq}) \xrightarrow{\bullet} \operatorname{NO}_2(\operatorname{aq}) + \operatorname{OH}(\operatorname{aq}), \\ 70 & \operatorname{NO}_2(\operatorname{aq}) + \operatorname{H}^+(\operatorname{aq}) \xrightarrow{\bullet} \operatorname{HONO}(\operatorname{aq}), \\ \end{array}$   $\begin{array}{ll} E3 \\ E4 \\ E5 \\ E5 \end{array}$ 

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72 HONO produced in E5 can rapidly photolyze to produce NO and OH in the interstitial air 73 or the atmospheric boundary layer [Anastasio and Chu, 2009]. Reactions involving NO<sub>2</sub><sup>-</sup> 74 are intermediate reactions for NO<sub>3</sub><sup>-</sup> photolysis because NO<sub>3</sub><sup>-</sup> photolysis is required for 75 NO<sub>2</sub> formation and the end products of E1-E5 are all N<sub>r</sub>. Once produced, NO<sub>2</sub> and NO 76 can be efficiently transported to the overlying atmosphere via windpumping [Zatko et al., 77 2013] and enter into rapid NO<sub>x</sub>-cycling reactions. In the atmosphere, the relative 78 abundance of NO and NO<sub>2</sub> will be determined by local atmospheric conditions, 79 specifically oxidant concentrations (e.g., O<sub>3</sub>, HO<sub>2</sub>, RO<sub>2</sub>, BrO, and ClO) [Frey et al., 80 2013]. The snow-sourced NOx can be re-oxidized to HNO3 via E6 under sunlit 81 conditions.

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$$NO_2(g) + OH(g) \rightarrow HNO_3(g),$$
 E6

The HNO<sub>3</sub> produced in E6 can undergo wet or dry deposition to the snow surface [*Dibb et al.*, 2004] within a day [*Slusher et al.*, 2002, *Wang et al.*, 2008]. Evidence for HNO<sub>3</sub> re-deposition is seen in the snow NO<sub>3</sub><sup>-</sup> concentration profile at many polar locations, where NO<sub>3</sub><sup>-</sup> concentrations may be over an order of magnitude higher in the top two centimeters (cm) of snow compared to NO<sub>3</sub><sup>-</sup> concentrations below [*Dibb et al.*, 2004, *Frey et al.*, 2009, *Mayewski and Legrand*, 1990, *Rothlisberger et al.*, 2000].

92 Once HNO<sub>3</sub> is deposited back to the snow, it is available for photolysis again. NO<sub>3</sub><sup>-</sup> can 93 be recycled multiple times between the boundary layer and the snow before burial below 94 the photochemically-active region, known as the snow photic zone [Davis et al., 2008, 95 Erbland et al., 2015]. Only two previous studies have attempted to quantify the degree of 96 nitrogen recycling between the air and snow. Davis et al. [2008] use estimates of 97 atmospheric NO<sub>x</sub> overhead-column burdens and average atmospheric NO<sub>x</sub> lifetimes 98 along with primary nitrogen deposition measurements from Legrand and Kirchner [1990] 99 to estimate that nitrogen is recycled 1.8 times on average between the air and snow in one 100 photochemical season in East Antarctica, although this value may be 3 to 5 times higher 101 due to uncertainties in primary nitrogen deposition estimates. Erbland et al. [2015] use a 102 multi-layer, one-dimensional snow model (TRANSITS) and calculate that nitrogen is 103 recycled 4 times on average before burial beneath the snow-photic zone. Erbland et al. 104 [2015] observe an inverse relationship between snow accumulation rate and nitrogen recycling in regions where snow accumulation rates are greater than 50 kg m<sup>-2</sup>  $a^{-1}$ , and 105 suggest that in these regions, the degree of  $NO_3^-$  recycling is governed by the time  $NO_3^-$ 106 107 remains in the snow photic zone.

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109 The photolysis of snow  $NO_3^-$  and subsequent recycling between the air and snow alters the concentration and isotopic (e.g.,  $\delta^{15}N$ ) signature of NO<sub>3</sub><sup>-</sup> that is ultimately preserved 110 111 in polar ice sheets, which hampers the interpretation of ice-core NO<sub>3</sub> records [Wolff et 112 al., 2008]. Such records have been sought to reconstruct the past history of the abundance 113 of NO<sub>x</sub> in the atmosphere [Wolff, 1995]. It has also been suggested that the nitrogen  $(\delta^{15}N)$  and oxygen  $(\Delta^{17}O)$  isotopic composition of ice-core NO<sub>3</sub> can provide information 114 on past variability in atmospheric NO<sub>x</sub> sources and oxidant abundances [e.g., Alexander 115 et al., 2004, Hastings et al., 2005]. Different sources of NO<sub>x</sub> have different  $\delta^{15}$ N 116 117 signatures (~ -19‰ to 25‰, [Geng et al., 2014a, Walters et al., 2015]), giving ice-core  $\delta^{15}N(NO_3)$  measurements the potential to track NO<sub>x</sub>-source changes over time. 118

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120 Ice-core  $\delta^{15}N(NO_3^{-})$  values will be altered if there is photolysis-driven loss of NO<sub>3</sub><sup>-</sup> from the snow when snow-sourced  $NO_x$  is transported away from the site of primary 121 122 deposition. Nitrate photolysis in snow is associated with a large fractionation constant ( $\varepsilon$ , e.g., -47.9% [Berhanu et al., 2014]), providing the boundary layer with a source of NO<sub>x</sub> 123 that is highly depleted in  $\delta^{15}N(NO_3^-)$ , leaving highly enriched  $\delta^{15}N(NO_3^-)$  in the snow. In 124 Antarctica, atmospheric  $\delta^{15}N(NO_3)$  values at the coast are as low as -40‰, indicating 125 126 transport of snow-sourced NO<sub>x</sub> from the continental interior [Morin et al., 2009], while on the East Antarctica plateau, snow  $\delta^{15}N(NO_3)$  up to 480% has been reported [Blunier 127 et al., 2005, Erbland et al., 2013, Frey et al., 2009, Shi et al., 2014], indicating net loss of 128 129 NO<sub>3</sub><sup>-</sup> driven by photolysis. In Greenland, atmospheric  $\delta^{15}N(NO_3)$  values are much less depleted (as low as -15‰) and snow  $\delta^{15}N(NO_3)$  values are much less enriched (as high 130 131 as 15‰) compared to these extreme values observed in coastal Antarctica and on the East 132 Antarctic plateau [Geng et al., 2014a, Hastings et al., 2004, Jarvis et al., 2009].

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134 If snow-sourced NO<sub>x</sub> is simply re-deposited back to the snow surface at the site of 135 emission, a vertical profile in  $\delta^{15}$ N(NO<sub>3</sub><sup>-</sup>) within the snow photic zone will develop due to 136 vertical redistribution of NO<sub>3</sub><sup>-</sup> [*Erbland et al.*, 2013, *Frey et al.*, 2009]; however, the 137 depth-integrated  $\delta^{15}$ N(NO<sub>3</sub><sup>-</sup>) will not be impacted, even with active photolysis-driven

recycling between the atmosphere and the snow. Enrichment in ice-core  $\delta^{15}N(NO_3^{-1})$ 138 139 requires photolysis-driven loss from snow from net atmospheric transport of snowsourced NO<sub>x</sub> away from the locations of its production. In addition to photolysis, ice-core 140 141  $\delta^{15}N(NO_3)$  values are also influenced by evaporation of HNO<sub>3</sub> [*Mulvaney et al.*, 1998] 142 from snow and by atmospheric processing, such as NO<sub>x</sub> cycling [Frever et al., 1993] and 143 gas-particle partitioning [Heaton et al., 1997, Geng et al., 2014a]; however, these impose 144 a fractionation in  $\delta^{15}N(NO_3)$  at least an order of magnitude smaller than photolysis, and are thus not able to explain the large enrichments in snow  $\delta^{15}N(NO_3)$  observed on the 145 East Antarctic plateau [Blunier et al., 2005, Erbland et al., 2013, Frev et al., 2009, Shi et 146 147 al., 2014].

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149 Here we incorporate an idealized snowpack with a NO<sub>3</sub><sup>-</sup> photolysis parameterization into a global chemical transport model. The idealized Antarctic and Greenland ice sheets in 150 151 the model have similar properties as the real ice sheets, but are subject to assumptions 152 about the chemical and physical properties of the snow. The idealized snowpacks in this 153 modeling framework attempt to account for the spatial variability in parameters important 154 to snow NO<sub>3</sub><sup>-</sup> photolysis in order to investigate the potential spatial variability in snow-155 sourced NO<sub>x</sub> fluxes, reactive nitrogen recycling and redistribution, and preservation of 156 ice-core NO<sub>3</sub><sup>-</sup> across Antarctica and Greenland, where observations of these parameters 157 over large spatial scales are difficult to obtain. A major advantage of using a global 158 chemical transport model framework is the ability to examine the potential redistribution 159 and loss of reactive nitrogen due to photolysis-driven loss of snow NO<sub>3</sub><sup>-</sup> across large 160 spatial scales. The sensitivity of many parameters, such as snow accumulation rate, on the 161 flux of snow-sourced NO<sub>x</sub>, nitrogen recycling, and loss of snow NO<sub>3</sub><sup>-</sup> is elucidated by comparing modeled results in Antarctica and Greenland. Section 2 describes the inclusion 162 163 of an idealized snowpack with a snow NO<sub>3</sub><sup>-</sup> photolysis parameterization into a global chemical transport model, GEOS-Chem. Section 3 explores the implications of 164 165 photolysis-driven reactive nitrogen recycling and redistribution for boundary layer 166 chemistry and the alteration of NO<sub>3</sub><sup>-</sup> concentration and its nitrogen isotopes ( $\delta^{15}N$ ) 167 ultimately archived in ice cores. In section 3, we also compare model results in Antarctica 168 and Greenland to examine the sensitivity of the flux of snow-sourced  $NO_x$  and associated 169 photolysis-driven processes on meteorological parameters and various chemical, optical, 170 and physical properties of snow. We end section 3 by discussing results from our model 171 sensitivity studies that highlight the largest uncertainties in our ability to model these 172 processes. Section 4 summarizes our conclusions and provides recommendations for 173 future laboratory and field studies based on our model sensitivity results. 174

### 175 **2. Methods**

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## 177 2.1. Incorporating Snow NO<sub>3</sub><sup>-</sup> Photolysis into a Global Chemical Transport Model

Table 1 provides a glossary of the variables used throughout this paper.

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# 179180 2.1.1. Global Chemical Transport Model Description

181 GEOS-Chem is a global 3-dimensional (3-D) model of coupled aerosol-oxidant 182 chemistry with detailed  $HO_x$ -NO<sub>x</sub>-VOC-O<sub>3</sub>-BrO<sub>x</sub> tropospheric chemistry originally 183 described in Bey et al. [2001]. The model uses assimilated meteorological data from the

NASA Goddard Earth Observing System (GEOS-5) including winds, convective mass 184 185 fluxes, boundary layer depths, temperature, precipitation, and surface properties. 186 Meteorological data have 6-hour temporal resolution (3-hour for surface variables and 187 mixing depths). The TPCORE advection algorithm [Lin and Rood, 1996] is the transport routine in GEOS-Chem and is based on the calculation of the slopes between neighboring 188 189 grid boxes. At the poles, neighboring grid boxes are used to estimate transport of 190 chemical species into and out of the circular polar grid box. In Figures 2-11, data in each 191 grid box are smoothed using bilinear interpolation. The spectral direct and diffuse downwelling surface irradiance and photolysis frequencies are calculated using the Fast-192 193 JX radiative transfer module [Bian and Prather, 2002, Mao et al., 2010, Wild et al., 194 2000]. In GEOS-Chem, aerosols can be wet deposited via scavenging in convective 195 updrafts and by rainout from convective anvils and large-scale precipitation [Liu et al., 2001]. The wet deposition scheme for gases is described by Amos et al. [2012] and the 196 197 scavenging of aerosol by snow and cold/mixed precipitation is described by Wang et al. 198 [2011]. Dry-deposition velocities for coarse mode aerosols (radii between 1-10 mm) are 199 calculated based on aerosol size and hydroscopic growth as described in Zhang et al. 200 [2001]. Aerosol deposition to snow and ice surfaces is described by Fisher et al. [2011]. 201 For smaller aerosols (radii less than 1 µm), dry deposition velocities are calculated with a 202 standard resistance-in-series scheme [Wang et al., 1998, Wesely, 1989].

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204 Anthropogenic NO<sub>x</sub> emissions are from the EDGAR 3.2-FT2000 global inventory for the 205 year 2000 [Oliver et al., 2005], scaled by country on the basis of energy statistics as 206 described by van Donkelaar et al. [2008]. The monthly inventory of emissions from 207 biomass burning are from the Global Fire Emissions Database (GFED2) [van der Werf et 208 al., 2009]. Soil NO<sub>x</sub> emissions are computed using a parameterization described in Hudman et al. [2012], which is a function of vegetation type, temperature, soil moisture, 209 210 precipitation, and fertilizer emissions. Emissions of NO<sub>x</sub> from lightning are linked to deep convection following the parameterization of Price and Rind [1992] and are scaled 211 212 globally as described by Murray et al. [2012] to match OTD/LIS climatological 213 observations of lightning flashes. The stratospheric source of NO<sub>v</sub> (=NO<sub>v</sub> +HNO<sub>3</sub>) utilizes 214 monthly climatological 3-D production and loss rates from the Global Modeling Initiative 215 (GMI) model [Allen et al., 2010], which captures the formation of the polar vortex and 216 PSC sedimentation [Murray et al., 2012].

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218 For this work, GEOS-Chem version v9-01-01 was run at 2°x2.5° horizontal resolution 219 with 72 hybrid vertical levels using GEOS-5 meteorology from May 2009 to May 2010. 220 The model was spun up for six months prior to May 2009. There are no sub-surface 221 (snow) layers in GEOS-Chem and the three lowest vertical levels are each roughly 100 222 meters in height above Antarctica. The boundary layer in GEOS-Chem is calculated 223 using a parameterization involving the bulk Richardson number with surface friction, a 224 turbulent velocity scale, and non-local fluxes of heat and moisture [Holtslag and Boville, 1993] as implemented by Lin and McElroy [2010]. The mixing of emissions, dry 225 226 deposition, and concentrations of individual species within the boundary layer are 227 determined by static instability. In a stable boundary layer, the local scheme based on 228 eddy diffusivity-theory is used, and the mixing is weak. In an unstable boundary layer, 229 boundary layer mixing is triggered by large eddies. Average boundary layer mixing ratios

- 230 (ppbv) of species reported in this study (e.g.,  $NO_3^-$ ,  $NO_x$ , OH,  $O_3$ ) are mixing ratios in the 231 lowest vertical grid box (total height ~ 100 m).
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Figure 1 illustrates the nitrogen recycling associated with snow NO<sub>3</sub><sup>-</sup> photolysis as included in the model. The total flux of snow-sourced NO<sub>x</sub> from the snow,  $F_{NOx}$  (molec cm<sup>-2</sup> s<sup>-1</sup>), is calculated using the wavelength-dependent absorption cross-section for NO<sub>3</sub><sup>-</sup> photolysis ( $\sigma_{NO3}$ , cm<sup>2</sup> molec<sup>-1</sup>), the temperature (T)- and pH-dependent quantum yield for NO<sub>3</sub><sup>-</sup> photolysis ( $\phi$ , molec photon<sup>-1</sup>), the depth- and wavelength-dependent actinic flux in the snow photic zone (*I*, photons cm<sup>-2</sup> s<sup>-1</sup> nm<sup>-1</sup>), and the average NO<sub>3</sub><sup>-</sup> concentration ([NO<sub>3</sub><sup>-</sup>], molec cm<sup>-3</sup>) integrated over the depth of the photic zone.  $F_{NOx}$  (molec cm<sup>-2</sup> s<sup>-1</sup>), is calculated in E7 and converted into units of ng N m<sup>-2</sup> yr<sup>-1</sup> in E9 and E10.

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$$F_{NOx} = \int_{\lambda_0}^{\lambda_1} \int_{z_0}^{z_{3e}} J(\lambda, z) \cdot [NO_3^-] d\lambda dz,$$
243 E7

where  $[NO_3^-]$  is the average photolabile NO<sub>3</sub><sup>-</sup> concentration over the depth of the snow photic zone ( $z=z_0$  to  $z=z_{3e}$ ) and J is the photolysis rate constant (s<sup>-1</sup>), which is calculated in E8:

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 $J = \sigma_{NO_3^-}(\lambda) \cdot \phi(T, pH) \cdot I(\lambda, z)$ E8

250 In E8,  $\sigma_{NO3}$  is from Sander et al. [2006]. The temperature-dependent  $\phi$  from Chu and 251 Anastasio [2003] (assumed constant pH of 5) is calculated using the modeled air temperature in the lowest vertical grid box. The actinic flux (1) is calculated at 1-cm 252 253 intervals and integrated from the snow surface  $(z_0)$  to the depth of the photic zone  $(z_{3e})$ . 254 The snow photic zone is defined as three times the e-folding depth of ultraviolet (UV) 255 actinic flux in snow  $(z_{3e})$ , where 1 e-folding depth is  $z_e$ . Below  $z_{3e}$ , more than 95% of the 256 radiation has been attenuated and minimal photochemistry occurs. The flux of snow-257 sourced NO<sub>x</sub> is integrated over several ultraviolet wavelength bands (298-307 nm, 307-258 312 nm, 312-320 nm, 320-345 nm), which are then summed to calculate total  $F_{NOx}$  from 259 the photolysis of snow NO<sub>3</sub><sup>-</sup> between  $\lambda$ =298-345 nm. We assume that all NO<sub>x</sub> formed in 260 E7 is immediately evaporated and transported into the overlying boundary layer via wind 261 pumping and diffusion [Zatko et al., 2013].

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#### 263 2.1.2 Calculating Radiative Transfer in Snow

264 A 2-stream, plane parallel snowpack actinic flux parameterization based on a 4-stream 265 radiative transfer model [Grenfell, 1991] was developed and described in detail in Zatko 266 et al. [2013] and has been implemented into GEOS-Chem for the purposes of this study. 267 The parameterization is simple, broadly applicable, and allows for variation in snow and sky properties (e.g., solar zenith angle, cloud fraction) over time. Ice grains are assumed 268 269 to be spherical in shape and light-absorbing impurities (LAI), including black carbon, 270 brown carbon, dust, and organics, are assumed to be homogeneously distributed 271 throughout the snow and always external to the ice grain. The snowpack actinic flux parameterization is used to calculate the UV actinic flux (photons  $\text{cm}^{-2} \text{ s}^{-1} \text{ nm}^{-1}$ ) and the 272 mean summer e-folding depths (cm) across Antarctica and Greenland,, which are used to 273 calculate  $F_{NOx}$ . The snowpack actinic flux parameterization is most sensitive to radiation 274 275 equivalent mean ice grain radii  $(r_e)$  and insoluble LAI in snow [Zatko et al., 2013]; higher 276 concentrations of LAI in the snow and smaller  $r_e$  lead to shallower e-folding depths ( $z_e$ ). 277 Vertical  $r_e$  and snow density ( $\rho_{snow}$ ) profiles at Dome C, Antarctica from Gallet et al. 278 [2011] are used across Antarctica for all seasons, which range from 86 to 235 µm and 260 to 360 kg m<sup>-3</sup>, from the snow surface to 300-cm depth, respectively. Vertical  $r_e$  and 280  $\rho_{snow}$  profiles at Summit, Greenland from Carmagnola et al. [2013] are used in Greenland, 281 ranging from 73 to 211 µm and 235 to 350 kg m<sup>-3</sup>, from the snow surface to 300-cm 282 depth, respectively.

283

284 The concentration of black carbon (BC) concentrations in snow ( $C_{BC}$ ) are calculated using modeled total annual black carbon (hydrophilic + hydrophobic) deposition (ng BC 285  $m^{-2} yr^{-1}$ ) and total annual snow accumulation rates (g H<sub>2</sub>O m<sup>-2</sup> yr<sup>-1</sup>) in GEOS-Chem. At 286 some locations in coastal Antarctica, high accumulation rates (up to 700 kg m<sup>-2</sup> yr<sup>-1</sup>) lead 287 to unrealistically low  $C_{BC}$  (as low as 0.01 ng g<sup>-1</sup>), therefore the minimum  $C_{BC}$  value used 288 in the model is 0.08 ng g<sup>-1</sup>, which is comparable to the  $C_{BC}$  values measured in high snow 289 accumulation rate regions, such as in the East Antarctic sea ice zone (0.1 ng  $g^{-1}$ ) [Zatko 290 291 and Warren, 2015]. Insoluble non-black carbon species (nonBC) including dust, brown 292 carbon, and organics, are responsible for the majority (up to 89% at  $\lambda$ =305 nm) of the 293 absorption of radiation at UV wavelengths [Zatko et al., 2013] in snow. These nonBC 294 species and their concentrations have not been well quantified in snow. Based on 295 observations reported in Zatko et al. [2013], we scale UV-absorption by insoluble nonBC 296 to the absorption by insoluble black carbon in snow by assuming that at  $\lambda$ =650-700 nm, 297 which is a wavelength range where black carbon dominates absorption, insoluble black 298 carbon is responsible for 70% of the particulate absorption. We also assume that nonBC 299 material has an absorption Ångstrom exponent of 5 [Doherty et al., 2010].

300

301 We neglect the influence of soluble LAI in the snow and only consider the influence of 302 insoluble LAI for calculations of actinic flux profiles in snow. To determine whether 303 soluble LAI contribute significantly to light-absorption in the snow, we calculate the total 304 extinction coefficient for insoluble BC, insoluble nonBC, and soluble LAI following 305 section 2.1 of Zatko et al. [2013] and using the absorption coefficients for soluble material in snow reported in Beine et al., [2011] in northern Alaska. To our knowledge, 306 307 observations of soluble light-absorbing impurities in Antarctic and Greenland snow are 308 unavailable. We use soluble LAI observations from Alaska to provide a relative estimate of the importance of soluble LAI in polar snow. The absorption coefficients (0.028  $m^{-1}$  at 309 310  $\lambda$ =307 nm) from Beine et al. [2011] are identical to the extinction coefficients because it is assumed that there is no scattering by soluble species. Insoluble  $C_{BC}$  (9 ng g<sup>-1</sup>) from 311 Barrow, Alaska [Doherty et al., 2010] were used to calculate extinction coefficients for 312 313 insoluble BC and nonBC material and therefore the amount of nonBC absorption in the 314 UV and near-visible wavelengths following Zatko et al. [2013]. Insoluble nonBC 315 material is responsible for 9-14 times more absorption than soluble material in the 316 wavelength range  $\lambda$ =298-345 nm. Insoluble BC material is responsible for 1.5-10 times 317 more absorption than soluble material in the wavelength range  $\lambda$ =298-345 nm. The 318 extinction coefficient is not influenced by the addition of a soluble absorber because 319 scattering by snow grains dominates the extinction in snow. The effective co-albedo of 320 single scattering is increased by 6-15% when soluble absorbers are included. The resulting change in  $z_e$  is at most 0.5 cm, which represents an increase of 4-9% in the 321

#### 322 wavelength region of $\lambda$ =298-345 nm.

#### 323 2.1.3. Calculating NO<sub>3</sub><sup>-</sup> Concentrations in Snow

The median value of sub-surface snow NO<sub>3</sub><sup>-</sup> concentrations is 60 ng g<sup>-1</sup> in Antarctica 324 [Bertler et al., 2005] and 132 ng g<sup>-1</sup> in Greenland [Burkhart et al., 2009]. Snow NO<sub>3</sub><sup>-</sup> was 325 326 collected over depths corresponding to between 1 and 70 year(s) of snow accumulation in 327 Antarctica and between 1 and 148 year(s) of snow accumulation in Greenland. The 328 observed median values of sub-surface snow NO3<sup>-</sup> concentrations are used for modeled 329 sub-surface (from 2-cm depth to the bottom of the snow photic zone,  $z_{3e}$ ) snow NO<sub>3</sub><sup>-</sup> 330 concentrations  $(NO_3^{-1})_{bot}$  across Antarctica and Greenland. Although there is a large variation in observed snow NO<sub>3</sub><sup>-</sup> concentrations from the ITASE campaign [Bertler et al., 331 332 2005], there is no clear spatial pattern across Antarctica. In Greenland, the relationship 333 between snow accumulation rate and snow NO3<sup>-</sup> concentrations is non-linear and snow accumulation alone cannot account for the spatial variability in NO3<sup>-</sup> concentrations in 334 335 lower snow accumulation rate regions [Burkhart et al., 2009]. In Antarctica, snow NO<sub>3</sub> 336 concentrations in the top 2 cm of snow are up to 10 times higher than  $NO_3^{-1}$ 337 concentrations below 2-cm depth [Dibb et al., 2004, Erbland et al., 2013, Frey et al., 338 2009, Mayewski and Legrand, 1990, Rothlisberger et al., 2000], while in Greenland, 339 surface snow layers are at most 2 times higher compared to sub-surface snow layers 340 [*Dibb et al.*, 2007]. In this study, NO<sub>3</sub><sup>-</sup> concentrations in the top 2 cm of snow ( $[NO_3^-]_{top}$ ) in Antarctica are calculated by enhancing  $[NO_3]_{bot}$  by a factor of 6 in the top 2 cm of 341 342 snow, which is the median of observed  $NO_3^-$  enhancement factors (*EF*) in Antarctica [Dibb et al., 2004, Erbland et al., 2013, Frey et al., 2009, Mayewski and Legrand, 1990, 343 344 Rothlisberger et al., 2000]. Since  $NO_3^-$  concentrations in Antarctica are enhanced by a factor of 6 in the top 2 cm of snow, an equal amount of NO<sub>3</sub><sup>-</sup> has been removed from the 345 346 remainder of the photic-zone depth to maintain mass balance of NO<sub>3</sub><sup>-</sup> within the snow 347 column. In the modeled Greenland snowpack,  $[NO_3]_{top}$  is not enhanced (*EF*=1) although 348 EF is varied in a sensitivity study to assess the impact of EF on snow-sourced NO<sub>x</sub> in 349 Greenland (section 3.7).

350

351 As mentioned in the introduction, the measured quantum yields for the dominant  $NO_3^{-1}$ photolysis pathway (E1) range from 0.003 molec photon<sup>-1</sup> [*Chu and Anastasio*, 2003] to 352 0.6 molec photon<sup>-1</sup> [Zhu et al., 2010] at T=253K and exhibit a dependency on 353 354 temperature (see temperature-dependent equation in Chu and Anastasio [2003]). A higher 355 fraction of  $NO_3^-$  was likely present on ice surfaces in the Zhu et al. [2010] study 356 compared to the Chu and Anastasio [2003] study due to the different sample preparation 357 methods, and likely explains the 3 order-of-magnitude difference in quantum yields. This 358 interpretation suggests NO<sub>3</sub> on the surface of ice grains is much more photolabile 359 compared to NO<sub>3</sub><sup>-</sup> embedded within ice grains, consistent with results from Meusinger et 360 al. [2014]. In this study, we assume that  $NO_3^-$  that is wet deposited to the snow surface is 361 more likely to be embedded in the interior of a snow grain compared to  $NO_3^-$  that is dry deposited to the surface of the snow grain. To simulate this effect in an idealized 362 363 snowpack, we do not adjust the quantum yields, but instead scale snow  $NO_3^{-1}$ concentrations by the fraction of dry relative to total (wet + dry) deposition to the 364 365 Antarctic and Greenland snow surface, assuming that only the fraction of dry deposited 366  $NO_3^-$  is photolabile ( $F_n$ ). The degree of migration of  $NO_3^-$  within a snow grain after deposition due to snow metamorphism is unknown, which may influence the photolability of  $NO_3^-$  [*Domine and Shepson*, 2002].

369

370 Other modeling studies have attempted to calculate the fraction of photolabile  $NO_3^{-1}$  in 371 snow by estimating the concentration of  $NO_3^-$  contained within the liquid-like region 372 (LLR) on the surface of ice grains (e.g., Thomas et al., 2012). In this work, we do not 373 explicitly calculate  $NO_3^-$  photolysis within the LLR because there are still many 374 unknowns about the LLR [Domine et al., 2013], including the distribution of NO<sub>3</sub><sup>-</sup> 375 between the bulk snow and the LLR. This distribution is better understood for some 376 species, such as chloride [*Cho et al.*, 2002], but it is unclear if NO<sub>3</sub><sup>-</sup> behaves similarly. In 377 this study, we have assumed that all NO<sub>x</sub> formed from the photolysis of photolabile NO<sub>3</sub><sup>-</sup> 378 is transferred to the boundary layer, which neglects any recombination chemistry (e.g., Erbland et al., 2015) that prevents NO<sub>x</sub> from leaving the ice matrix, and may lead to 379 380 overestimates in the modeled  $F_{NOx}$  values presented in this study. The quantum yield for 381  $NO_3$  photolysis is dependent on the location of  $NO_3$  in snow, and although there are 382 uncertainties surrounding the location of  $NO_3^{-1}$  in snow, we perform model sensitivity 383 studies using the full range of measured quantum yields to provide bounds for the amount 384 of  $NO_x$  produced from snow  $NO_3^-$  photolysis.

385

#### 386 2.2. Model Sensitivity Studies

387 Model results from Greenland and Antarctica are compared in section 3, which sheds 388 light on the influence of chemical and meteorological parameters on  $F_{NOx}$ , nitrogen 389 recycling and redistribution, and NO<sub>3</sub><sup>-</sup> loss from snow. Additionally, due to uncertainties 390 in our understanding of snow photochemistry [Domine et al., 2013], we perform a variety 391 of model sensitivity studies, as shown in Table 3. The range of values used for the 392 parameters varied in the sensitivity studies reflects our estimates of their uncertainties. 393 The majority of these sensitivity studies focus on Antarctica, but the sensitivity of  $F_{NOx}$  to 394 EF is tested in Greenland. The quantum yield is varied from 0.002 molec photon<sup>-1</sup> 395 (corresponding to T=244 K) [Chu and Anastasio, 2003] to 0.6 molec photon<sup>-1</sup> [Zhu et al., 396 2010]. Snow NO<sub>3</sub><sup>-</sup> concentrations below 2 cm ( $/NO_3^-/_{bot}$ ) are halved and doubled with 397 respect to the base case scenario and the impact of scaling NO<sub>3</sub><sup>-</sup> concentrations by the 398 fraction of photolabile NO<sub>3</sub> ( $F_p$ ) is investigated. The NO<sub>3</sub> enhancement factor in the top 399 2 cm of snowpack is varied from 1 to 10, based upon the range of reported observations 400 [Dibb et al., 2004, Frey et al., 2009, Mayewski and Legrand, 1990, Rothlisberger et al., 2000].  $C_{BC}$  is halved and doubled with respect to the base case scenario. Since field and 401 satellite measurements suggest significant increases in surface  $r_e$  throughout austral 402 403 summer in Antarctica [Jin et al., 2008, Klein, 2014], the impact of increasing surface  $r_e$ during austral summer on  $F_{NOx}$  in Antarctica is evaluated in this study. The  $r_e$  profiles are 404 405 varied in three sensitivity studies to examine its influence on  $F_{NOx}$ . The bulk extinction coefficient for snow (Kext<sub>tot</sub>) is increased and decreased by 20% with respect to the base 406 407 case scenario because Libois et al. [2013] suggest that the spherical snow grain 408 assumption overestimates e-folding depths by a factor of 1.2. These sensitivity studies are 409 used to provide estimates of the influence the uncertainty in these parameters on 410 calculations of  $F_{NOx}$ .

#### 412 2.3. Estimating the Impact of Snow NO<sub>3</sub><sup>-</sup> Photolysis on Boundary Layer Chemistry 413 and Ice-Core NO<sub>3</sub><sup>-</sup> Records

414 NO<sub>3</sub><sup>-</sup> photolysis, followed by oxidation, recycling, and redistribution of snow-sourced 415 reactive nitrogen, influences both boundary layer chemistry and the concentration and 416 isotopic signature of  $NO_3^-$  that is ultimately preserved in ice-core records. The 417 preservation of NO<sub>3</sub><sup>-</sup> in ice cores is most dependent on the amount of NO<sub>3</sub><sup>-</sup> lost from the 418 snow through photolysis via transport of snow-sourced NO<sub>x</sub> away from the site of 419 primary deposition. The methods used to explore and quantify nitrogen recycling and 420 photolysis-driven loss of  $NO_3^-$  in snow are described in the following sections.

421

#### 422 2.3.1. Reactive Nitrogen Recycling Between the Air and Snow

423 The Nitrogen Recycling Factor (NRF) is a metric originally proposed by Davis et al. 424 [2008] to quantify the degree of reactive nitrogen recycling in snow over 1 year. The 425 *NRF* is calculated in E9:

426

$$427 \qquad NRF = \frac{F_{NOX}}{F_{PRI}},$$

$$428$$

In E9,  $F_{NOx}$  (ng N m<sup>-2</sup> yr<sup>-1</sup>) is the annual sum of NO<sub>x</sub> released from the snow and  $F_{PRI}$  (ng 429 N m<sup>-2</sup> vr<sup>-1</sup>) is the annual sum of primary NO<sub>3</sub> deposited to the snow. Davis et al. [2008] 430 431 use the NRF to describe nitrogen recycling on both macro-scale (e.g., across the East 432 Antarctic plateau) and micro-scale (e.g., the number of times one molecule of NO<sub>3</sub><sup>-</sup> is 433 recycled) levels. An NRF greater than 1 suggests that multiple nitrogen recycling events 434 occur in the snow. NRF represents the average, or "bulk" degree of nitrogen recycling in 435 snow because it does not resolve the degree of nitrogen recycling on a molecular level in 436 the snow; some  $NO_3^-$  molecules may never be photolyzed while other  $NO_3^-$  molecules 437 may be photolyzed and recycled many times greater than NRF. The NRF has 438 implications for boundary layer chemistry because the continual re-emission of NO<sub>x</sub> 439 enhances the effective concentration of  $NO_x$  in the boundary layer [Davis et al., 2008]. 440 Additionally, nitrogen recycling between the air and snow may alter the preservation of 441  $NO_3^-$  in ice-core records.

442

#### 443 2.3.2. Export of Snow-sourced Nitrate Away from the Original Site of Photolysis

444 Once snow-sourced  $NO_x$  is emitted to the atmosphere, it is subject to transport away from 445 the original site of photolysis. If snow-sourced NO<sub>x</sub> is oxidized to HNO<sub>3</sub> and re-deposited 446 back to the snow surface, then there is no net photolysis-driven loss of NO<sub>3</sub><sup>-</sup> from the 447 snow. However, if some of the snow-sourced  $NO_x$  is transported away from the site of 448 primary deposition, there is a net photolysis-driven loss of NO<sub>3</sub><sup>-</sup> from the snow. The 449 fraction (f) of total  $NO_3^-$  (photolabile + non-photolabile) lost from the snow driven by 450 photolysis is calculated in E10:

451

452 
$$f = \left(\left(\frac{F_R}{F_{NOx}}\right)^{\tau_{NO_3}} burial} - 1\right) \times F_P \times F_{P\,photo}$$
E10

453

In E10, negative values of f represent loss of  $NO_3^-$  from the snow and positive values of f 454 represent gain of NO<sub>3</sub><sup>-</sup> to the snow. In E10,  $F_R$  (ng N m<sup>-2</sup> yr<sup>-1</sup>) is the total annual flux of recycled NO<sub>3</sub><sup>-</sup> to the snow surface and  $F_{NOx}$  (ng N m<sup>-2</sup> yr<sup>-1</sup>) is the total annual flux of NO<sub>x</sub> 455 456

E9

released from the snow from photolysis of snow NO<sub>3</sub><sup>-</sup>.  $F_R$  is calculated by subtracting the 457 458 depositional flux of NO3<sup>-</sup> from a model run without snow photochemistry from the 459 depositional flux of NO<sub>3</sub><sup>-</sup> from a model run with snow photochemistry. The ratio of  $F_R$  to 460  $F_{NOx}$  represents the fraction of photolabile NO<sub>3</sub><sup>-</sup> remaining in the snow after 1 year. As long as  $NO_3^-$  remains in the photic zone,  $NO_3^-$  can continually be lost from the snow by 461 462 photolysis-driven processes. The preservation of  $NO_3^-$  in ice cores is dependent on the 463 fraction of  $NO_3^{-1}$  lost from the snow through photolysis during the entire time that  $NO_3^{-1}$ remains in the photic zone. Provided that there are no major changes in parameters that 464 influence snow photochemistry (e.g., LAI, overhead ozone abundance) from year to year, 465 466 the fraction of photolabile  $NO_3^-$  lost from the snow will be stable from year to year.

467

468  $\tau_{NO_{3}^{-}burial}$  represents the time that NO<sub>3</sub><sup>-</sup> remains in the photic zone (years) and in E10, 469  $\tau_{NO_{3}^{-}burial}$  accounts for the loss of NO<sub>3</sub><sup>-</sup> that occurs during the entire time that it remains 470 in the photic zone. When NO<sub>3</sub><sup>-</sup> remains in the photic zone for less than a year ( $\tau_{NO_{3}^{-}burial}$ 471 < 1),  $\tau_{NO_{3}^{-}burial}$  in E10 is set equal to 1.  $\tau_{NO_{3}^{-}burial}$  is calculated according to E11, where 472 both the depth of the photic zone (cm) and the total annual snow accumulation ( $\alpha_r$ ) (cm 473 yr<sup>-1</sup>) are considered.

$$475 \quad \tau_{NO_3^- burial} = \frac{z_e}{\alpha_r}, \tag{E11}$$

476

In E11,  $z_e$  (cm) is 1 e-folding depth of UV actinic flux and is used instead of  $z_{3e}$  because 477 87-91% of snow-sourced  $NO_x$  is produced within the top 1 e-folding depth. To convert total annual snow accumulation rate from kg m<sup>-2</sup> yr<sup>-1</sup> to cm, a typical polar snow density 478 479 (0.36 g cm<sup>-3</sup>) [Grenfell et al., 1994] is assumed.  $\tau_{NO_{3 burial}}$  is the minimum amount of 480 time on average that NO3<sup>-</sup> remains in the top one-third of the snow photic zone before 481 482 burial beneath because nitrogen recycling, which effectively redistributes NO<sub>3</sub><sup>-</sup> upwards in the snow, is not factored into E11.  $\tau_{NO_3^- burial}$  thus represents the lifetime of NO<sub>3</sub><sup>-</sup> in snow in an average sense and does not resolve photolysis and recycling of individual 483 484 485 NO<sub>3</sub><sup>-</sup>molecules.

486

487 In E10,  $\left(\left(\frac{F_R}{F_{NOx}}\right)^{\tau_{NO_3^-}burial} - 1\right)$  represents the fraction of photolabile NO<sub>3</sub><sup>-</sup> lost from the 488 snow through photolysis. This fraction is multiplied by  $F_p$  to calculate the fraction of total 489 (photolabile + non-photolabile) NO<sub>3</sub><sup>-</sup> lost from the snow through photolysis (f). If the 490 lifetime of NO<sub>3</sub><sup>-</sup> against burial is shorter than the lifetime of NO<sub>3</sub><sup>-</sup> against photolysis,  $F_p$  is 491 multiplied by  $F_{p_{photo}}$ , which represents the fraction of photolabile NO<sub>3</sub><sup>-</sup> that is buried 492 below the snow photic zone before photolysis.  $F_{p_{photo}}$  is calculated in E12:

493

494 
$$F_{p_{photo}} = \frac{\tau_{NO_3^-burial}}{\tau_{NO_3^-photolysis}}$$
, E12

495

496 where  $\tau_{NO_3^- burial}$  represents the lifetime of NO<sub>3</sub><sup>-</sup> against burial below the photic zone and 497  $\tau_{NO_3^- photolysis}$  is lifetime of NO<sub>3</sub><sup>-</sup> against photolysis.  $\tau_{NO_3^- photolysis}$  is calculated in E13:

498  
499 
$$\tau_{NO_{3}^{-}photolysis} = \frac{1}{J}$$
, E13  
500

501 In E13, J (E8) has been integrated from  $\lambda$ =298-345 nm and averaged over the top e-502 folding depth in snow ( $z_e$ ).  $\tau_{NO_3^-photolysis}$  represents the average lifetime of NO<sub>3</sub><sup>-</sup> in the 503 snow photic zone against loss by photolysis.

504

505 If f is 0, then all snow-sourced NO<sub>x</sub> is redeposited to the snow and there is no net loss of 506  $NO_3$ . f is also 0 if the net export of snow-sourced  $NO_x$  away from the site of original 507 photolysis is balanced by net import of snow-sourced NO<sub>x</sub> from other Antarctic or 508 Greenland locations. If  $-1 \le f \le 0$ , the export of local snow-sourced NO<sub>x</sub> is higher than the 509 deposition of snow-sourced NO<sub>x</sub> from elsewhere in Antarctica or Greenland, resulting in 510 net photolysis-driven loss of NO<sub>3</sub><sup>-</sup> from the snow. If f > 0, the export of local snow-511 sourced NO<sub>x</sub> is lower than the deposition of snow-sourced NO<sub>x</sub> from elsewhere in 512 Antarctica or Greenland, resulting in net photolysis-driven gain of  $NO_3^-$  to the snow.

513

514 *f* is used to calculate the enrichment in ice-core  $\delta^{15}N(NO_3^-)$  due solely to the impact of 515 photolysis-driven loss of NO<sub>3</sub><sup>-</sup> in snow when  $-1 \le f < 0$ . We use a Rayleigh fractionation 516 equation used to calculate  $\delta^{15}N(NO_3^-)$  [*Blunier et al.*, 2005]: 517

518 
$$\delta^{15}N(NO_3^-) = ((\delta^{15}N(NO_3^-)_{air} + 1) \cdot (1+f)^{\epsilon}) - 1$$
 E14

519

In E14,  $\delta^{15}N(NO_3^-)_{air}$  is the annual-averaged  $\delta^{15}N$  value of boundary layer NO<sub>3</sub><sup>-</sup> and  $\epsilon$ 520 is the fractionation constant (-47.9‰ [Berhanu et al., 2014]). In this work, we set 521  $\delta^{15}N(NO_3^-)_{air}$  equal to 0% to investigate the enrichment in  $\delta^{15}N(NO_3^-)$  only from 522 photolysis-driven loss of NO<sub>3</sub><sup>-</sup> from snow. The model-calculated  $\delta^{15}N(NO_3)$  enrichments 523 524 represent values in snow below the photic zone (i.e., at depths where sunlight is minimal 525 and NO<sub>3</sub> photolysis is not occurring) because these values are indicative of the loss of 526 snow NO<sub>3</sub><sup>-</sup> over the total time that it spent in the snow photic zone. When NO<sub>3</sub><sup>-</sup> is buried below the snow photic zone, no more photolysis or alteration of  $\delta^{15}N(NO_3^{-1})$  occurs; 527 hence, the  $\delta^{15}N(NO_3)$  is further unaltered and is preserved in the ice-core record. The 528 529 chemical and physical properties of snow within the photic zone are used to estimate what the enrichment in  $\delta^{15}N(NO_3)$  would be in ice cores at depths below the snow photic 530 531 zone.

532

#### 533 **3. Results and Discussion**

#### 534 **3.1.** Parameters that Influence *F<sub>NOX</sub>* and its Spatial Redistribution

535 Figure 2a and 2b present modeled air temperature in the lowest vertical grid boxes, which 536 range from 237-271 K in Antarctica and 257-280 K in Greenland; lowest temperatures 537 are located at the highest elevations. Figure 2c and 2d show modeled total annual snow accumulation rates from GEOS-Chem (kg m<sup>-2</sup> yr<sup>-1</sup>), ranging from 10-700 kg m<sup>-2</sup> yr<sup>-1</sup> in 538 Antarctica and 60-1400 kg m<sup>-2</sup> yr<sup>-1</sup> in Greenland. In both regions, the decrease in snow 539 540 accumulation rate from the coast to the top of the plateau is attributed to increased 541 distance from the ocean (moisture source), increased elevation, and lower temperatures. 542 Figure 2e and 2f show modeled annual mean surface wind divergence from May 2009 to

- May 2010. Figure 2e is consistent with Antarctic Mesoscale Prediction System surface wind output [Figure 3 in *Parish and Bromwich*, 2007], indicating that the large-scale airflow pattern in Antarctica flows from the East Antarctic plateau downslope towards the coast (katabatic winds). There are three major regions of wind convergence in Antarctica, located near the Ross, Ronne, and Amery ice shelves. In Greenland, air generally flows downwards from the continental interior towards the coasts.
- 549

Figure 3a and 3b present modeled snow  $C_{BC}$ , ranging from 0.08 to 0.6 ng g<sup>-1</sup> in Antarctica 550 and 0.8 to 5.5 ng g<sup>-1</sup> in Greenland. In Antarctica, the highest  $C_{BC}$  values are found on the 551 East Antarctic plateau and the spatial pattern of  $C_{BC}$  is governed by the snow 552 553 accumulation rate; higher snow accumulation rates dilute  $C_{BC}$  [Doherty et al., 2013]. The modeled boundary layer black carbon concentrations are relatively uniform across 554 555 Antarctica (0.1-0.6 pptv) because the majority of black carbon reaches Antarctica through 556 long-range transport (with the exception of local production from Antarctic research 557 stations). In Greenland, the highest  $C_{BC}$  values are found at the coasts due to their 558 proximity to pollution sources, even though snow accumulation rates are highest at the 559 coasts as well. Figure 3c and 3d show mean summer (DJF, Antarctica; JJA, Greenland) e-560 folding depths of UV actinic flux in snow  $(z_e)$ , which range from 24 to 69 cm in 561 Antarctica and 2 to 17 cm in Greenland. The shallowest e-folding depths are located in 562 regions of relatively high  $C_{BC}$ , which is on the plateau in Antarctica and at the coasts in 563 Greenland. Higher  $C_{BC}$  in snow results in shallower  $z_e$  because UV absorption in snow is enhanced as the concentration of LAI increases [Zatko et al., 2013]. In this study, coastal 564 565 grid boxes are a mixture of water, sea ice, and snow-covered surfaces, and since actinic 566 flux profiles are only calculated for snow-covered surfaces, the average  $z_e$  in coastal grid 567 boxes are artificially shallow.

568

Figure 4a and 4b show the fraction of dry-deposited NO<sub>3</sub><sup>-</sup> compared to total deposited NO<sub>3</sub><sup>-</sup> across Antarctica and Greenland. The ratio of dry deposition to total deposition ranges from 0.05 in coastal Antarctica and coastal Greenland to 0.92 in central Greenland and 0.99 on the East Antarctic plateau. Figure 4c and 4d show annual mean sub-surface (from 2-cm depth to the bottom of the photic zone,  $z_{3e}$ ) snow NO<sub>3</sub><sup>-</sup> concentrations ([NO<sub>3</sub><sup>-</sup> J<sub>bot</sub>) in the model scaled by  $F_p$ , ([NO<sub>3</sub><sup>-</sup>]<sub>bot</sub> x  $F_p$ ), which ranges from 13-60 ng g<sup>-1</sup> across Antarctica and 8-121 ng g<sup>-1</sup> across Greenland.

576

# 577 3.2. Emission, Recycling, and Redistribution of Nitrogen Across Antarctica and 578 Greenland

Figure 5 shows model-calculated mean summer snow-sourced NO<sub>x</sub> flux,  $\overline{F_{NOx}}$ , across Antarctica and Greenland. The spatial patterns of  $\overline{F_{NOx}}$  in Figure 5 are governed by the depth of the photic zone ( $z_e$ ) (especially in Antarctica) and also the concentration of photolabile NO<sub>3</sub><sup>-</sup> (Figure 4a and 4b), which is lowest at the coasts in both Antarctica and Greenland in the model. Additionally, spatial patterns of  $\overline{F_{NOx}}$  are influenced by the temperature-dependent quantum yield; the highest quantum yield values are located at the coasts where temperatures are highest.  $\overline{F_{NOx}}$  ranges from 0.01-6.4x10<sup>8</sup> molec cm<sup>-2</sup> s<sup>-1</sup> in Antarctica and 0.5-11x10<sup>8</sup> molec cm<sup>-2</sup> s<sup>-1</sup> in Greenland.

588 Figure 6a and 6b present the total annual depositional flux of primary  $NO_3^-(F_{PRI})$ , which ranges from  $0.9-35 \times 10^5$  ng N m<sup>-2</sup> yr<sup>-1</sup> in Antarctica and  $14-1000 \times 10^5$  ng N m<sup>-2</sup> yr<sup>-1</sup> in 589 590 Greenland and is highest at the coasts due to its relative proximity to NO<sub>x</sub>-source regions 591 in lower latitudes. An adjoint modeling study by Lee et al. [2014] suggests that boundary 592 layer  $NO_3^-$  abundance in Antarctica is dominated by  $NO_3^-$  transport to Antarctica 593 originating from NO<sub>x</sub> emissions from 25-65°S during austral winter and by thermal 594 decomposition of peroxyacyl nitrate (PAN) as it descends from the free troposphere in all 595 other seasons. In Greenland, boundary layer nitrate is predominately in the gas-phase 596 (HNO<sub>3</sub>) [Bergin et al., 1995, Dibb et al., 1994] and it has been suggested that NO<sub>3</sub><sup>-</sup> in 597 Greenland snow originates from both the troposphere [Geng et al., 2014a, Wespes et al., 598 2012] and stratosphere [Davidson et al., 1989].

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Figure 6c and 6d show the total annual depositional flux of recycled NO<sub>3</sub> ( $F_R$ ), which 600 ranges from  $0.5-11 \times 10^5$  ng N m<sup>-2</sup> yr<sup>-1</sup> in Antarctica and  $0.4-9 \times 10^5$  ng N m<sup>-2</sup> yr<sup>-1</sup> in 601 Greenland and is highest in areas of wind convergence (Greenland coasts and Antarctic 602 603 ice shelves). Figure 6e shows the ratio of deposition of recycled nitrogen ( $F_R$ ) to total 604 deposition  $(F_{PRI} + F_R)$  in Antarctica. In Antarctica, recycled nitrogen  $(F_R)$  is the dominant 605 (50-85%) form of NO<sub>3</sub><sup>-</sup> deposition along the Transantarctic mountains and in regions of 606 wind convergence, such as the Ronne, Ross, and Amery ice shelves. Along the Antarctic 607 coast,  $F_R$  represents as little as 11% of the deposition flux. Figure 6f shows the ratio of 608 total annual primary  $NO_3^-$  deposition to total  $NO_3^-$  deposition (primary + recycled) in 609 Greenland. Primary NO<sub>3</sub><sup>-</sup> deposition is responsible for at least 85% of total NO<sub>3</sub><sup>-</sup> 610 deposition across most of Greenland, and up to 100% in southern Greenland. This ratio is 611 set equal to 0 over the ocean and sea ice, which leads to artificially low ratios at the coast 612 because the model grid boxes at the coasts are an average of ocean, sea-ice, and 613 continental values.

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615 Figure 7 shows the Nitrogen Recycling Factor (*NRF*), which ranges from 0.2 to 12 across 616 Antarctica and 0.01-1.6 across Greenland. Nitrogen is recycled multiple times over the 617 course of 1 year across most of Antarctica, with the exception of the coasts. In contrast, 618 *NRF* values are less than 1.5 across all of Greenland. The spatial pattern of *NRF* is governed by the flux of snow-sourced NO<sub>x</sub>, which is influenced by the depth of the 619 620 photic zone  $(z_e)$ , the concentration of photolabile NO<sub>3</sub>, and the temperature-dependent 621 quantum yield for NO<sub>3</sub><sup>-</sup> photolysis. The spatial pattern of NRF is also dependent on  $F_{PRI}$ , 622 which is highest at the coasts and lowest at the top of the plateaus. In both Antarctica and 623 Greenland, NRF values are lowest near the coast because the fraction of photolabile  $NO_3^{-1}$ 624 is small and  $F_{PRI}$  values are high. The maximum NRF values generally occur in regions 625 corresponding to maximum  $F_{NOx}$  values. The average NRF value presented in Erbland et 626 al. [2015] for Dome C is 4, and the difference in model-estimates of nitrogen recycling at 627 Dome C in Erbland et al. [2015] and in this study (NRF = 6 at Dome C) is at least 628 partially due to the assumption in Erbland et al. [2015] that 20% of snow-sourced  $NO_3^-$  is 629 transported away from Dome C via katabatic winds. In GEOS-Chem, 25% of snow-630 sourced NO<sub>3</sub><sup>-</sup> is transported away at Dome C, which is slightly larger than the assumption in Erbland et al. [2015]. Larger NO<sub>3</sub><sup>-</sup> export fractions will lead to larger loss of snow 631 632  $NO_3^{-}$ , which may also lead to a larger number of recycling events via transport and re-633 deposition of snow-sourced NO<sub>x</sub> throughout East Antarctica. Davis et al. [2008] estimate 634 an NRF of 1.8, which is roughly 3 to 6 times lower than the modeled East Antarctic NRF 635 values in this study (NRF=5-10), although Davis et al. state that their estimated NRF 636 value could be factors of 3 to 5 times higher due to uncertainties in primary nitrogen 637 deposition estimates. Erbland et al. [2015] found a relationship between inverse snow 638 accumulation rates and nitrogen recycling in regions where the snow accumulation rates  $(\alpha_r)$  are higher than 50 kg m<sup>-2</sup> a<sup>-1</sup> in Antarctica. The relationship between NRF and  $1/\alpha_r$  in 639 our model simulations for  $\alpha_r > 50$  kg m<sup>-2</sup> a<sup>-1</sup> in Antarctica suggests that only 22% of the 640 spatial variability of NRF can be explained by  $\alpha_r$  (Figure S1), likely due to the 641 642 redistribution of snow-sourced reactive nitrogen across Antarctica by winds. In 643 Greenland, which is a region where snow accumulation rates are greater than 50 kg m<sup>-2</sup> a<sup>-2</sup> 644 , our model simulations suggest that only 30% of the spatial variability of NRF can be 645 explained by  $\alpha_r$  (Figure S2).

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#### 647 **3.3. Impact of Reactive Nitrogen Recycling on Boundary Layer Chemistry**

648 The height of the boundary layer will strongly influence the abundance of NO<sub>3</sub>, reactive 649 nitrogen oxides, and oxidants emitted or formed at or near the surface. At many polar 650 stations (e.g., Neumayer, South Pole, Dome C, Halley, Kohnen, Summit) there is a wide 651 range of observed boundary layer heights during summer (10-600 m [Casasanta et al., 652 2014, Cohen et al., 2007, Davis et al., 2004, Drue and Heinemann, 2007, Handorf, 1996, 653 Helmig et al., 2002, Jones et al., 2006, 2008, King et al., 2006, Kodama et al., 1985, 654 Konig-Langlo et al., 1998, Neff et al., 2008, Oncley et al., 2004, Travouillon et al., 2008, 655 *Weller et al.*, 1999]), and although modeled boundary layer heights are not systematically 656 biased in one direction compared to observations, they often do not agree well. Therefore, 657 only the relative impacts of snow photochemistry on reactive nitrogen and oxidant 658 abundances are evaluated in this study. The impact of snow NO<sub>3</sub><sup>-</sup> photolysis on boundary 659 layer chemistry can be examined by considering factor changes in boundary layer NO<sub>x</sub>, 660  $NO_3^-$ , OH, and  $O_3$  mixing ratios between simulations with and without snow  $NO_3^-$ 661 photolysis. Other snow photochemical reactions mentioned in the introduction but not 662 included in this modeling study will also impact oxidant abundances, but the effects of 663 each photochemical reaction are not additive due to the highly non-linear nature of 664 oxidant cycling.

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666 As shown in Figure 8, the inclusion of a snow NO<sub>x</sub> source leads to factor increases in 667 boundary layer mixing ratios of NO<sub>x</sub> from 1.2-24.5, gas-plus aerosol-phase nitrate from 668 1.0-11.8, OH from 1.8-5.3, and  $O_3$  from 1.1-1.8 in Antarctica. The largest factor increases 669 are in West Antarctica, particularly near the Ross and Ronne ice shelves, where winds 670 carrying photo-produced species converge. The surface transport pattern is especially 671 important for the redistribution of the longer-lived species NO<sub>3</sub><sup>-</sup> and O<sub>3</sub>. Figure 9 shows 672 that the inclusion of a snow NO<sub>x</sub> source leads to factor increases in boundary layer 673 mixing ratios of NO<sub>x</sub> from 1.0-6.3, gas-plus aerosol-phase nitrate from 1.0-2.2, OH from 674 1.1-2.4, and O<sub>3</sub> from 1.0-1.14 in Greenland. The largest factor increases for short-lived 675 species (NO<sub>x</sub> and OH) are in central Greenland where  $F_{NOx}$  is highest (Figure 5) while the largest factor increases for longer-lived species (NO<sub>3</sub><sup>-</sup> and O<sub>3</sub>) are located in regions of 676 677 wind convergence.

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#### 679 **3.4. Implications for Ice-Core Records of** NO<sub>3</sub><sup>-</sup> **Concentrations and Isotopes**

680 Figure 10a and 10b show the minimum amount of time that snow NO<sub>3</sub><sup>-</sup> remains in the photic zone on average,  $\tau_{NO_3 \text{ burial}}$  (E11) as calculated in the model. NO<sub>3</sub><sup>-</sup> remains in the 681 snow photic zone for 4 months near the Antarctic coasts and up to 6.5 years on the East 682 683 Antarctic plateau before burial below the photic zone. In Greenland, NO<sub>3</sub><sup>-</sup> remains in the 684 photic zone for 0.1 months at the coasts and up to 7 months in central Greenland. The spatial pattern of  $\tau_{NO_3^-burial}$  is governed by the snow accumulation rate, both directly and indirectly through its influence on  $C_{BC}$ . The spatial patterns of  $\tau_{NO_3^-burial}$  are in agreement 685 686 with the expectation that NO<sub>3</sub><sup>-</sup> remains in the photic zone the longest in areas with low 687 snow accumulation rates. 688

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690 In Antarctica, the lifetime of NO<sub>3</sub><sup>-</sup> against burial is always longer than the lifetime of 691  $NO_3^-$  against photolysis. However in Greenland, the lifetime of  $NO_3^-$  against burial may be shorter than the lifetime against photolysis due to the much shorter  $\tau_{NO_3 burial}$  values. 692 Figure 10c presents the lifetime of nitrate against photolysis in Greenland 693  $(\tau_{NO_{3}^{-}photolysis}, E13)$ , which ranges from less than a month in northern Greenland to 37 694 months in southern Greenland. The spatial pattern of  $\tau_{NO_3^- photolysis}$  is dependent on the 695 mean summer (JJA) daily-averaged downwelling surface irradiance values (not shown), which are highest in northern Greenland. Figure 10d shows  $F_{p_{photo}}$  (E12), which ranges 696 697 from 0.003 to 1 across Greenland. Larger  $F_{p_{photo}}$  values in north-central Greenland 698 699 suggest that the lifetime of photolabile NO<sub>3</sub> against photolysis is sufficiently short 700 relative to its lifetime in the snow photic zone, allowing for a potentially large fraction of 701 snow  $NO_3^{-1}$  to be photolyzed before burial beneath the photic zone. In southeastern 702 Greenland, the lifetime of photolabile NO<sub>3</sub><sup>-</sup> against photolysis is longer than its lifetime 703 in the snow photic zone, which will reduce post-depositional photolysis of snow NO<sub>3</sub><sup>-</sup> in 704 this region.

705

706 Figure 11a and 11b show the fraction of NO<sub>3</sub><sup>-</sup> gained or lost from the snow through 707 photolysis (f, E10), which ranges from -0.99 to 0.30 in Antarctica and -0.83 to 0.02 in 708 Greenland. Throughout most of Antarctica and Greenland, values of f are negative 709 indicating that the photolysis of snow NO<sub>3</sub> leads to net loss of NO<sub>3</sub> from the snowpack 710 in most locations. Positive f values indicate regions with net gain of  $NO_3^-$  to the snow 711 resulting from the spatial redistribution of  $NO_3^-$  driven by snow photochemistry. In some 712 regions of convergence, such as over the Ronne Ice Shelf in Antarctica, there is a net gain 713 of snow-sourced  $NO_3^-$ . There are sharp gradients in f between the plateaus and the coasts. 714 The largest loss of snow NO<sub>3</sub><sup>-</sup> occurs at the top of the plateaus, where most photolyzed 715  $NO_3^-$  is transported away by katabatic winds. Along the coasts, the photolysis-driven loss 716 of NO<sub>3</sub><sup>-</sup> from the snow is minimal due to high snow accumulation rates and transport of snow-sourced  $NO_3^-$  from higher elevations. The spatial pattern of f is influenced by the 717 time that NO<sub>3</sub><sup>-</sup> remains in the photolytic zone  $(\tau_{NO_3^-burial})$ , the concentration of 718 photolabile  $NO_3^-(F_n)$ , and wind patterns across Antarctica and Greenland. 719

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Figure 11c and 11d show modeled enrichments in ice-core  $\delta^{15}N(NO_3^-)$  from photolysisdriven loss of NO<sub>3</sub><sup>-</sup> in snow. The  $\delta^{15}N(NO_3^-)$  enrichments presented in Figures 11c and 11d represent enrichments in  $\delta^{15}N(NO_3^-)$  below the snow photic zone where NO<sub>3</sub><sup>-</sup>

724 photolysis is not occurring, effectively integrating the total loss of NO<sub>3</sub><sup>-</sup> during its lifetime in the snow photic zone. Model-calculated ice-core  $\delta^{15}N(NO_3)$  values range 725 from 0 to 363‰ in Antarctica and 0 to 90‰ in Greenland and are highest at the tops of 726 the plateaus. The modeled ice-core  $\delta^{15}N(NO_3)$  enrichments resulting from the photolysis-727 728 driven loss of snow nitrate are sensitive to the fractionation constant ( $\epsilon$ ). In this study, the 729 fractionation constant is varied over the full range of values reported in Erbland et al 730 [2013], Frey et al., [2009], and Shi et al. [2015]; an  $\varepsilon$  of –90‰ increases modeled icecore  $\delta^{15}N(NO_3)$  by a factor of 2 and an  $\epsilon$  of -10% decreases modeled ice-core  $\delta^{15}N(NO_3)$ 731 732 ) by a factor of 5 across Antarctica and Greenland.

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#### 734 **3.5.** Relationship Between Nitrogen Recycling and Photolytic-loss of NO<sub>3</sub><sup>-</sup> in Snow

735 The degree of photolysis-driven loss of snow  $NO_3^-$  is determined by both rates of 736 photolysis and transport patterns across Antarctica and Greenland. The spatial patterns of 737 recycling (NRF, Figure 7) and loss (f, Figure 11a and 11b) differ across Antarctica and 738 Figure 12 shows the relationship between f and NRF across Antarctica. The magnitude of 739 nitrogen recycling and degree of photolysis-driven loss of snow NO<sub>3</sub><sup>-</sup> in Antarctica are well correlated ( $r^2 = 0.74$ , p< 0.001) in regions where NO<sub>3</sub><sup>-</sup> remains in the photic zone for 740 less than 2 years ( $\tau_{NO_3^-burial} < 2$ ) (Figure 12a). The relationship between recycling and loss breaks down ( $r^2 = 0.03$ , p< 0.001) in locations where NO<sub>3</sub><sup>-</sup> remains in the photic zone 741 742 743 for more than 2 years (Figure 12b). The relationship between recycling and loss weakens with increasing  $\tau_{NO_3^- burial}$  because recycling of reactive nitrogen occurs at or near the surface only, while loss of NO<sub>3</sub><sup>-</sup> occurs throughout the depth of snow photic zone. The 744 745 746 time that  $NO_3^-$  remains in the snow photic zone is dependent on snow accumulation rates 747 and the concentrations of LAI in snow, the latter of which is partially governed by snow accumulation rates. In the present climate,  $\tau_{NO_3^-burial}$  less than 2 years corresponds to snow accumulation rates higher than 130 kg m<sup>-2</sup> a<sup>-1</sup> in Antarctica. Figure 13 shows the 748 749 750 relationship between f and NRF in Greenland. Nitrogen recycling and the degree of photolysis-driven loss of snow NO<sub>3</sub> are well correlated ( $r^2 = 0.71$ , p < 0.001) in 751 Greenland, likely because NO<sub>3</sub><sup>-</sup> remains in the photic zone for less than 2 years across all 752 of Greenland. Snow accumulation rates are higher than 130 kg m<sup>-2</sup> a<sup>-1</sup> across much of 753 Greenland, except in parts of central Greenland, where snow accumulation rates are 754 between 60 and 100 kg m<sup>-2</sup> a<sup>-1</sup>. Differences in the relationship between snow 755 accumulation rate and  $\tau_{NO_3 burial}$  between Greenland and Antarctica are due to the higher 756 757 concentrations of LAI in Greenland snow.

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#### 759 **3.6. Comparison between Greenland and Antarctica**

Comparing model results in Greenland and Antarctica reveals information about how different parameters influence snow-sourced NO<sub>x</sub> fluxes, nitrogen recycling, and photolysis-driven loss of snow NO<sub>3</sub><sup>-</sup>. Air temperatures (Figure 2a, 2b) and annual snow accumulation rates (Figure 2c, 2d) are generally higher in Greenland compared to Antarctica, and Greenland is also closer to pollution sources, which lead to differences in the magnitudes and spatial patterns of  $F_{NOx}$ , *NRF*, and *f* between these two regions.

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The magnitude and spatial patterns of  $F_{NOx}$  are influenced by snow photic zone depths (which are governed by snow LAI concentrations), snow NO<sub>3</sub><sup>-</sup> concentrations, and the 769 quantum yield for  $NO_3^-$  photolysis. The e-folding depths of UV actinic flux are shallower 770 in Greenland (2-17 cm) compared to Antarctica (24-69 cm) because concentrations of light-absorbing impurities in snow are higher in Greenland ( $C_{BC} = 0.8-5.5 \text{ ng g}^{-1}$ ) 771 compared to Antarctica ( $C_{BC} = 0.08-0.6 \text{ ng g}^{-1}$ ). Similar to snow  $C_{BC}$ , the proximity of 772 Greenland to natural and anthropogenic NO<sub>x</sub> sources leads to higher observed snow NO<sub>3</sub> 773 774 values; the median of observed sub-surface snow  $NO_3^-$  concentrations in Greenland (132) 775 ng g<sup>-1</sup>, [Burkhart et al., 2009]) is over 2 times higher than observed sub-surface snow NO<sub>3</sub><sup>-</sup> concentrations in Antarctica (60 ng g<sup>-1</sup>, [Bertler et al., 2005]). Similarly, modeled 776 sub-surface snow NO<sub>3</sub> concentrations ( $[NO_3]_{bot} \ge F_p$ ) are higher across much of 777 778 Greenland compared to Antarctica. However, in coastal Antarctica and Greenland, 779 modeled sub-surface snow NO<sub>3</sub><sup>-</sup> concentrations ( $/NO_3^-/_{bot} \ge F_p$ ) are similar in magnitude 780 because the fraction of dry-deposited NO<sub>3</sub><sup>-</sup> to total deposited NO<sub>3</sub><sup>-</sup> ( $F_p$ ) is lower in Greenland, which lowers the concentration of photolabile NO3<sup>-</sup> in the model. 781 Additionally, modeled snow NO<sub>3</sub> in the top 2 cm of snow  $([NO_3]_{top})$  is generally higher 782 across Antarctica ( $[NO_3]_{top}$ =78-360 ng g<sup>-1</sup>) compared to Greenland ( $[NO_3]_{top}$ =8-121 ng g<sup>-1</sup>) because *EF*=6 in Antarctica and *EF*=1 in Greenland. Lastly, summer temperatures 783 784 785 are higher in Greenland (257-280 K) compared to Antarctica (237-271 K), which leads to higher quantum yields for NO<sub>3</sub><sup>-</sup> photolysis in Greenland ( $\phi$ =0.0032-0.0069) compared to 786 787 Antarctica ( $\phi$ =0.0015-0.0052).

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Fluxes of snow-sourced NO<sub>x</sub> ( $F_{NOx}$ ) are somewhat higher in Greenland (0.5-11x10<sup>8</sup>) 789 790 molec cm<sup>-2</sup> s<sup>-1</sup>) compared to Antarctica (0.01-6.4x10<sup>8</sup> molec cm<sup>-2</sup> s<sup>-1</sup>) because  $\phi$ , and often sub-surface snow NO<sub>3</sub><sup>-</sup> concentrations ( $/NO_3^-/_{bot} \ge F_p$ ), are higher in Greenland 791 792 compared to Antarctica, even though snow photic zones are shallower across Greenland 793 due to higher snow LAI concentrations. In Antarctica, the highest  $F_{NOx}$  values occur midway up the plateau because photic zone depths and the quantum yields for NO<sub>3</sub> 794 795 photolysis increase towards the coast while the concentrations of photolabile NO<sub>3</sub> 796 decrease towards the coast. In Greenland, the highest  $F_{NOx}$  values occur at the top of the 797 plateau because both photic zone depths and concentrations of photolabile  $NO_3^-$  are at a 798 maximum there. The quantum yield of NO3<sup>-</sup> photolysis increases towards the coast, 799 partially dampening the decreasing  $F_{NOx}$  gradient towards the coast.

800

801 The Nitrogen Recycling Factors (NRF) are much smaller in Greenland (NRF=0.01-1.6) 802 compared to Antarctica (NRF=0.2-12), suggesting that the degree of nitrogen recycling is much lower in Greenland. Although  $F_{NOx}$  values are generally higher in Greenland 803 804 compared to Antarctica, the degree of nitrogen recycling is lower in Greenland because 805 of its proximity to pollution sources. Primary deposition of NO<sub>3</sub><sup>-</sup> is over an order of magnitude larger in Greenland compared to Antarctica. The total nitrate, NO<sub>x</sub>, OH, and 806 807 O<sub>3</sub> boundary layer mixing ratio factor increases in Greenland are 16, 7, 3, and 2 times 808 lower, respectively, compared to Antarctica due to its proximity to pollution sources.

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810 NO<sub>3</sub><sup>-</sup> remains in the snow photic zone for a shorter period of time in Greenland (up to 7 811 months) compared to Antarctica (up to 6.5 years), because photic zones are shallower and 812 snow accumulation rates are higher in Greenland. The assumed fraction of photolabile 813 NO<sub>3</sub><sup>-</sup> ( $F_p$ ) in the model is lower in Greenland compared to Antarctica because a higher 814 fraction of NO<sub>3</sub><sup>-</sup> is wet-deposited in Greenland. Additionally, the lifetime of photolabile

- 815 NO<sub>3</sub> against burial beneath the snow photic zone  $(\tau_{NO_3 burial})$  is generally shorter than its 816 lifetime against photolysis  $(\tau_{NO_3 photolysis})$  in Greenland, further limiting post-817 depositional loss and recycling of snow NO<sub>3</sub>.
- 818

819 In both Greenland and Antarctica, the highest fractions of photolysis-driven loss of snow 820  $NO_3^{-}(f)$  are located at the top of the plateaus and the lowest f values are found at the 821 coasts. Photolysis-driven loss of snow NO<sub>3</sub> is generally lower in Greenland compared to 822 Antarctica because the fraction of photolabile  $NO_3^-(F_p)$  and the time spent in the snow photic zone  $(\tau_{NO_{3}})$  are lower in Greenland. Similar to the spatial patterns of f, 823 modeled enrichments in ice-core  $\delta^{15}N(NO_3^-)$  are highest on the plateau and lowest at the coasts in both Antarctica and Greenland.  $\delta^{15}N(NO_3^-)$  values in Antarctica are up to 4 824 825 times larger compared to Greenland because a higher fraction of  $NO_3^-$  is lost from the 826 827 snow via photolysis in Antarctica.

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#### 829 **3.7.** Exploring the Influence of Chemical, Optical, and Physical Parameters in Snow 830 on $F_{NOx}$

831 Snow accumulation rates have a major influence on snow-sourced NO<sub>x</sub> fluxes, nitrogen 832 recycling and redistribution, and photolysis-driven loss of snow  $NO_3^-$  via two 833 mechanisms. By impacting burial rates, the snow accumulation rate impacts the time 834 NO<sub>3</sub> spends in the snow photic zone. Snow accumulation rates influence light-absorbing 835 impurity concentrations in snow, which impact the depth of the snow photic zone. Snow 836 accumulation rates also influence concentrations of photolabile  $NO_3^{-1}$  in snow; both 837 directly through dilution effects and indirectly by controlling the partitioning of wet and 838 dry deposited NO<sub>3</sub><sup>-</sup>. Temperature also influences  $F_{NOx}$  via the temperature-dependent 839 quantum yield ( $\phi$ ), with higher temperatures leading to higher values of  $\phi$ .

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841 In addition to exploring the sensitivity of snow NO<sub>3</sub><sup>-</sup> photolysis to meteorological 842 parameters, the sensitivity of mean summer snow-sourced NO<sub>x</sub> fluxes ( $\overline{F_{NOx}}$ ) to a variety 843 of chemical, optical, and physical parameters in snow is explored in this section. 844 Although the vast majority of these sensitivity studies focus on Antarctica because model 845 computation time was limited, one sensitivity study was performed in Greenland and will 846 be described below.

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Table 3 shows the dependence of  $\overline{F_{NOX}}$  on uncertainties in the quantum yield for NO<sub>3</sub><sup>-</sup> 848 photolysis ( $\phi$ ), the fraction of photolabile NO<sub>3</sub><sup>-</sup> ( $F_p$ ), sub-surface snow NO<sub>3</sub><sup>-</sup> 849 850 concentrations  $([NO_3]_{bot})$ , radiation equivalent mean ice grain radii  $(r_e)$ , the bulk snow 851 extinction coefficient ( $Kext_{tot}$ ), the NO<sub>3</sub><sup>-</sup> concentration enhancement factor in the top 2 cm 852 of snow (EF), and snow black carbon concentrations in Antarctica. The range of values 853 for each of these parameters is determined by their estimated degree of uncertainty, in order to highlight the largest uncertainties in calculations of  $\overline{F_{NOx}}$ . The sensitivity study 854 results are compared to  $\overline{F_{NOX}}$  from the standard scenario, which is also described in Table 855 3. The  $\overline{F_{NOx}}$  values from the standard scenario are slightly different than the  $\overline{F_{NOx}}$  values 856 presented in Figure 5a because a spatially-uniform  $\phi$  is used in the standard scenario 857 858 while a temperature-dependent  $\phi$  is used in the rest of the manuscript (Figures 5-14).

 $\overline{F_{NOx}}$  is most sensitive to uncertainties in  $\phi$ , which increases  $\overline{F_{NOx}}$  by up to a factor of 330 when  $\phi$  from Zhu et al. [2010] (0.6 molec photon<sup>-1</sup>) is used compared to the  $\phi$  in the 860 861 standard scenario ( $\phi$ =0.002 molec photon<sup>-1</sup> [*Chu and Anastasio*, 2003]). The second most 862 influential parameter is the concentration of photolabile NO<sub>3</sub> ( $/NO_3$  /  $_{bot}$  x  $F_p$ ). Assuming 863 that all NO<sub>3</sub> is photolabile ( $F_p=1$ ) increases  $\overline{F_{NOx}}$  by up to a factor of 7.4 (at the coasts) 864 with respect to the standard scenario. Use of the fraction of dry-deposited NO<sub>3</sub> ( $F_p$ ) to 865 scale the concentration of photolabile NO<sub>3</sub><sup>-</sup> lowers  $\overline{F_{NOx}}$  by up to 85% along the coast, 866 but has little impact on the East Antarctic plateau due to the high fraction of dry 867 deposited NO<sub>3</sub>. Uncertainties in  $r_e$ , Kext<sub>tot</sub>, EF, and  $C_{BC}$  influence  $\overline{F_{NOx}}$  by up to a factor 868 of 1.3 compared to the standard scenario. In Greenland, EF is varied between 1 and 2, 869 870 which is the range of observed EF presented in Dibb et al. [2007]. Similar to Antarctica, varying *EF* from 1 to 2 increases  $\overline{F_{NOx}}$  in Greenland by at most a factor of 1.2. 871

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873 Calculated snow-sourced NO<sub>x</sub> fluxes are by far most sensitive to uncertainties in the 874 quantum yield for  $NO_3^-$  photolysis and the concentration of photolabile  $NO_3^-$ , which are 875 likely related to one another. The results of Chu et al. [2003], Zhu et al. [2010], and 876 Meusinger et al. [2014] suggest that the quantum yield is largely influenced by the 877 location of NO<sub>3</sub><sup>-</sup> in the ice grain; quantum yields tend to be lowest when NO<sub>3</sub><sup>-</sup> is buried 878 within the ice grain and highest when  $NO_3^-$  is on the ice grain surface. Although we have 879 not explicitly varied the quantum yield based upon  $NO_3^{-1}$  location in ice grains due to lack 880 of information regarding the location, we alter the concentration of photolabile  $NO_3^{-1}$ 881 based upon the ratio of dry to total deposited NO<sub>3</sub><sup>-</sup> This effectively assumes that only the 882 dry-deposited fraction of  $NO_3^{-1}$  is on the surface of the snow grain and is photolabile, and 883 that the wet-deposited  $NO_3^-$  is trapped within an ice grain and thus is not photolabile. 884 Using conservative  $\phi$  values (Chu and Anastasio, [2003]) and assuming that only drydeposited NO<sub>3</sub><sup>-</sup> is photolabile may suggest that our calculated  $F_{NOx}$  values represent a 885 886 lower limit. However, no field-or laboratory-based information is available to assess our 887 assumption that only dry-deposited  $NO_3^{-1}$  is photolabile; it is possible that the fraction of 888  $NO_3^{-1}$  that is photolabile is higher or lower than what we assume.

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890 These sensitivity studies highlight the need for field, laboratory, and modeling studies to 891 investigate factors influencing the quantum yield and concentration of photolabile NO<sub>3</sub>, such as the location of NO<sub>3</sub><sup>-</sup> within ice grains. We note that  $F_{NOx}$  is also sensitive to the 892 depth of the snow photic zone, which in turn is most sensitive to snow LAI 893 894 concentrations and  $r_e$  [Zatko et al., 2013]. By scaling the concentrations of total snow 895 LAI to  $C_{BC}$  in the model based on observations in Greenland and Antarctica [Zatko et al., 896 2013], we assume the same factor of 2 uncertainty in total LAI concentrations as we do 897 for  $C_{BC}$ . More measurements of snow LAI other than black carbon are needed to confirm 898 our assumptions about snow LAI concentrations and their associated uncertainty.

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#### 900 4. Conclusions

We have incorporated an idealized snowpack along with a snow radiative transfer model into a global chemical transport model (GEOS-Chem) and used this modeling framework to simulate the photolysis of snow NO<sub>3</sub><sup>-</sup> and calculate the associated snow-sourced flux and redistribution of nitrogen across Antarctica and Greenland. An important goal of this study is to explore the sensitivity of various chemical, optical, and physical properties of 906 snow, as well as meteorology, on fluxes of snow-sourced  $NO_x$  and related nitrogen 907 recycling, redistribution, and loss of  $NO_3^-$  from the snow. This modeling framework is 908 also used to examine the impact of snowpack  $NO_3^-$  photolysis on boundary layer 909 chemistry and the preservation of  $NO_3^-$  concentration and nitrogen isotopes in ice cores 910 across Antarctica and Greenland.

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912 The calculated fluxes of snow-sourced NO<sub>x</sub> in Antarctica and Greenland range from 0.01-6.4x10<sup>8</sup> molec cm<sup>2</sup> s<sup>-1</sup> and 0.5-11x10<sup>8</sup> molec cm<sup>2</sup> s<sup>-1</sup>, respectively. The modeled 913 914 spatial patterns of snow-sourced NO<sub>x</sub> fluxes are determined by the spatial patterns of 915 light-absorbing impurity (e.g., insoluble black carbon, dust, organics) concentrations in 916 snow, photolabile NO<sub>3</sub><sup>-</sup> concentrations, and temperature-dependent quantum yields for 917  $NO_3$  photolysis. In the model, the spatial patterns of light-absorbing impurities are 918 influenced by snow accumulation rates and proximity to pollution sources, the spatial 919 patterns of photolabile NO<sub>3</sub><sup>-</sup> in the model are influenced by the amount of wet-deposited 920  $NO_3$  compared to total deposited  $NO_3$ , and the spatial patterns of quantum yields of  $NO_3$ 921 photolysis are influenced by modeled surface air temperatures. Model sensitivity studies 922 suggest that the magnitude of the snow-sourced NO<sub>x</sub> flux is most sensitive to 923 uncertainties in the quantum yield for NO3<sup>-</sup> photolysis and the concentration of 924 photolabile  $NO_3^-$ , which are likely related to one another. The concentration of light 925 absorbing impurities in snow is also important for the flux of snow-sourced NO<sub>x</sub>, but we assume a much smaller degree of uncertainty in this parameter relative to uncertainties in 926 927 the quantum yield and the concentrations of photolabile NO<sub>3</sub><sup>-</sup>. Fluxes of snow-sourced 928 NO<sub>x</sub> are somewhat higher in Greenland compared to Antarctica because the quantum 929 yields, and often photolabile NO<sub>3</sub><sup>-</sup> concentrations, are higher in Greenland because 930 Greenland is warmer and closer to anthropogenic NO<sub>x</sub> sources (e.g., northern hemisphere 931 mid-latitudes). This is counteracted by the higher concentrations of snow light-absorbing 932 impurities leading to shallower show photic zones in Greenland.

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934 The Nitrogen Recycling Factor (*NRF*) is dependent on the magnitude and spatial patterns 935 of fluxes of primary  $NO_3^-$  to the snow relative to fluxes of  $NO_x$  from the snow. NRF 936 values greater than 1 suggest that nitrogen is recycled multiple times between the air and 937 snow. NRF values range from 0.2 to 12 in Antarctica and are greater than 1 across most 938 of Antarctica, with the exception of the coasts. NRF values range from 0.01 to 1.6 in 939 Greenland and are only larger than 1 in central Greenland. The degree of nitrogen 940 recycling is lower in Greenland because NO<sub>3</sub><sup>-</sup> deposition to snow is dominated by 941 primary NO<sub>3</sub><sup>-</sup> deposition in Greenland, largely because Greenland is closer to pollution 942 sources. Similarly, boundary-layer mixing ratios of total nitrate, NO<sub>x</sub>, OH, and O<sub>3</sub> are less 943 influenced by snow-sourced NO<sub>x</sub> in Greenland because of its proximity to pollution 944 sources.

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This modeling framework can also be used to examine the impact of  $NO_3^-$  photolysis on the preservation of  $NO_3^-$  in ice cores. The time that  $NO_3^-$  remains in the snow photic zone is dependent on snow accumulation rates both directly, and indirectly through their influence on light-absorbing impurity concentrations in snow.  $NO_3^-$  remains in the snow photic zone for a much shorter period of time in Greenland (up to 7 months) compared to Antarctica (up to 6.5 years) because snow accumulation rates are higher in Greenland. 952 The fraction of  $NO_3^-$  lost from the snow through photolysis (f) ranges from -0.99 to 0.30 953 in Antarctica and -0.83 to 0.02 in Greenland, where negative values indicate net loss of 954  $NO_3^{-1}$  from the snow. The fraction of  $NO_3^{-1}$  lost from the snow is dependent on the time 955 that  $NO_3^-$  remains in the snow photic zone, the concentration of photolabile  $NO_3^-$ , and 956 wind patterns across Antarctica and Greenland. In both Antarctica and Greenland, net 957 loss of snow NO<sub>3</sub><sup>-</sup> is highest on top of the plateaus and lowest at the coasts; some regions 958 (e.g., Ronne and Ross Antarctic ice shelves) experience net gain of snow  $NO_3^-$  due to the 959 redistribution of snow-sourced nitrogen. The fraction of photolysis-driven loss of snow NO<sub>3</sub><sup>-</sup> is lower in Greenland compared to Antarctica because the fraction of photolabile 960 NO<sub>3</sub><sup>-</sup> and the time spent in the snow photic zone ( $\tau_{NO_3 burial}$ ) is lower in Greenland due to 961 the higher snow accumulation rates and higher concentrations of snow light absorbing 962 impurities. The fraction of  $NO_3^-$  lost from the snow through photolysis is used to 963 calculate the enrichment in ice-core  $\delta^{15}N(NO_3^-)$  solely from photolysis-driven NO<sub>3</sub><sup>-</sup> loss 964 in snow. The modeled enrichment in ice-core  $\delta^{15}N(NO_3)$  ranges from 0 to 363‰ in 965 Antarctica and 0 to 90% in Greenland: the highest  $\delta^{15}N(NO_3)$  enrichments are calculated 966 967 in central Antarctica and Greenland.

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969 A significant relationship exists between nitrogen recycling and photolysis-driven loss of 970 snow NO<sub>3</sub><sup>-</sup> in Antarctica when NO<sub>3</sub><sup>-</sup> remains in the photic zone for less than 2 years  $(\tau_{NO_{3}^{-}burial} < 2)$ , corresponding to a snow accumulation rate greater than 130 kg m<sup>-2</sup> a<sup>-1</sup> in 971 the present day. There is also a significant relationship between nitrogen recycling and photolysis-driven loss of snow  $NO_3$  throughout all of Greenland, which is a region where 972 973 snow accumulation rates are generally higher than 130 kg m<sup>-2</sup> a<sup>-1</sup>. Since the spatial 974 variability of ice-core  $\delta^{15}N(NO_3)$  is mainly determined by the fractional loss of snow 975  $NO_3^-$  due to the large fractionation factor associated with its photolysis [Berhanu et al., 976 2014], observations of  $\delta^{15}N(NO_3)$  in snow and ice can be used to estimate both the 977 978 degree of recycling and loss of snow NO<sub>3</sub><sup>-</sup> in Antarctica and Greenland as long as this 979 condition is met. The relationship between recycling and loss can be useful for the 980 interpretation of the oxygen isotopic composition of ice-core NO<sub>3</sub><sup>-</sup> (e.g., Sofen et al. [2014]), which is impacted by recycling of snow NO3<sup>-</sup>. We note that the relationship 981 between  $\tau_{NO_3^- burial}$  and snow accumulation rate may vary in different climates depending on the concentrations of light-absorbing impurities in snow [*Geng et al.*, 2015]. 982 983

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985 This is the first modeling study to incorporate an idealized snowpack along with a snow 986 radiative transfer model into a global chemical transport model. This modeling 987 framework is used to investigate the impacts of snow-sourced  $NO_x$  on boundary layer 988 chemistry and nitrogen recycling and redistribution, and its spatial variability, across 989 Antarctica and Greenland. The modeled spatial patterns of nitrogen recycling, photolysisdriven loss of NO<sub>3</sub><sup>-</sup> from snow, and ice-core  $\delta^{15}N(NO_3)$  are each sensitive to multiple 990 991 meteorological and chemical parameters, some of which are interdependent. Model 992 sensitivity studies suggest that future field, laboratory, and modeling studies continue to 993 focus on gaining a better understanding of the quantum yield for  $NO_3^-$  photolysis and the 994 concentration of photolabile  $NO_3^-$ , which are likely related. More observations of the 995 concentration of UV light absorbing impurities in snow are also needed. Updated 996 information about the quantum yield for NO<sub>3</sub><sup>-</sup> photolysis and the concentration of 997 photolabile  $NO_3^{-1}$  in snow along with additional snow photochemical reactions can be 998 incorporated into this modeling framework in the future, which will continue to improve 999 our understanding of the impacts of snow photochemistry on boundary layer chemistry 1000 and the preservation of  $NO_3^-$  and other photochemically-active species in ice cores.

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Table 1. Glossary of variables used in this paper.		
Variable	Unit	Description
λ	nm	Wavelength
$\phi$	molec photon <sup>-1</sup>	Quantum yield for NO <sub>3</sub> <sup>-</sup> photolysis
$\sigma_{NO3}$	$cm^2$	Absorption cross-section for NO <sub>3</sub> <sup>-</sup> photolysis
Ι	photons cm <sup>-2</sup> s <sup>-1</sup> nm <sup>-1</sup>	Actinic flux of UV radiation
Ze	cm	e-folding depth of UV actinic flux in snow
Z <sub>3e</sub>	cm	Depth of snow photic zone
$\alpha_r$	kg m <sup>-2</sup> yr <sup>-1</sup>	Total annual snow accumulation rate
$C_{BC}$	$ng g^{-1}$	Annual mean snow black carbon concentration
$r_e$	μm	Radiation equivalent mean ice grain radii
$\rho_{snow}$	kg m <sup>-3</sup>	Snow density
<i>Kext</i> <sub>tot</sub>	cm <sup>-1</sup>	Bulk extinction coefficient for snow
$[NO_3]_{top}$	ng g <sup>-1</sup>	Mean NO <sub>3</sub> <sup>-</sup> concentration in top 2 cm of snow
$[NO_3]_{bot}$	ng g <sup>-1</sup>	Mean NO <sub>3</sub> <sup>-</sup> concentration from 2-cm depth to the bottom of
		the snow photic zone
EF	unitless	NO <sub>3</sub> <sup>-</sup> enhancement factor in top 2 cm of snow
$F_p$	fraction	Fraction of photolabile NO <sub>3</sub> <sup>-</sup> in snow
$[NO_3]_{bot} \ge F_p$	ng g <sup>-1</sup>	Mean NO <sub>3</sub> <sup>-</sup> concentration from 2-cm depth to the bottom of
		the snow photic zone scaled by fraction of photolabile NO <sub>3</sub>
		in snow
8	<b>‰</b>	Fractionation constant for NO <sub>3</sub> <sup>-</sup> photolysis
$\overline{F_{NOx}}$	molec cm <sup>-2</sup> s <sup>-1</sup>	Mean summer flux of snow-sourced NO <sub>x</sub>
$F_{NOx}$	ng N m <sup>-2</sup> yr <sup>-1</sup>	Annual sum of snow-sourced NO <sub>x</sub> flux
$F_{PRI}$	$ng N m^{-2} yr^{-1}$	Annual sum of primary NO <sub>3</sub> <sup>-</sup> deposited to snow
$F_R$	ng N m <sup>-2</sup> yr <sup>-1</sup>	Annual sum of recycled NO <sub>3</sub> <sup>-</sup> to snow
NRF	unitless	Metric to assess degree of nitrogen recycling in 1 year
$ au_{NO_3^-burial}$	years/months	Lifetime of NO <sub>3</sub> <sup>-</sup> against burial below snow photic zone
$\tau_{NO_3^-photolysis}$	months	Lifetime of NO <sub>3</sub> <sup>-</sup> against photolysis in snow photic zone
F <sub>p</sub> <sub>photo</sub>	unitless	Fraction of photolabile NO <sub>3</sub> <sup>-</sup> photolyzed rather than buried
<sup>p</sup> photo		below the snow photic zone
f	fraction	Fraction of photolysis-driven loss of NO <sub>3</sub> <sup>-</sup> from snow
$\delta^{15}N(NO_3)$	%0	Nitrogen isotopic composition of NO <sub>3</sub>
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1498 Table 1. Glossary of variables used in this paper.

Variable		References	
$\phi$	0.0015-0.0052 molec photon <sup>-1a</sup> (Antarctica)	Chu and Anastasio [2003]	
	0.0032-0.0069 molec photon <sup>-1a</sup> (Greenland)		
$\sigma_{NO3}$	$2.7 \times 10^{-20} \text{ cm}^2 (\lambda = 298-307 \text{ nm})$	Sander et al. [2006]	
	$2.4 \times 10^{-20} \text{ cm}^2 (\lambda = 307 - 312 \text{ nm})$		
	$1.9 \times 10^{-20} \text{ cm}^2 (\lambda = 312 - 320 \text{ nm})$		
	$2.3 \times 10^{-21} \text{ cm}^2 (\lambda = 320 - 345 \text{ nm})$		
8	-47.9‰	Berhanu et al. [2014]	
Γ <sub>e</sub>	$86.0-235.0 \ \mu\text{m}^{b}$ (Antarctica)	Carmagnola et al. [2013]	
	$73.0-211.0 \ \mu m^b$ (Greenland)	Gallet et al. [2011]	
$\rho_{snow}$	260-360 kg m <sup>-3b</sup>	Gallet et al. [2011]	
-	235-350 kg m <sup>-3b</sup>	Carmagnola et al. [2013]	
EF	6 <sup>c</sup> (Antarctica)	Dibb et al. [2004, 2007]	
	1 <sup>c</sup> (Greenland)	Erbland et al. [2013]	
		Frey et al. [2009]	
		Mayewski and Legrand [1990	
		Rothlisberger et al. [2000]	
[NO <sub>3</sub> ] <sub>bot</sub>	$60 \text{ ng g}^{-1d}$	Bertler et al. [2005]	
	132 ng g <sup>-1e</sup>	Burkhart et al. [2009]	
		Dibb et al. [2007]	
		Honrath et al. [2002]	
		Geng et al. [2014b]	

1511 Table 2. Value(s) of parameters used in the model.

1512 <sup>a</sup>Temperature-dependent equation from Chu and Anastasio [2003] used. The modeled temperatures ranged from 237-271K across Antarctica and 257-280K across Greenland. 1513 1514 <sup>b</sup>Vertical profiles of  $r_e$  range from 86 µm at the snow surface to 332 µm at 300-cm depth in Antarctica and from 73 µm at the snow surface to 211 µm at 300-cm depth in 1515 Greenland and are not varied spatially or temporally. Vertical profiles of  $\rho_{snow}$  range from 260 kg m<sup>-3</sup> at the snow surface to 360 kg m<sup>-3</sup> at 300-cm depth in Antarctica and from 235 kg m<sup>-3</sup> at the snow surface to 350 kg m<sup>-3</sup> at 300-cm depth in Greenland and are not varied 1516 1517 1518 spatially or temporally. Observations from Gallet et al. [2011] (Dome C) are used across 1519 Antarctica and from Carmagnola et al. [2013] (Summit) are used across Greenland. The 1520 deepest observed  $r_e$  and  $\rho_{snow}$  values are used for all modeled  $r_e$  and  $\rho_{snow}$  values at deeper 1521 1522 snow depths.

<sup>c</sup>Median of observed  $NO_3^-$  enhancement factors in Antarctica (1-17) and low end of observed  $NO_3^-$  enhancement factors in Greenland (1-2).

<sup>d</sup>Median of observed sub-surface snow NO<sub>3</sub><sup>-</sup> mixing ratios from the ITASE campaign [*Bertler et al.*, 2005].

<sup>e</sup>Average NO<sub>3</sub><sup>-</sup> concentrations from buried snow layer presented in Table 3 are used.

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Table 3. Dependence of mean austral summer (DJF) flux of snow-sourced NO<sub>x</sub> ( $\overline{F_{NOx}}$ ) in Antarctica on quantum yield ( $\phi$ ), the fraction of photolabile NO<sub>3</sub><sup>-</sup> ( $F_p$ ), snow NO<sub>3</sub><sup>-</sup> concentrations below 2 cm ( $[NO_3^-]_{bot}$ ), the radiation equivalent ice grain radius ( $r_e$ ), the bulk snow extinction coefficient (*Kext<sub>tot</sub>*), the NO<sub>3</sub><sup>-</sup> concentration enhancement factor in the top 2 cm (*EF*), and snow black carbon concentration.

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Parameter	Base case values	Values used in sensitivity	$\overline{F_{NOx}}$ range in
	(where $\overline{F_{NOx}} = 0.5 - 7.8 \times 10^8$	studies	sensitivity studies
	molec $\text{cm}^{-2} \text{ s}^{-1}$ )		$(x10^8 \text{ molec cm}^2 \text{ s}^{-1})$
			(Factor changes
			compared to $\overline{F_{NOx}}$ in
			base case, unitless)
Quantum yield	0.002 molec photon <sup>-1 a</sup>	0.6 molec photon <sup>-1</sup>	5-2600
( <i>φ</i> )			(10.0-333.0)
Fraction of photolabile	0.01-0.99	Set to 1 everywhere	3.7-9.6
$NO_3(F_p)$	(spatial variation, Figure 3c)		(1.2-7.4)
Sub-surface snow NO <sub>3</sub>	60.0 ng g <sup>-1b</sup>	$30-120 \text{ ng g}^{-1}$	0.3-15.8
$([NO_3]_{bot})$			(0.6-2.0)
Radiation equivalent	Jan: 332.0 μm <sup>c</sup>	Study 1: 332.0 μm <sup>d</sup>	0.5-10.2
mean ice grain radii $(r_e)$	Dec-Feb: 198-332.0 μm <sup>c</sup>	Study 2: 198-332.0 µm <sup>d</sup>	(1.0-1.3)
	Mar-Nov: 86.0-332.0 μm <sup>c</sup>	Study 3: 86.0-332.0 µm <sup>d</sup>	
Bulk snow extinction	$1.7-6.9 \times 10^3 \text{ m}^{-1}$	$\pm 20\%$ with respect to	0.5-9.4
coefficient ( <i>Kext<sub>tot</sub></i> )	(spatial variation)	base case values	(1.0-1.2)
NO <sub>3</sub> <sup>-</sup> enhancement	6.0 <sup>e</sup>	1-10	0.5-9.3
factor in top 2 cm (EF)			(1.0-1.2)
Snow black carbon	$0.08-0.6 \text{ ng g}^{-1}$	$\pm$ factor of 2 with respect	0.5-8.6
$(C_{BC})$	(spatial variation, Figure 3b)	to base case values	(1.0-1.1)

- 1540 <sup>a</sup>from Chu and Anastasio [2003] at T=244K
- <sup>b</sup>median of ITASE campaign [*Bertler et al.*, 2005]

1542 <sup>c</sup>r<sub>e</sub> is varied vertically and temporally, but uniformly across Antarctica based on Gallet et

- al. [2011] and Klein [2014]. In January,  $r_e$  is constant with depth (332 µm), in December and February,  $r_e$  ranges from 198 µm at the snow surface to 332 µm at 300 cm depth, and
- 1545 from March to November,  $r_e$  ranges from 86 µm at the surface to 360 µm at 300 cm 1546 depth.
- <sup>d</sup> in  $r_e$  sensitivity study 1, the base-case 'January'  $r_e$  profile is applied for every month. In r<sub>e</sub> sensitivity study 2, the base-case 'December and February'  $r_e$  profile is applied for every month. In  $r_e$  sensitivity study 3, the base-case 'March-November'  $r_e$  profile is applied for every month.
- <sup>e</sup>median of observed *EF* [*Dibb et al.*, 2004, *Frey et al.*, 2009, *Mayewski and Legrand*,
- 1552 1990, Rothlisberger et al., 2000].
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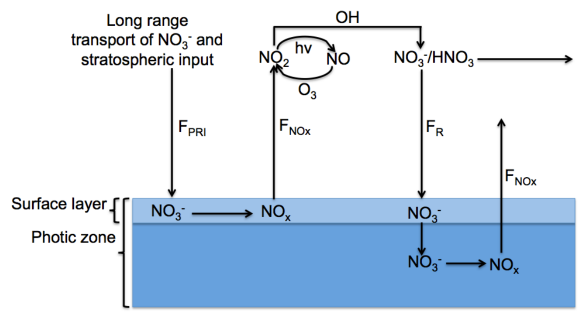


Figure 1. Schematic showing the nitrogen recycling associated with NO<sub>3</sub><sup>-</sup> photolysis as included in the model.  $F_{PRI}$  (ng N m<sup>-2</sup> yr<sup>-1</sup>) is the downward, primary flux of NO<sub>3</sub><sup>-</sup> to Antarctica and Greenland originating from long-range transport and the stratosphere,  $F_{NOx}$  (ng N m<sup>-2</sup> yr<sup>-1</sup>) is the upward flux of snow-sourced NO<sub>x</sub> to the boundary layer, and  $F_R$  (ng N m<sup>-2</sup> yr<sup>-1</sup>) is downward, recycled flux of HNO<sub>3</sub> to the snow surface. The surface snow layer (top 2 cm) is distinguished from the rest of the photic zone because 30-65% of snow-sourced NO<sub>x</sub> is produced in the top 2 cm of snowpack [Zatko et al., 2013], and because both NO<sub>3</sub><sup>-</sup> concentrations and actinic flux are much higher in the top surface layer compared to deeper layers.

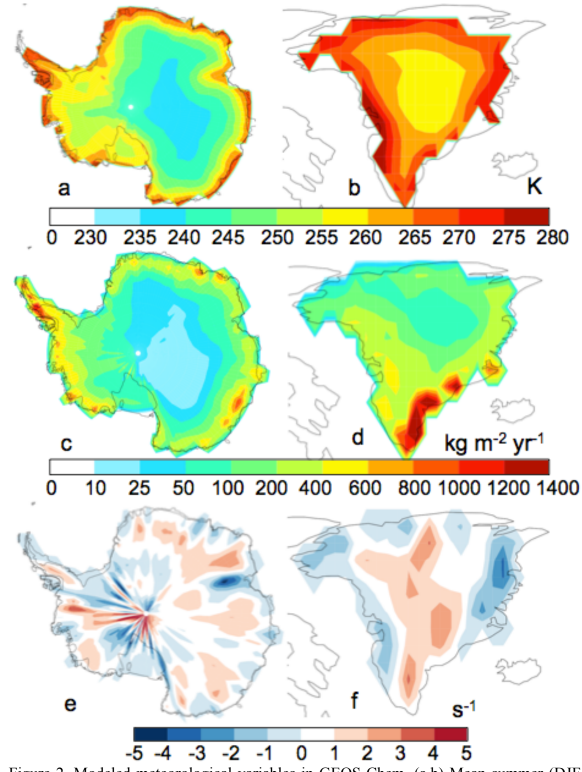
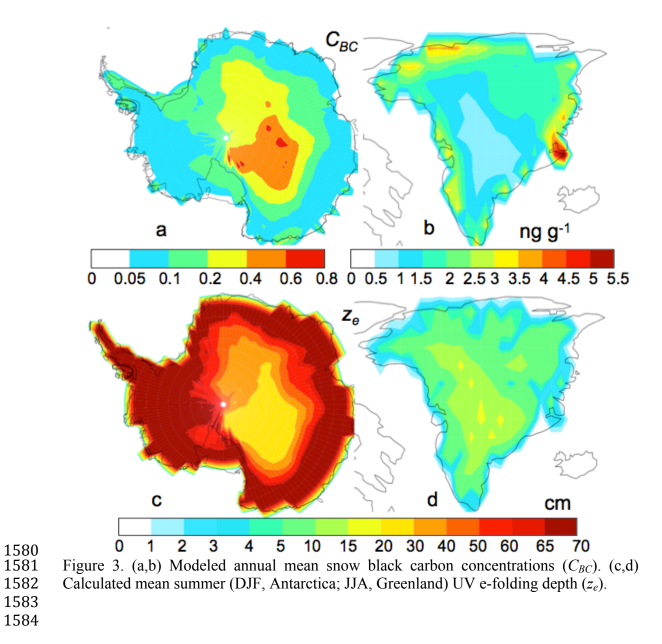
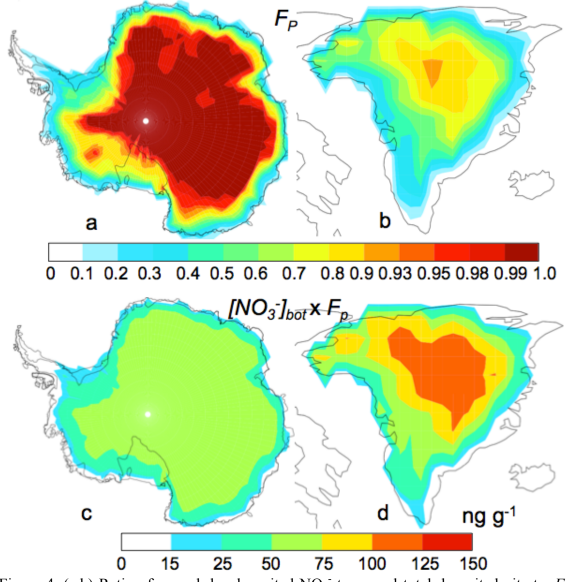


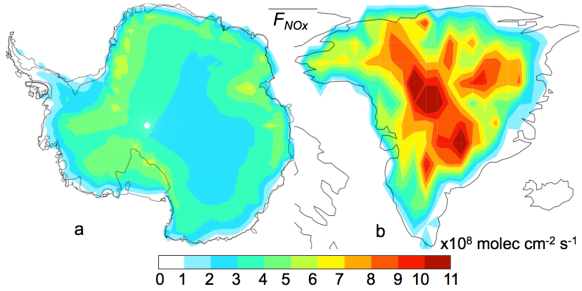


Figure 2. Modeled meteorological variables in GEOS-Chem. (a,b) Mean summer (DJF, Antarctica; JJA, Greenland) air temperature in lowest vertical grid box. (c,d) Annual 1576 total snow accumulation rate (kg m<sup>-2</sup> yr<sup>-1</sup>) from May 2009 to May 2010. (e,f) Annual mean surface wind divergence (s<sup>-1</sup>) from May 2009 to May 2010. Blue regions indicate 1577 1578 1579 regions of convergence.

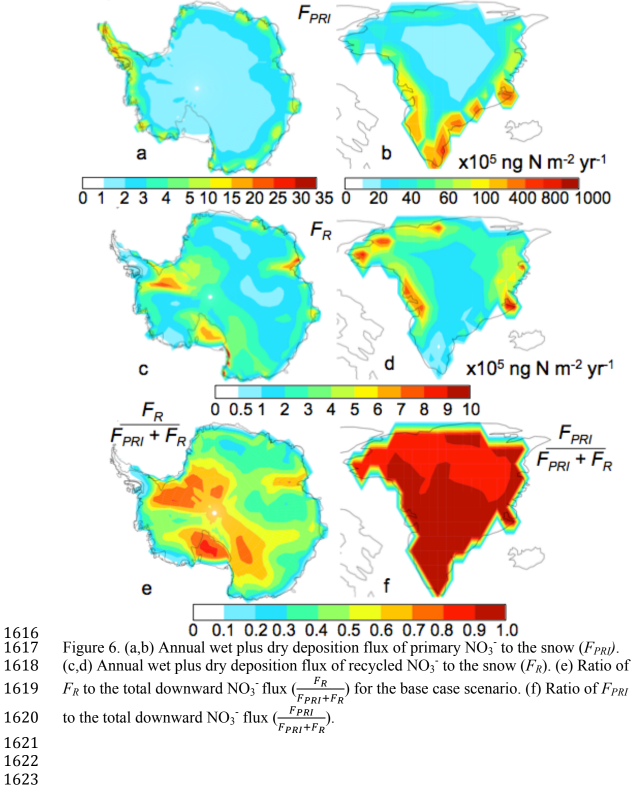


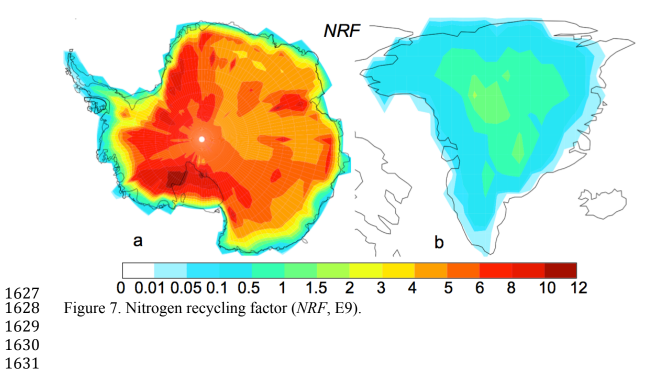


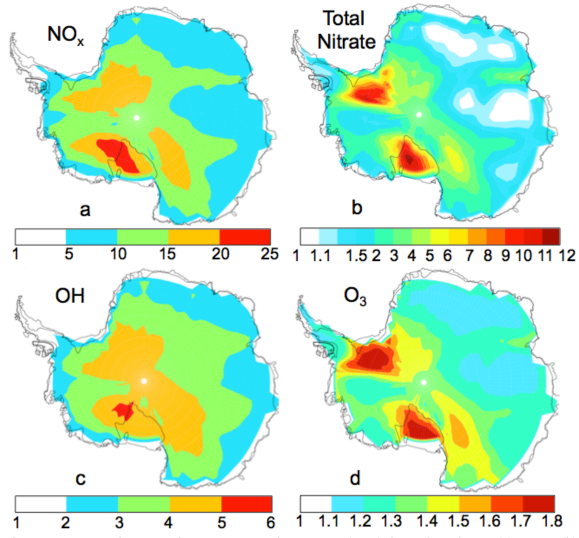
1585 0 15 25 50 75 100 125 150 1586 Figure 4. (a,b) Ratio of annual dry-deposited  $NO_3^-$  to annual total deposited nitrate,  $F_P$ . (c,d) Annual sub-surface snow  $NO_3^-$  concentrations ( $[NO_3^-]_{bot} \ge F_p$ ) used in the model. 



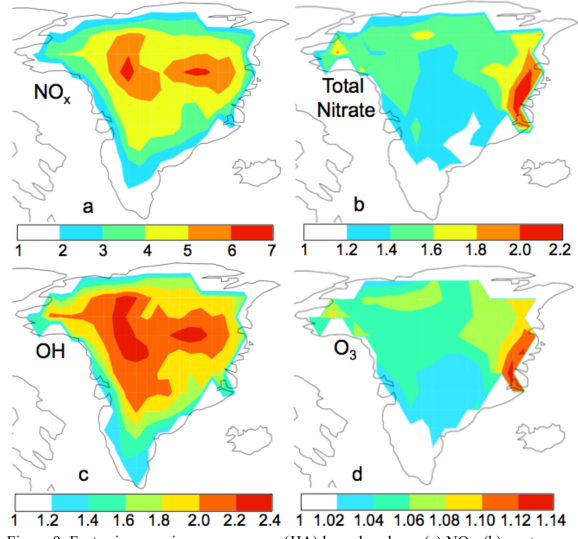
1593 Figure 5. Mean summer (DJF, Antarctica; JJA, Greenland) flux of snow-sourced NO<sub>x</sub> from the snow  $(\overline{F_{NOx}})$ .



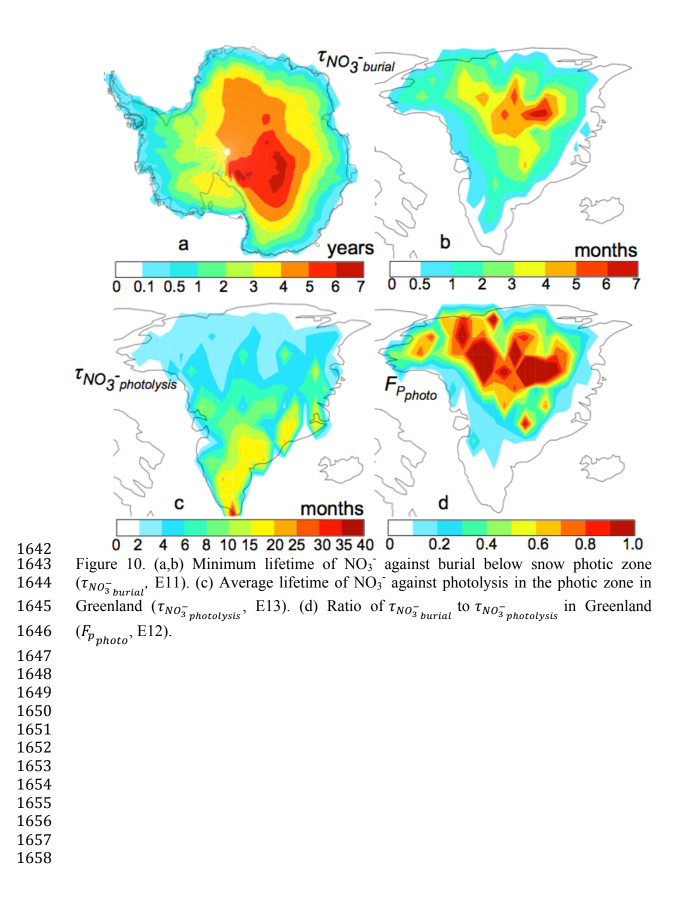


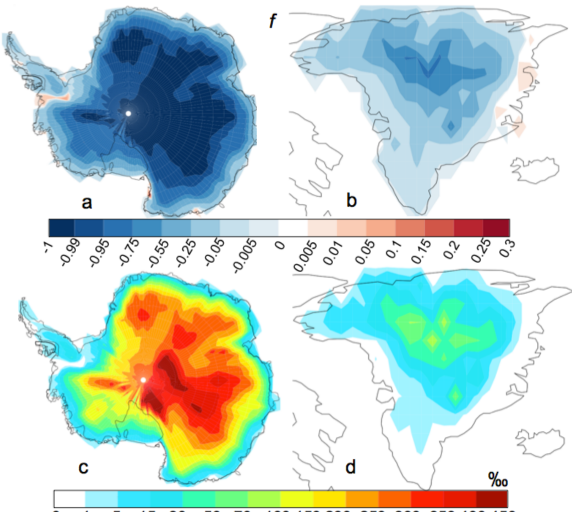


1 2 3 4 5 6 1 1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 Figure 8. Factor increase in mean austral summer (DJF) boundary layer (a) NO<sub>x</sub>, (b) gas+aerosol phase NO<sub>3</sub>, (c) OH, and (d) O<sub>3</sub> mixing ratios between model runs with  $F_{NOx}$ to mpared to without  $F_{NOx}$ .

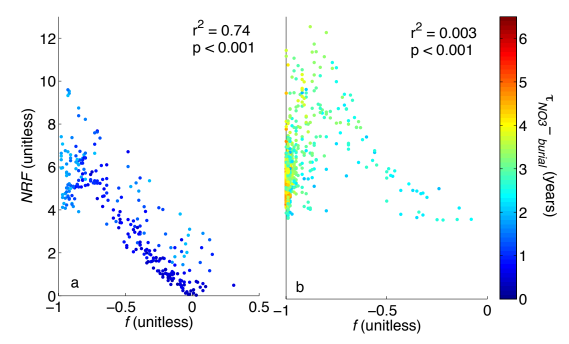


1 1.2 1.4 1.6 1.8 2.0 2.2 2.4 1 1.02 1.04 1.06 1.08 1.10 1.12 1.14 Figure 9. Factor increase in mean summer (JJA) boundary layer (a) NO<sub>x</sub>, (b) gas+aerosol phase nitrate, (c) OH, and (d) O<sub>3</sub> mixing ratios between model runs with  $F_{NOx}$  compared to without  $F_{NOx}$ .



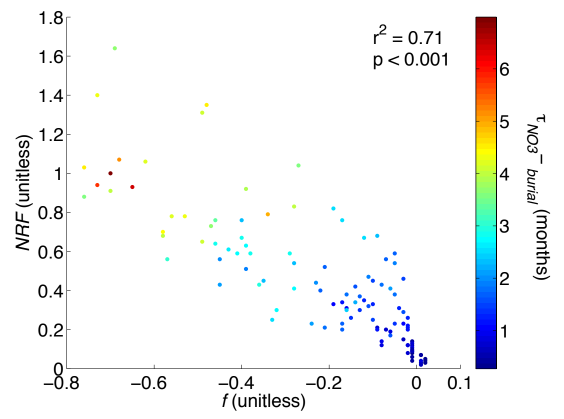


1659 0 1 5 15 30 50 70 100 150 200 250 300 350 400 450 1660 Figure 11. (a,b) Fraction of NO<sub>3</sub><sup>-</sup> gained (positive values) or lost (negative values) from 1661 the snow through photolysis (f, E10). (c,d) Modeled enrichment in ice-core  $\delta^{15}N(NO_3^{-})$ 1662 (E14) due to photolysis-driven loss of NO<sub>3</sub><sup>-</sup> in snow. Note the varied color bar scale in 1663 Figures 11a and 11b.



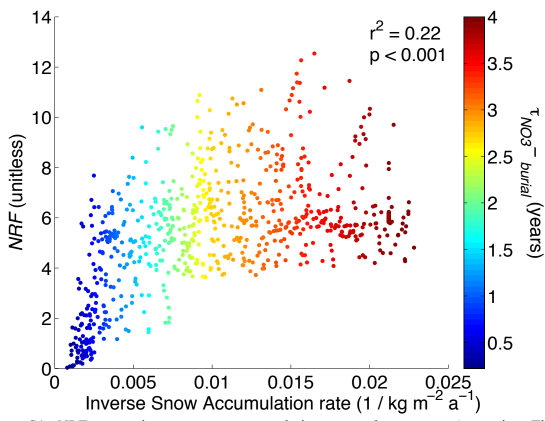


1680 Figure 12. NRF versus f values across Antarctica. (a) Regions where  $NO_3^-$  remains in the photic zone for 2 years or less. (b) Regions where  $NO_3^-$  remains in the photic zone for more than 2 years. The color scale represents the number of years NO<sub>3</sub><sup>-</sup> remains in the photic zone  $(\tau_{NO_3 burial})$ . 



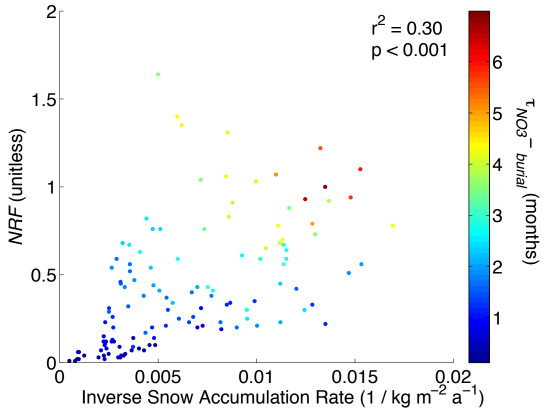
1688 Figure 13. *NRF* versus *f* values across Greenland. The color scale represents the minimum number of months that NO<sub>3</sub><sup>-</sup> remains in the photic zone  $(\tau_{NO_3^-burial})$ . 





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Figure S1. NRF versus inverse snow accumulation rate values across Antarctica. The color scale represents the minimum number of years that NO<sub>3</sub><sup>-</sup> remains in the photic zone  $(\tau_{NO_3^-burial}).$ 



1719 1720 Figure S2. *NRF* versus inverse snow accumulation Rate (1 / kg m<sup>-</sup> a<sup>-</sup>) 1720 Figure S2. *NRF* versus inverse snow accumulation rate values across Greenland. The 1721 color scale represents the minimum number of months that NO<sub>3</sub><sup>-</sup> remains in the photic 1722 zone ( $\tau_{NO_3^- burial}$ ).

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