# Response to the referees

Again, we thank Prof. Joel Savarino (the handling editor) and the anonymous reviewer for their time in reviewing our revised manuscript. Below, we give a point-by-point response to the comments and suggestions of the reviewers, in the order of (1) comments from Referees, (2) author's response, and (3) author's changes in manuscript (referee comments in black; author's response and changes in manuscript in blue).

## The Reviewer #1

We thank the reviewer very much for his/her thoughtful review of our revised manuscript. All of the comments and suggestions have been taken into account, and are included in the revised version. Below, we give a point-by-point response to the comments and suggestions of the reviewer, in the order of (1) comments from Referees, (2) author's response, and (3) author's changes in manuscript (referee comments in black; author's response and changes in manuscript in blue).

#### (1) comments from Referees

The authors are to be commended for taking all reviewer comments seriously and implementing significant revisions to the manuscript. From my point of view I'd be happy to recommend publication after addressing the points below and carefully rechecking grammar/spelling:

## (1) author's response

Again, we greatly appreciate the reviewer for his/her time in reviewing the revised version of the manuscript. We have carefully read the whole manuscript and improved grammar and spelling.

## (1) author's changes in manuscript

Following the reviewer's comments, we carefully read the manuscript. Please see the revised version.

## (2) comments from Referees

- please add to the introduction & discussion (4.2) the Halley study, something like:

Significant concentrations of organic nitrates (PAN and alkyl nitrates) were observed in the lower atmosphere at Halley in coastal Antarctica consistent with an oceanic source (Jones et al., 2011). They dominated the NOy budget during the winter, and were on a par with inorganic nitrate compounds during the summer. Although not a direct source of snowpack nitrate, organic nitrates would act as a source of NOx to coastal Antarctica that would ultimately contribute to nitrate within the snowpack (Jones et al., 2011). However, multi-seasonal measurements of surface snow nitrate correlate strongly with inorganic NOy species (especially HNO3) rather than organic (Jones et al., 2011).

## (2) author's response

We thank the reviewer for the suggestion. The inorganic  $NO_3^-$  plays an important role in the atmospheric  $NO_v$  budget.

## (2) author's changes in manuscript

Following the reviewer's suggestion, in the section of introduction, the statement was added, as follows,

At Halley station in coastal Antarctica, significant concentrations of organic nitrates (peroxyacetyl nitrate (PAN) and alkyl  $NO_3^-$ ) were observed in the lower atmosphere (Jones et al., 2011). Organic nitrates dominated the  $NO_y$  (sum of reactive nitrogen oxide compounds) budget during the winter, and were on a par with inorganic nitrate compounds during the summer. Although not a direct source of snowpack  $NO_3^-$ , organic nitrates could act as source of  $NO_x$  to coastal Antarctica that would ultimately contribute to  $NO_3^-$  within the snowpack (Jones et al., 2011).

As the reviewer suggested, this point was also included in Discussion (4.2), as follows,

Although the organic nitrates can play an important role in the atmospheric  $NO_y$  budget, multi-seasonal measurements of surface snow  $NO_3^-$  correlate strongly with inorganic  $NO_y$  species (especially HNO<sub>3</sub>) rather than organic (Jones et al., 2011).

For the changes, please see the revision-tracked version of manuscript, sections **1. Introduction**, and **4.2 Effects of coexisting ions on NO**<sub>3</sub><sup>-</sup>

#### (3) comments from Referees

- It is great that you include now the atmospheric observations. However, in my view they need to go into the main manuscript not just the supplementary material; e.g. add a panel to Fig.1 showing the variation with distance from the coast and make corresponding amendments to methods, results and discussion. Table S1 (move to main manuscrpt) must also include columns with site ID and sampling day.

As I pointed out before atmospheric NO3- is key to discuss and interpret an air-snow study of nitrate in snow. In fact, this is probably the only Antarctic traverse, which produced HiVol filter samples of atmospheric NO3-, very relevant to the entire discussion in this manuscript.

## (3) author's response

We agree with the reviewer. The sampling and analysis methods of atmospheric  $NO_3^-$  were included in the section of methodology. The section of result and Figure 1 were revised accordingly. The full information about sampling location, sampling data, sample ID and chemical ion concentrations ( $NO_3^-$  and  $SO_4^{2-}$ ) is present in Table S1 in supporting information.

## (3) author's changes in manuscript

Following the reviewer's suggestion, the sampling method for atmospheric  $NO_3^-$  was included in section **2.2 Sample collection**, as follows,

To support understanding of the air-snow transfer of  $NO_3^-$  on the traverse, atmospheric  $NO_3^-$  was collected on glass fiber filters (Whatman G653) using a high volume air sampler (HVAS), with a flow rate of ~1.0 m<sup>3</sup> min<sup>-1</sup> for 12-15 hr, during the inland traverse campaign in 2015/2016. The  $NO_3^-$  collected on glass fiber filters are expected to equal the sum of particulate  $NO_3^-$  and gaseous HNO<sub>3</sub>, based upon previous investigations in East Antarctica (Savarino et al., 2007; Frey et al., 2009; Erbland et al., 2013). In total, 34 atmospheric samples were collected on the traverse. In addition, two field blanks were collected from filters installed in the HVAS without pumping and treated as samples thereafter. Detailed information about the atmospheric sampling is presented in Table S1 in supporting information.

After sample collection, all filters and snow samples were sealed in clean PE bags and preserved in clean thermal insulated boxes. All of the samples were transported to the laboratory under freezing conditions (< -20 °C).

Details on the analytical processing of atmospheric  $NO_3^-$  samples were added in **2.3 Sample** analysis, as follows,

In the laboratory, three quarters of individual filters were cut into pieces using pre-cleaned scissors that were rinsed between samples, placed in ~100 ml of Milli-Q water, ultrasonicated for 40 min and leached for 24 hr under shaking. The sample solutions were then filtered through 0.22  $\mu$ m ANPEL PTFE filters for NO<sub>3</sub><sup>-</sup> concentration.

Accordingly, Figure 1 in the manuscript was revised, as follows, and the main results of the atmospheric  $NO_3^-$  investigation was included in section **3.1**.



**Figure 1.** Concentrations of  $NO_3^-$  in snow (surface snow, crystal ice and snowpits; on the primary *y*-axis) and atmosphere (on the secondary *y*-axis), with error bars representing one standard deviation of  $NO_3^-$  (1 $\sigma$ ) for individual snowpits. Also shown is the annual snow accumulation rate on the traverse (red solid line; based on Ding et al. (2011)). Note that  $NO_3^-$  concentration in one crystal ice sample (red dot), is higher than the maximum value of the primary *y*-axis ( $NO_3^-$  concentration = 16.7 µeq L<sup>-1</sup> in the parentheses).

## (4) comments from Referees

177-80 weird phrasing. Better: The late winter/early secondary maximum of nitrate observed in surface snow at coastal and inland locations has been attributed to the stratospheric source based on the nitrate stable isotopic composition (Legrand, 1989; Savarino, 2007; Frey, 2009).

## (4) author's response

We thank the reviewer very much.

## (4) author's changes in manuscript

Following the suggestion of the reviewer, the sentence was rephrased, as follows,

The late winter/early spring secondary maximum of  $NO_3^-$  observed in the atmosphere at coastal and inland locations has been attributed to the stratospheric source based on the  $NO_3^-$  stable isotopic composition (Legrand et al., 1989; Savarino et al., 2007; Frey et al., 2009).

## (5) comments from Referees

184-90 I think you really need to distinguish between no3- spikes and no3- variablitly on decadal

to centennial or millennial time scales. A statistically significant link between the former and SPEs (solar proton events) has now been refuted. However, a link between the long-term variability of no3- and solar cycles as suggested in Traversi et al. (2012) is very different in terms of time scales and likelyhood of physical processes to be aligned, and may be present at some locations.

## (5) author's response

We agree with the reviewer and thanks for the comment.

#### (5) author's changes in manuscript

Following the reviewer's comment, this paragraph was rephrased, as follows,

In addition, while some studies suggested that snow/ice  $NO_3^-$  is possibly linked with extraterrestrial fluxes of energetic particles and solar irradiation, with solar flares corresponding to  $NO_3^-$  spikes (Zeller et al., 1986; Smart et al., 2014), other observations and recent modeling studies have established that there is not a clear connection between solar variability and  $NO_3^-$  concentrations (Legrand et al., 1989; Legrand and Kirchner, 1990; Wolff et al., 2008; Wolff et al., 2012; Duderstadt et al., 2014; Duderstadt et al., 2016; Wolff et al., 2016). However, the potential link between the long-term (e.g., centennial to millennial time scales) variability of  $NO_3^-$  and solar cycles may be present at some locations (Traversi et al., 2012).

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## End of the responses.

1	Nitrate deposition and preservation in the snowpack along a traverse
2	from coast to the ice sheet summit (Dome A) in East Antarctica
3	
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22 Abstract. Antarctic ice core nitrate (NO<sub>3</sub>) can provide a unique record of the atmospheric reactive 23 nitrogen cycle. However, the factors influencing the deposition and preservation of NO<sub>3</sub><sup>-</sup> at the ice sheet 24 surface must first be understood. Therefore, an intensive program of snow sample collections and 25 atmospheric sampling was made on a traverse from the coast to the ice sheet summit, Dome A, East 26 Antarctica. Snow samples in this observation include 120 surface snow samples (top ~3\_cm), 20 27 snowpits with depths of 150 to 300\_cm, and 6 crystal ice samples (the topmost needle like layer on 28 Dome A plateau). The main purpose of this investigation is to characterize the distribution pattern and 29 preservation of NO<sub>3</sub><sup>-</sup> concentrations in the snow in different environments. Results show that an increasing trend of NO3<sup>-</sup> concentrations with distance inland is present in surface snow, and NO3<sup>-</sup> is 30 31 extremely enriched in the crystal ice (with a maximum of 16.1  $\mu$ eq L<sup>-1</sup>). NO<sub>3</sub><sup>-</sup> concentration profiles for 32 snowpits vary between coastal and inland sites. On the coast, the deposited  $NO_3^-$  was largely preserved, 33 and the archived  $NO_3^-$  fluxes are dominated by snow accumulation. The relationship between the 34 archived NO3<sup>-</sup> and snow accumulation rate can be well depicted by a linear model, suggesting a homogeneity of atmospheric NO<sub>3</sub><sup>-</sup> levels. It is estimated that dry deposition contributes 27-44 % of the 35 archived NO<sub>3</sub><sup>-</sup> fluxes, and the dry deposition velocity and scavenging ratio for NO<sub>3</sub><sup>-</sup> was relatively 36 37 constant near the coast. Compared to the coast, the inland snow shows a relatively weak correlation 38 between archived NO<sub>3</sub><sup>-</sup> and snow accumulation, and the archived NO<sub>3</sub><sup>-</sup> fluxes were more concentration dependent. The relationship between NO3<sup>-</sup> and coexisting ions (nssSO4<sup>2-</sup>, Na<sup>+</sup> and Cl<sup>-</sup>) was also 39 investigated, and the results show a correlation between  $nssSO_4^{-2}$  (fine aerosol particles) and  $NO_3^{-1}$  in 40 41 surface snow, while the correlation between NO3<sup>-</sup> and Na<sup>+</sup> (mainly associated with coarse aerosol 42 particles) is not significant. In inland snow, there were no significant relationships found between NO<sub>3</sub> 43 and the coexisting ions, suggesting a dominant role of NO3 recycling in determining the 44 concentrations. 45

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#### 48 1 Introduction

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As the major sink of atmospheric nitrogen oxides ( $NO_x = NO$  and  $NO_2$ ), nitrate ( $NO_3^-$ ) is one of the major chemical species measured in polar snow and ice. The measurements of  $NO_3^-$  in ice cores may offer potential for understanding the complex atmospheric nitrogen cycle as well as oxidative capacity of the atmosphere through time (Legrand and Mayewski, 1997; Alexander et al., 2004; Hastings et al., 2009; Geng et al., 2017). However, the sources, transported pathways, and preservation of  $NO_3^-$  in Antarctic snowpack are still not well understood-in Antarctic snowpack, hampering the interpretation of ice core  $NO_3^-$  records.

57 The accumulation of NO<sub>3</sub><sup>-</sup> in snow is associated with various environmental factors and continental, 58 tropospheric and stratospheric sources could influence NO<sub>3</sub> concentrations (Legrand and Kirchner, 59 1990; McCabe et al., 2007; Wolff et al., 2008; Lee et al., 2014). In surface snow, NO<sub>3</sub><sup>-</sup> levels are 60 thought to be linked with snow accumulation rate, and higher values are usually present in areas with 61 low accumulation, e.g., East Antarctic plateaus (Qin et al., 1992; Erbland et al., 2013; Traversi et al., 62 2017). Unlike sea salt related ions (e.g., chloride (Cl<sup>+</sup>), sodium (Na<sup>+</sup>), and occasionally sulfate (SO<sub>4</sub><sup>2-</sup>)), 63 NO<sub>3</sub><sup>-</sup> does not usually show an elevated level in coastal Antarctic snow (Mulvaney and Wolff, 1994; 64 Bertler et al., 2005; Frey et al., 2009), suggesting a negligible contribution from sea salt aerosols. 65 However, the marine emissions of alkyl NO3, particularly methyl and ethyl NO3, produced in surface 66 oceans by microbiological and/or photochemical processes, are thought to be a possible contribution to 67 Antarctic NO<sub>3</sub><sup>-</sup> (Jones et al., 1999; Liss et al., 2004). At Halley station in coastal Antarctica, significant 68 concentrations of organic nitrates (peroxyacetyl nitrate (PAN) and alkyl NO<sub>3</sub><sup>-</sup>) were observed in the 69 lower atmosphere (Jones et al., 2011). Organic nitrates dominated the NO<sub>v</sub> (sum of reactive nitrogen 70 oxide compounds) budget during the winter, and were on par with inorganic nitrate compounds during 71 the summer. Although not a direct source of snowpack nitrate, organic nitrates could act as source of 72 NO, to coastal Antarctica that would ultimately contribute to NO<sub>3</sub> within the snowpack (Jones et al., 73 2011).

74 While industrial and/or agricultural emissions have contributed to increasing NO<sub>3</sub><sup>-</sup> levels in 75 Greenland snow and ice over recent decades to hundreds of years, the anthropogenic contribution to 76 Antarctic NO3<sup>-</sup> is less clear (Mayewski and Legrand, 1990; Hastings et al., 2009; Felix and Elliott, 77 2013; Geng et al., 2014). Lightning and NO<sub>x</sub> produced in the lower stratosphere have long been 78 thought to play a major role (Legrand et al., 1989; Legrand and Kirchner, 1990). Recently, adjoint 79 model simulations proposed that tropospheric transport of  $NO_3^-$  from mid-low latitude  $NO_x$  sources is 80 an important source to the Antarctice year round, though less so in austral spring/summer (Lee et al., 81 2014). A recent treatment of NO3<sup>-</sup> in snow in the same global chemical transport model suggests that 82 the recycling of NO<sub>3</sub><sup>-</sup> and/or transport of NO<sub>x</sub> due to photolysis of NO<sub>3</sub><sup>-</sup> in the surface snow layer is 83 important in determining summertime concentrations (Zatko et al., 2016). The stratospheric inputs of 84  $NO_3^-$  are thought to be the result of from N<sub>2</sub>O oxidation to NO, then and formation of NO<sub>3</sub><sup>-</sup> that is 85 deposited via polar stratospheric cloud (PSC) sedimentation (Legrand et al., 1989; Legrand and 86 Kirchner, 1990). The late winter/early spring secondary maximum of NO3<sup>-</sup> observed in the atmosphere 87 at coastal and inland locations has been attributed to the stratospheric source based on the NO3 stable 88 isotopic composition This source has been used to explain sporadic NO3 -concentration peaks and its 89 associated isotopic composition in later winter and/or early spring at both coastal and inland locations 90 (Legrand et al., 1989; Savarino et al., 2007; Frey et al., 2009). At some sites, the snow/ice core NO<sub>3</sub>

91 concentrations were found to be linked with regional atmospheric circulation (e.g., sea level pressure

92 gradient; Goodwin et al., 2003; Russell et al., 2006). In general, atmospheric circulation appears not to 93 affect snow NO<sub>3</sub><sup>-</sup> concentrations directly, but indirectly through an influence on the air mass transport 94 and/or snow accumulation rate (Russell et al., 2004; Russell et al., 2006). In addition, while some 95 studies suggested that snow/ice NO3<sup>-</sup> is possibly linked with extraterrestrial fluxes of energetic particles 96 and solar irradiation, with solar flares corresponding to NO3<sup>-</sup> spikes (Zeller et al., 1986; Smart et al., 97 2014), other observations and recent modeling studies have established that there is not a clear 98 connection between solar variability and NO3<sup>-</sup> concentrations (Legrand et al., 1989; Legrand and 99 Kirchner, 1990; Wolff et al., 2008; Wolff et al., 2012; Duderstadt et al., 2014; Duderstadt et al., 2016; 100 Wolff et al., 2016). However, the potential link between the long-term (e.g., centennial to millennial 101 time scales) variability of NO<sub>3</sub> and solar cycles may be present at some locations (Traversi et al., 2012). 102 In summary, factors influencing  $NO_3^-$  levels in snow/ice are complicated, and the significance of the 103 relationship between NO<sub>3</sub><sup>-</sup> and controlling factors varies temporally and spatially.

104 Gas phase and snow concentration studies, and recent isotopic investigations and modeling have 105 shown that  $NO_3$ , particularly in snow on the Antarctic plateau, is a combination of deposition of HNO<sub>3</sub> 106 and post-depositional loss or recycling of NO3<sup>-</sup> (e.g., R öthlisberger et al., 2002; Davis et al., 2004; 107 Dibb et al., 2004; Erbland et al., 2013; Erbland et al., 2015; Shi et al., 2015; Bock et al., 2016; Zatko et 108 al., 2016). Based upon a suite of isotopic studies in the field and laboratory, it has been demonstrated 109 that under cold, sunlit conditions ultraviolet photolysis dominates NO<sub>3</sub><sup>-</sup> post-depositional processing, 110 whereas HNO3 volatilization may become more important at warmer temperatures > -20 °C 111 (Röthlisberger et al., 2002; Frey et al., 2009; Erbland et al., 2013; Berhanu et al., 2015). In snowpack, 112 the solar radiation decreases exponentially, with attenuation described in terms of an e-folding depth  $(z_e)$ 113 where the actinic flux is reduced to 37 % (i.e., 1/e) of the surface value. Thus, about 95 % of snowpack 114 photochemistry is expected to occur above the depth of three times  $z_e$  (Warren et al., 2006). Field measurements at Dome C on the East Antarctic plateau suggest a  $z_e$  of 10 to 20 cm (France et al., 2011), 115 116 and the depth is dependent upon the concentration of impurities contained in the snow (Zatko et al., 117 2013). In the inland regions with low snow accumulation rates, particularly on the East Antarctic 118 plateaus, photolysis has been shown to lead to significant post-depositional loss of NO<sub>3</sub>, demonstrated by significant enrichment in <sup>15</sup>N of snow NO<sub>3</sub><sup>-</sup> (i.e., high  $\delta^{15}$ N) (Frey et al., 2009; Erbland et al., 2013; 119 Berhanu et al., 2015; Erbland et al., 2015; Shi et al., 2015), as well as a decrease in  $\delta^{18}$ O and  $\Delta^{17}$ O due 120 to reformation of NO3<sup>-</sup> in the condensed phase (Erbland et al., 2013; Shi et al., 2015 and references 121 122 therein). The transport and recycling of NO<sub>x</sub> sourced from photolysis of snow NO<sub>3</sub><sup>-</sup> in the summertime 123 has been invoked to model the distribution of snowpack NO3<sup>-</sup> across the Antarctic plateau (Zatko et al., 124 2016). However, snow physical characteristics play a crucial role in  $NO_3^-$  deposition and preservation. 125 For instance, summertime concentrations in the surface skin layer of snow (the uppermost ~4 mm) can be explained as the result of co-condensation of HNO3 and water vapour, with little to no photolytic 126 127 loss in this microlaver (Bock et al., 2016). The combination of concentration and isotopic studies, along 128 with physical aspects of the snow, could lead to the reconstruction and interpretation of atmospheric 129 NO<sub>3</sub> over time (e.g., Erbland et al., 2015; Bock et al., 2016), if there is detailed understanding of the 130 NO3<sup>-</sup> deposition and preservation in different environments in Antarctica.

The effects of volatilization of  $NO_3^-$  are uncertain, given that one field experiment suggests that this process is an active player in  $NO_3^-$  loss (17 % (-30 °C) to 67 % (-10 °C) of  $NO_3^-$  lost after two weeks' physical release experiments; Erbland et al., 2013), while other laboratory and field studies show that volatilization plays a negligible role in  $NO_3^-$  loss (Berhanu et al., 2014; Berhanu et al., 2015). Further investigations are needed to quantify the effects of volatilization for a better understanding of  $NO_3^-$  136 preservation in the snow/ice. Based on  $z_e$ , NO<sub>3</sub><sup>-</sup> at deeper depths in Antarctic snow (e.g., > 100 cm), 137 well beyond the snow photic zone, may be taken as the archived fraction. Thus, NO<sub>3</sub><sup>-</sup> in deeper snow 138 possibly provides an opportunity to investigate the archived fraction and potential influencing factors 139 (e.g., snow accumulation rate). Given that an extensive array of ice core measurements is unavailable 140 in most of Antarctica, the deeper snowpits (with depth > 100 cm) may offer a useful way to investigate 141 the archived NO<sub>3</sub><sup>-</sup>.

142 In the atmosphere in Antarctica, particularly during spring and summer, NO<sub>3</sub><sup>-</sup> is found to be mainly 143 in the form of gas phase  $HNO_3$ , with  $NO_3^-$  concentration several times higher in gas phase than in the particulate phase (Piel et al., 2006; Legrand et al., 2017b; Traversi et al., 2017). During-the 144 post-depositional processes, the uptake of gaseous HNO<sub>3</sub> is thought to be important in NO<sub>3</sub> 145 146 concentrations in surface snow layers (Udisti et al., 2004; Traversi et al., 2014; Traversi et al., 2017). 147 Due to the high concentration in summer, HNO<sub>3</sub> appears to play an important role in acidifying sea-salt 148 particles, possibly accounting for the presence of NO3<sup>-</sup> in the particulate phase in summer (Jourdain and 149 Legrand, 2002; Legrand et al., 2017b; Traversi et al., 2017). It is noted that the significant increase of 150 NO<sub>3</sub><sup>-</sup> during the cold periods (e.g., Last Glacial Maximum) could be associated with its attachment to 151 dust aerosol, instead of the formation of gas phase HNO<sub>3</sub> (Legrand et al., 1999; Wolff et al., 2010).

152 To date, investigations on spatial and temporal patterns of snow NO<sub>3</sub><sup>-</sup> have been performed on 153 several traverses in Antarctica (e.g., 1990 International Trans-Antarctica Expedition, and DDU to 154 Dome C; Qin et al., 1992; Bertler et al., 2005; Frey et al., 2009; Erbland et al., 2013; Pasteris et al., 155 2014), but these provide an uneven distribution of snow NO<sub>3</sub><sup>-</sup> concentrations, and leave<del>leaving</del> large 156 regions un-sampled (e.g., Lambert Glacier basin and Dome A plateau). Over the past few decades, 157 while several glaciological observations have been carried out on the Chinese inland Antarctic traverse 158 route from Zhongshan to Dome A, East Antarctica (Hou et al., 2007; Ding et al., 2010; Ma et al., 2010; 159 Ding et al., 2011; Li et al., 2013; Shi et al., 2015), the data on snow chemistry are still rare, particularly 160 detailed information on NO3. From 2009 to 2013, we therefore conducted surface snow and snowpit 161 sampling campaigns along the traverse route, and with the main objectives were to (1) to describe NO<sub>3</sub> 162 distribution in surface snow and snowpits, (2)-to characterize the relationship between archived NO<sub>3</sub>-163 and snow accumulation rate, and (3)-to examine the potential effects of coexisting ions on NO<sub>3</sub>-164 preservation. The results of this study may help to better understand NO<sub>3</sub><sup>-</sup> deposition and preservation 165 in the snowpack, which is critical to the interpretation of ice core  $NO_3^-$  records.

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## 167 2 Methodology

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#### 169 2.1 Study area (Zhongshan to Dome A traverse)

170 The Zhongshan to Dome A CHINARE (Chinese National Antarctic Research Expedition) inland 171 traverse is an important leg of the ITASE (International Trans-Antarctic Scientific Expedition). The 172 traverse is in the Indian Ocean sector of East Antarctica, passing through the Lambert Glacier, the 173 largest glacier in Antarctica. In January 1997 the first Chinese Antarctic inland expedition reached an 174 area ~300 km from the coast; in January 1998 the traverse was extended to 464 km, and in December 175 1998, to the Dome A area ~1100 km from the coast. In the austral 2004/2005 summer for the first time, 176 the traverse extended to the ice sheet summit, Dome A, a total distance of ~1260 km. In January 2009, 177 the Chinese inland research base, Kunlun station (80°25'1.7"S and 77 °6'58.0"E, 4087 m above mean 178 sea level), was established at Dome A, mainly aiming aimed at deep ice core drilling and astronomical 179 observations. Now, Kunlun base is a summer station, and the CHINARE team typically conducts an

#### annual inland traverse from the coastal Zhongshan station to Dome A.

181 In January 2010, the Dome A deep ice core project was started, and the construction of basic 182 infrastructure (including drill trench and scientific workroom) took 4 summer seasons. The deep ice 183 core drilling began in January 2013, and in total 801 m ice core was recovered by <u>the</u> 2016/2017 season. 184 The investigation of  $NO_3^-$  deposition and preservation in the snowpack will be of help to the 185 interpretation of Dome A deep ice core  $NO_3^-$  records.

# 187 **2.2 Sample collection**

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188 During the 2010/2011 CHINARE, surface snow samples (the topmost ~3 cm) were collected at an 189 interval of ~10 km along the traverse route from Zhongshan to Dome A, using 3.0 cm diameter 190 high-density polyethylene (HDPE) bottles (volume = 100 ml). The bottles were pre-cleaned with 191 Milli-Q ultrapure water (18.2 M $\Omega$ ), until electrical conductivity of the water stored in bottles (> 24 h) 192 decreased to  $<0.5 \ \mu\text{S cm}^{-1}$ . Then, the bottles were dried under a class 100 super clean hood at 20 °C. Immediately after the drying procedure, the bottles were sealed in clean PE bags that were not opened 193 194 until the field sampling started. At each sampling site (typically > 500 m away from the traverse route), 195 the bottles were pushed into surface snow layers in the windward direction. In total, 120 surface snow 196 samples were collected. In addition, at each sampling site, the upper snow density (~10 cm) was 197 measured using a density scoop with the a volume of 1000 cm<sup>3</sup>. As the field blanks, pPre-cleaned 198 bottles filled with Milli-Q water were taken to the field and treated to the same conditions as field 199 samples to represent field blanks (n = 3).

200 On the Dome A plateau, the snow is soft and non-cohesive, and morphology of the surface snow is 201 different from other areas on the traverse, with a needle ice crystal layer extensively developed, in 202 particular on the sastrugi (Fig. S1 in supporting information). The depth of the needle-like crystal ice 203 layer (referred to as-the "crystal ice" in the following context) is generally < 1.0 cm. In order to 204 investigate air-snow transfer of NO3<sup>-</sup> in this uppermost ~1 cm layer, the crystal ice was collected using 205 a clean HDPE scoop, and then was poured into the clean wide mouth HDPE bottles. Approximately 30 206 g of crystal ice was collected for each sample. In total, 6 crystal ice samples were collected on the 207 traverse near Dome A plateau.

208 In addition to surface snow, snowpit samples were collected during CHINARE inland traverse 209 campaigns in 2009/2010, 2010/2011, and 2012/2013. The snowpits were excavated manually, and the 210 snow wall in the windward direction was scraped clean and flat with a clean HDPE scraper. Then the 211 bottles were pushed horizontally into the snow wall. Snowpit samples were collected from the base 212 towards the top layer along a vertical line. During the sampling process, all personnel wore PE gloves 213 and facemasks to minimize potential contamination. Note that the snowpits are generally > 1 km from 214 the traverse route to avoid possible contamination from the expedition activities. The full information 215 about individual snowpits, including location, distance from the coast, elevation, snowpit depth, 216 sampling resolution, collection date, and annual snow accumulation rate, is summarized in Table 1. All 217 together, 20 snowpits (SP1 to SP20 in Fig. 2, with SP20 corresponding to the location of Kunlun 218 station at Dome A), as 1741 snow samples, were collected.

219 To support understanding of the air-snow transfer of NO<sub>3</sub> on the traverse, atmospheric NO<sub>3</sub> was
 220 collected on glass fiber filters (Whatman G653) using a high volume air sampler (HVAS), with a flow

- **221** rate of  $\sim 1.0 \text{ m}^3 \text{ min}^{-1}$  for 12-15 hr, during the inland traverse campaign in 2015/2016. The NO<sub>3</sub><sup>-5</sup>
- 222 <u>collected on glass fiber filters are expected to equal the sum of particulate NO<sub>3</sub><sup>-</sup> and gaseous HNO<sub>3</sub>.</u>
- 223 based upon previous investigations in East Antarctica (Savarino et al., 2007; Frey et al., 2009; Erbland

224 et al., 2013). In total, 34 atmospheric samples were collected on the traverse. In addition, two field

225 blanks were collected from filters installed in the HVAS without pumping and treated as samples

thereafter. Detailed information about the atmospheric sampling is presented in Table S1 in supporting
 information.

After sample collection, all filters and snow samples. After snow collection, the bottles were sealed in
 clean PE bags-again and preserved in clean thermal insulated boxes. All of the samples were
 transported to the laboratory under freezing conditions (< -20 °C).</li>

In the laboratory, three quarters of individual filters were cut into pieces using pre-cleaned scissors

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## 232 2.3 Sample analysis

234 that were rinsed between samples, placed in ~100 ml of Milli-Q water, ultrasonicated for 40 min and 235 leached for 24 hr under shaking. The sample solutions were then filtered through 0.22 µm ANPEL 236 PTFE filters for concentration analysis. Snow samples were melted in the closed sampling bottles on a super clean bench (class 100) before chemical measurements. Analyses of Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, Ca<sup>2+</sup>, 237 Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup> and SO<sub>4</sub><sup>2-</sup> were performed using a Dionex ICS-3000 ion chromatography system. The column 238 used for cation analysis (Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup> and Ca<sup>2+</sup>) was a Dionex column CS12 (2×250 mm), with 239 a guard column CG12 (2×50 mm); while the anions (Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup> and SO<sub>4</sub><sup>2</sup>) were analyzed using a 240 Dionex column AS11 (2×250 mm) with a guard column AG11 (2×50 mm). The eluent for cations was 241 242 18.0 mM methanesulfonic acid (MSA), and the gradient elution method was employed for anion 243 analysis, with eluent of potassium hydroxide (KOH). More details on this method are described in a previous report (Shi et al., 2012). During sample analysis, duplicated samples and field blanks were 244 synchronously analyzed. The pooled standard deviation  $(\sigma_p, \sigma_p = \sqrt{\sum_{i=1}^{k} (n_i - 1)s_i^2 / \sum_{i=1}^{k} (n_i - 1)},$ 245

where  $n_i$  and  $s_i^2$  are the size and variance of the *i*th samples respectively, and k is the total number of sample sets) of all replicate samples run at least twice in two different sample sets is 0.019 (CI), 0.023 (NO<sub>3</sub><sup>-</sup>), 0.037 (SO<sub>4</sub><sup>2-</sup>), 0.022 (Na<sup>+</sup>), 0.039 (NH<sub>4</sub><sup>+</sup>), 0.006 (K<sup>+</sup>), 0.006 (Mg<sup>2+</sup>) and 0.006 (Ca<sup>2+</sup>) µeq L<sup>-1</sup> respectively (n = 65 pairs of samples). Ion concentrations in field blanks (n = 3) are generally lower

than the detection limit (DL, 3 standard deviations of water blank in the laboratory).

For Antarctic snow samples, the concentrations of  $H^+$  are usually not measured directly, but deduced from the ion-balance disequilibrium in the snow. Here,  $H^+$  concentration is calculated through ion balance.

254  $[H^+] = [Cl^-] + [NO_3^-] + [SO_4^{2-}] - [Na^+] - [NH_4^+] - [Mg^{2+}] - [Ca^{2+}] (Eq. 1),$ 

where ion concentrations are in  $\mu$ eq L<sup>-1</sup>. In addition, the non-sea salt fractions of SO<sub>4</sub><sup>2-</sup> (nssSO<sub>4</sub><sup>2-</sup>) and Cl<sup>-</sup> (nssCl<sup>-</sup>) can be calculated from the following expressions, by assuming Na<sup>+</sup> exclusively from sea salt (in  $\mu$ eq L<sup>-1</sup>).

258  $[nssSO_4^{2-}] = [SO_4^{2-}] - 0.12 \times [Na^+] (Eq. 2),$ 

**259**  $[nssCl^{-}] = [Cl^{-}] - 1.17 \times [Na^{+}] (Eq. 3).$ 

260 It is noted that  $SO_4^{2-}$  fractionation (the precipitation of mirabilite (Na<sub>2</sub>SO<sub>4</sub> 10H<sub>2</sub>O)) may introduce a 261 bias in nssSO<sub>4</sub><sup>2-</sup>, particularly during the winter half year (Wagenbach et al., 1998a).

- 262
- 263 3 Results
- 264

#### 265 3.1 NO<sub>3</sub><sup>-</sup> concentration in surface snow

266 Concentrations of  $NO_3^-$  in surface snow are shown in Fig. 1, ranging from 0.6 to 5.1  $\mu$ eq L<sup>-1</sup>, with a

267 mean of 2.4  $\mu$ eq L<sup>-1</sup>. One standard deviation (1 $\sigma$ ) of NO<sub>3</sub><sup>-</sup> concentration in surface snow is 1.1  $\mu$ eq L<sup>-1</sup>, 268 with coefficient of variation (Cv, 1 $\sigma$  over mean) of 0.5, indicating a moderate spatial variability. On the 269 coastal ~450 km,  $NO_3^-$  shows a slightly increasing trend towards the interior, with a-low variability, while NO3<sup>-</sup> concentrations are higher in the inland region, with a large fluctuation. It is notable that in 270 the area ~800 km from the coast, where snow accumulation is relatively high, NO3<sup>-</sup> concentrations 271 272 decrease to  $< 2.0 \ \mu eq L^{-1}$ , comparable to the values on the coast. Near the Dome A plateau (> 1000 km 273 from coast), there is a tendency for higher NO<sub>3</sub><sup>-</sup> concentrations (> 5.0  $\mu$ eq L<sup>-1</sup>). Similarly, atmospheric  $NO_3^-$  concentrations increase from the coast towards the plateau, ranging from 8 to 183 ng m<sup>-3</sup> (mean = 274 275 55 ng m<sup>-3</sup>) (Fig. 1).

The percentage that surface snow NO<sub>3</sub><sup>-</sup> contributes to total ions (i.e., total ionic strength, sum of Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, Ca<sup>2+</sup>, Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>2-</sup> and H<sup>+</sup>, in µeq L<sup>-1</sup>) varies from 6.7 to 37.6 % (mean = 27.0 %; Fig. S2 in supporting information), with low values near the coast and high percentages on the plateau. A strong relationship was found between NO<sub>3</sub><sup>-</sup> and the total ionic strength in surface snow ( $R^2 = 0.55$ , p < 0.01).

In the crystal ice, the means (ranges) of Cl<sup>+</sup>, NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>-2</sup>, Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, Ca<sup>2+</sup> and H<sup>+</sup> 281 concentrations are 0.98 (0.62 - 1.27), 10.40 (8.35 - 16.06), 1.29 (0.87 - 2.13), 0.27 (0.21 - 0.33), 0.24282 283 (0.03 - 0.56), 0.05, (0.03 - 0.08), 0.18, (0.15 - 0.22), 0.18, (0.05 - 0.57) and 11.75, (9.56 - 18.12) µeq L<sup>-1</sup>, 284 respectively. H<sup>+</sup> and NO<sub>3</sub><sup>-</sup> are the most abundant species, accounting for 46.4 and 41.0 % of the total ions, followed by  $SO_4^{2-}$  (5.1 %) and  $Cl^-$  (3.9 %). The other 5 cations,  $Na^+$ ,  $NH_4^+$ ,  $K^+$ ,  $Mg^{2+}$  and  $Ca^{2+}$ , 285 286 only represent 3.6 % of the total ion budget. A significant linear relationship was found between NO<sub>3</sub> and the total ionic strength ( $R^2 = 0.99$ , p < 0.01), possibly suggesting that NO<sub>3</sub><sup>-</sup> is the species 287 288 controlling ion abundance by influencing acidity of the crystal ice (i.e., H<sup>+</sup> levels). In comparison with surface snow, concentrations of  $H^+$  and  $NO_3^-$  are significantly higher in crystal ice (Independent 289 Samples T Test, p<0.01), while concentrations of Cl<sup>-</sup>, SO<sub>4</sub><sup>-2</sup>, Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup> and Ca<sup>2+</sup> are 290 291 comparable in the two types of snow samples (Fig. S2 in supporting information). To date, the 292 information on the chemistry of ice crystal is rather limited but data from the so-called skin layer at Dome C (top ~4 mm snow), where NO<sub>3</sub> concentrations are in the range of 9 – 22  $\mu$ eq L<sup>-1</sup> in 293 294 summertime (Erbland et al., 2013), are generally comparable to our observations.

295 NO3<sup>-</sup> concentrations in surface snow have been widely measured across Antarctica (Fig. 2), and the 296 values vary from 0.2 to 12.9  $\mu$ eq L<sup>-1</sup>, with a mean of 2.1  $\mu$ eq L<sup>-1</sup> (n = 594,  $1\sigma = 1.7 \mu$ eq L<sup>-1</sup>) and a median of 1.4  $\mu$ eq L<sup>-1</sup>. Most of the data (87 %) fall in the range of 0.5 - 4.0  $\mu$ eq L<sup>-1</sup>, and only 7 % of the 297 298 values are above 5.0  $\mu$ eq L<sup>-1</sup>, mainly distributed on the East Antarctic plateaus. Spatially, NO<sub>3</sub><sup>-1</sup> 299 concentrations show an increasing trend with distance inland, and the values are higher in East than in 300 West Antarctica. Overall, this spatial pattern is opposite to that of the annual snow accumulation rate 301 (Arthern et al., 2006), i.e., low (high) snow accumulation corresponds to high (low) NO3-302 concentrations. It is difficult to compare with NO<sub>3</sub><sup>-</sup> concentrations derived from the "upper snow layer" 303 in different studies because each study sampled a different depth (Fig. 2), e.g., 2 - 10 cm for 304 DDU-Dome C traverse (Frey et al., 2009; Erbland et al., 2013), 25 cm for the 1989-1990 International 305 Trans-Antarctica Expedition (Qin et al., 1992) and 3 cm for this study. The different sampling depths 306 can result in large differences in NO3<sup>-</sup> concentration, especially on the East Antarctic plateaus (e.g., the 307 values of the topmost 1 cm of snow, the crystal ice in this study, can be up to >15  $\mu$ eq L<sup>-1</sup>; Fig. 1). In 308 this ease Because of this, any comparison of NO<sub>3</sub><sup>-</sup> concentrations in surface snow collected in different 309 campaigns should be made with caution.

#### 311 3.2 Snowpit NO<sub>3</sub><sup>-</sup> concentrations

312 Mean  $NO_3^-$  concentrations for snowpits are shown in Fig. 1. On-From the coast to the coastal ~450 313 km\_inland, snowpit  $NO_3^-$  means are comparable to those of surface snow; whereas,  $NO_3^-$  means are 314 lower in inland snowpits than in surface snow with the exception of sites ~800 km from the coast. In 315 general, the differences between snowpit  $NO_3^-$  means and the corresponding surface snow values are 316 small at sites with high snow accumulation (e.g., close to coast), while the differences are large in low 317 snow accumulation areas (e.g., near Dome A).

318 The profiles of NO<sub>3</sub><sup>-</sup> for all snowpits are shown in Fig. 3. NO<sub>3</sub><sup>-</sup> concentrations vary remarkably with 319 depth in pits SP1 - SP5, which are located near the coast. Although SP2 and SP5 show high NO3-320 concentrations in the topmost sample, the data from deeper depths can be compared with the surface 321 values. In addition, NO3<sup>-</sup> means for the entire snowpits are close to the means of the topmost layer 322 covering a full annual cycle of accumulation (i.e., the most recent year of snow accumulation) at 323 SP1-SP5 (Fig. 4). Given the high snow accumulation (Fig. 1), NO<sub>3</sub><sup>-</sup> variability in coastal snowpits is likely suggestive of a seasonal signature (Wagenbach et al., 1998b; Grannas et al., 2007; Shi et al., 324 325 2015). Among the coastal snowpits, water isotope ratios ( $\delta^{18}$ O of H<sub>2</sub>O) of samples at SP $\theta$ <sup>2</sup> were also 326 determined, thus allowing for investigating NO3<sup>-</sup> seasonal variability (Fig. S3 in supporting 327 information). In general, the  $\delta^{18}O(H_2O)$  peaks correspond to high NO<sub>3</sub><sup>-</sup> concentrations (i.e., NO<sub>3</sub><sup>-</sup> peaks 328 present in summer). This seasonal pattern is in agreement with previous observations of  $NO_3^-$  in 329 snow/ice and atmosphere in coastal Antarctica (Mulvaney and Wolff, 1993; Mulvaney et al., 1998; 330 Wagenbach et al., 1998b; Savarino et al., 2007).

In contrast, most of the inland snowpits show high  $NO_3^-$  concentrations in the top layer, and then fall sharply from > 2.0 µeq L<sup>-1</sup> in top snow to < 0.2 µeq L<sup>-1</sup> in the first meter of depth (Fig. 3). NO<sub>3</sub><sup>-</sup> means for the entire snowpits are typically lower than those of the most recent year snow layer (Fig. 4). Similar NO<sub>3</sub><sup>-</sup> profiles for snowpits have been reported elsewhere in Antarctica, as a result of post-depositional processing of NO<sub>3</sub><sup>-</sup> (R öthlisberger et al., 2000; McCabe et al., 2007; Erbland et al., 2013; Shi et al., 2015).

337 Comparison of the NO<sub>3</sub><sup>-</sup> profile patterns reveals significant spatial heterogeneity, even for 338 neighboring sites. For instance, sites SP11 and SP12, 14 km apart, feature similar snow accumulation rate (Table 1). If it is assumed that snow accumulation is relatively constant during the past several 339 340 years at SP11 (sampled in 2012/2013), snow in the depth of ~54 cm corresponds to the deposition in 2009/2010 (snow density = 0.45 g cm<sup>-3</sup>, from field measurements). NO<sub>3</sub><sup>-</sup> concentrations are much 341 342 higher in the top snow of SP12 (sampled in 2009/2010) than in the depth of ~54 cm in SP11 (Fig. 3). 343 This variation in NO<sub>3</sub><sup>-</sup> profiles at a local scale has been reported, possibly related to local morphologies associated with sastrugi formation and wind drift (Frey et al., 2009; Traversi et al., 2009). It is 344 345 interesting that higher NO3<sup>-</sup> concentrations were not found in the uppermost layer at sites SP7 and SP8 346 (~600 km from coast; Fig. 3), where large sastrugi with hard smooth surfaces was extensively 347 developed (from field observations; Fig. S4 in supporting information). Snow accumulation rate in this 348 area fluctuates remarkably, and the values of some sites are rather small or close to zero due to the strong wind scouring (Fig. 1) (Ding et al., 2011; Das et al., 2013). In this case, the snowpit NO<sub>3</sub>-349 350 profiles appear to be largely influenced by wind scour on snow, possibly resulting in missing years 351 and/or intra-annual mixing.

352

#### 353 4 Discussion

#### 355 4.1 Accumulation influence on NO<sub>3</sub><sup>-</sup>

356 The preservation of NO<sub>3</sub><sup>-</sup> is thought to be closely associated with snow accumulation, where most of 357 the deposited  $NO_3^-$  is preserved at sites with higher snow accumulation (Wagenbach et al., 1994; 358 Hastings et al., 2004; Fibiger et al., 2013). Whereas, NO3<sup>-</sup> may be altered significantly at sites with low 359 snow accumulation, largely due to photolysis (Blunier et al., 2005; Grannas et al., 2007; Frey et al., 360 2009; Erbland et al., 2013; Erbland et al., 2015). In the following discussion, we divide the traverse 361 into two zones, i.e., the coastal zone (-<-450 km from the coast, including SP1-SP5 and Core 1; Table 362 1) and the inland region (~450 km to Dome A, including pits SP6-SP20 and Core 2; Table 1), following 363  $NO_3$  distribution patterns in surface snow and snowpits (sections 3.1 and 3.2) as well as the spatial 364 pattern of snow accumulation rate (Fig. 1).

365 As for snowpits,  $NO_3^-$  levels in top and deeper layers are comparable near the coast, while  $NO_3^-$ 366 differs considerably between the upper and deeper snow at inland sites (Figs. 3 and 4). It is 367 demonstrated that <u>p</u>Photochemical processing is responsible for NO<sub>3</sub><sup>-</sup> distribution in inland snowpits 368 (Erbland et al., 2013; Berhanu et al., 2015). Considering that the actinic flux is always negligible below 369 the depth of 1 m, the bottom layers of the snowpits (i.e., > 100 cm; Table 1) are well below the 370 photochemically active zone (France et al., 2011; Zatko et al., 2013). In this case, NO<sub>3</sub><sup>-</sup> in the bottom 371 snowpit, i.e., below the photic zone, can be taken as the archived fraction without further modification, 372 as also suggested by on the basis of previous observations (Frey et al., 2009; Erbland et al., 2013; 373 Erbland et al., 2015). Here, we define  $NO_3^-$  in the bottom layer covering a full annual cycle of 374 deposition as an approximation of the annual mean of archived NO3<sup>-</sup> (i.e., beyond photochemical 375 processing; denoted as " $C_{archived}$ " in the following context; Fig. 4), thus allowing for calculating the 376 archived annual NO<sub>3</sub><sup>-</sup> flux (i.e., the product of  $C_{\text{archived}}$  and annual snow accumulation rate). Although 377 there is uncertainty over in the calculation of archived  $NO_3^-$  flux due to interannual variability in  $NO_3^-$ 378 inputs and snow accumulation, this assumption provides a useful way to investigate the relationship 379 between preservation of  $NO_3^-$  and physical factors considering that an extensive array of ice core 380 measurements is unavailable in most of Antarctica. It is noted that Carchived is generally close to (lower 381 than) the NO<sub>3</sub><sup>-</sup> means for entire snowpits in coastal (inland) Antarctica (Fig. 4).

#### 382

#### 383 4.1.1 NO<sub>3</sub><sup>-</sup> in coastal snowpack

The simplest plausible model to relate flux and concentration of  $NO_3^-$  in snow to its atmospheric concentration (Legrand, 1987; Alley et al., 1995) can be expressed as  $\frac{1}{2}$ 

 $F_{\text{total}} = K_1 C_{\text{atm}} + K_2 C_{\text{atm}} A \text{ (Eq. 4)},$ 

$$F_{\text{total}} = C_{\text{firm}} \times A \text{ (Eq. 5)},$$

388 where  $F_{\text{total}}$  is snow NO<sub>3</sub><sup>-</sup> flux (µeq m<sup>-2</sup> a<sup>-1</sup>);  $C_{\text{atm}}$  is atmospheric concentration of NO<sub>3</sub><sup>-</sup> (µeq m<sup>-3</sup>); *A* is 389 annual snow accumulation rate (kg m<sup>-2</sup> a<sup>-1</sup>);  $C_{\text{firn}}$  is measured firn NO<sub>3</sub><sup>-</sup> concentration (µeq L<sup>-1</sup>, here 390  $C_{\text{firn}} = C_{\text{archived}}$ ;  $K_1$  is the dry deposition velocity (cm s<sup>-1</sup>); and  $K_2$  is the scavenging ratio for precipitation 391 (m<sup>3</sup> kg<sup>-1</sup>), which allows to convert conversion of atmospheric concentration to snow concentration of 392 NO<sub>3</sub><sup>-</sup> in this study. From Eqs. 4 and 5, firn NO<sub>3</sub><sup>-</sup> concentration can be expressed as,

393  $C_{\text{firm}} = K_1 C_{\text{atm}} \times 1/A + K_2 C_{\text{atm}} (\text{Eq. 6})$ 

394 If  $K_1$  and  $K_2$  are constants, a linear relationship between measured NO<sub>3</sub><sup>-</sup> concentration ( $C_{\text{firn}}$ ) and snow 395 accumulation (A) can be interpreted using Eq. 6, which assumes <u>regional</u> spatial homogeneity of fresh 396 snow NO<sub>3</sub><sup>-</sup> levels and dry deposition flux<u>in</u> the regions. The slope ( $K_1C_{\text{atm}}$ ) of the linear model 397 represents an approximation of dry deposition flux of NO<sub>3</sub><sup>-</sup> (i.e., an apparent dry deposition flux), while 398 the intercept ( $K_2C_{\text{atm}}$ ) stands for NO<sub>3</sub><sup>-</sup> concentration in fresh snowfall. If dry deposition ( $K_1C_{\text{atm}}$ ) is much larger than wet deposition ( $K_2C_{atm}A$ ), the concentration of NO<sub>3</sub><sup>-</sup> in snow will be proportional to its concentration in the atmosphere. In the condition of a constant atmospheric concentration, larger snow accumulation will increase the flux of NO<sub>3</sub><sup>-</sup> but decrease its concentration in snow. While this linear model is a gross over-simplification of the complex nature of air-snow exchange of NO<sub>3</sub><sup>-</sup>, it provides a simple approach to compare the processes occurring on the coast versus those inland. In addition, this model can provides useful parameter values in modeling NO<sub>3</sub><sup>-</sup> deposition/preservation at large scales, considering that observations remain sparse across Antarctica (e.g., Zatko<sub>7</sub> et al., 2016).

The relationship between  $C_{\text{archived}}$  of NO<sub>3</sub><sup>-</sup> and snow accumulation rate is shown in Fig. 5. The linear fit of  $C_{\text{archived}}$  vs. inverse snow accumulation ( $R^2$ =0.88, p<0.01; Fig. 5a) supports the assumptions of spatial homogeneity. The intercept and slope of the linear fit suggest a NO<sub>3</sub><sup>-</sup> concentration in fresh snow and an apparent NO<sub>3</sub><sup>-</sup> dry deposition flux of 0.7±0.07 µeq L<sup>-1</sup> and 45.7±7.8 µeq m<sup>-2</sup> a<sup>-1</sup> respectively. The apparent dry deposition flux is opposite to the observation in Dronning Maud Land (DML) region, where the <u>a</u> negative dry deposition flux suggested net losses of NO<sub>3</sub><sup>-</sup> (Pasteris et al., 2014).

Figure 5b shows the archived fluxes of NO<sub>3</sub><sup>-</sup> on the coast, with values from 104 (at the lowest 413 accumulation site) to 169  $\mu$ eq m<sup>-2</sup> a<sup>-1</sup> (at the highest accumulation site). Taking the calculated NO<sub>3</sub><sup>-</sup> dry 414 deposition flux of 45.7  $\mu$ eq m<sup>-2</sup> a<sup>-1</sup>, dry deposition accounts for 27-44 % (mean = 36 %) of total NO<sub>3</sub><sup>-1</sup> 415 416 inputs, with higher (lower) percentages at lower (higher) snow accumulation sites. This result is in line 417 with the observations in Taylor Valley (coastal West Antarctica), where the snowfall was found to be 418 the primary driver for  $NO_3^-$  inputs (Witherow et al., 2006). This observation also generally agrees with, 419 but is greater than that in the modeling study of Zatko et al. (2016), which predicts a ratio of dry 420 deposition to total deposition of  $NO_3^-$  in Antarctica as < 20 % close to the coast, increasing towards the 421 plateaus.

422 In Figs. 5a and b, the strong linear relationships between  $NO_3^-$  and snow accumulation support that  $K_1$  and  $K_2$  are relatively constant on the coast (Eqs. 4 and 6). The average atmospheric concentration of 423 424  $NO_3$  in the coastal ~450 km region is 19.4 ng m<sup>-3</sup> in summer (Table S1 in supporting information). Taking  $C_{\text{atm}}=19.4 \text{ ng m}^{-3}$ ,  $K_1$  is estimated to be 0.5 cm s<sup>-1</sup>, identical to a typical estimate for HNO<sub>3</sub> 425 deposition velocity to a snow/ice surface (0.5 cm s<sup>-1</sup>; Seinfeld and Pandis, 1997). This predicted  $K_1$ 426 427 value is lower than that calculated for the dry deposition of HNO<sub>3</sub> at South Pole ( $0.8 \text{ cm s}^{-1}$ ; Huey et al., 2004). It is noted that the true  $K_1$  value could be larger than the prediction (0.5 cm s<sup>-1</sup>) due to the higher 428 429 values of  $C_{\text{atm}}$  atmospheric NO<sub>3</sub>-concentrations during in summer (i.e., 19.4 ng m<sup>-3</sup> for the calculation 430 of K<sub>1</sub>)time than in other seasons (Mulvaney et al., 1998; Wagenbach et al., 1998b; Savarino et al., 431 2007). The scavenging ratio for precipitation ( $K_2$ ) is calculated to be  $0.2 \times 10^4$  m<sup>3</sup> kg<sup>-1</sup>, i.e., 2 m<sup>3</sup> g<sup>-1</sup>.

432 If it is assumed that  $NO_3^-$  concentration in snow is related to its concentration in the atmosphere, the 433 scavenging ratio for  $NO_3^-$  (*W*) can be calculated on a mass basis from the following expression 434 (Kasper-Giebl et al., 1999),

435

$$W = \rho_{\rm atm} \times (C_{\rm f-snow} / C_{\rm atm})$$
 (Eq. 7),

436 where  $\rho_{atm}$  is air density (g m<sup>-3</sup>), and  $C_{f-snow}$  and  $C_{atm}$  are NO<sub>3</sub><sup>-</sup> concentrations in fresh snow (ng g<sup>-1</sup>) and 437 atmosphere (ng m<sup>-3</sup>) respectively. If taking  $\rho_{atm} \approx 1000$  g m<sup>-3</sup> (on average, ground surface temperature *t* 438  $\approx 255$  k, ground pressure  $P \approx 0.08$  MPa, in the coastal region),  $C_{f-snow} = 43$  ng g<sup>-1</sup> (see discussion above 439 and section 4.2 below), and  $C_{atm} = 19.4$  ng m<sup>-3</sup>, W is calculated to be ~2200, generally comparable to 440 previous reports (Barrie, 1985; Kasper-Giebl et al., 1999; Shrestha et al., 2002). It is noted that the 441 calculation here may be subject to uncertainty, due to the complex transfer of atmospheric NO<sub>3</sub><sup>-</sup> into the 442 snow. However, the scavenging ratio provides useful-valuable insights into the relation between NO<sub>3</sub><sup>-</sup> 443 concentrations in the atmosphere and snow, which might be useful in modeling NO<sub>3</sub><sup>-</sup> deposition at <u>a</u>
444 large-scale.

Figure 5c shows the distribution of flux is negatively correlated with  $C_{\text{archived}}$  of NO<sub>3</sub><sup>-</sup>, which is not surprising since  $C_{\text{archived}}$  is positively related to inverse accumulation (Fig. 5a). Based on the observed strong linear relationship between NO<sub>3</sub><sup>-</sup> flux and snow accumulation (Fig. 5b), the archived NO<sub>3</sub><sup>-</sup> flux is more accumulation dependent compared to  $C_{\text{archived}}$ . This is compatible with the observations in Greenland (Burkhart et al., 2009), where accumulation is generally above 100 kg m<sup>-2</sup> a<sup>-1</sup>, similar to the coastal values in this study.

451 In terms of surface snow on the coast,  $NO_3^-$  may be disturbed by the katabatic winds and wind 452 convergence located near the Amery Ice Shelf (that is, the snow-sourced  $NO_x$  and  $NO_3$  from the 453 Antarctic plateau possibly contributes to coastal snow NO<sub>3</sub>) (Parish and Bromwich, 2007; Ma et al., 454 2010; Zatko et al., 2016). In addition, the sampled ~3 cm surface layer roughly corresponds to the net 455 accumulation in the past 0.5-1.5 months assuming an even distribution of snow accumulation in the 456 course of a single year. This difference in exposure time of the surface snow at different sampling sites, 457 could possibly affect the concentration of  $NO_3$ , although the post-depositional alteration of  $NO_3$  was 458 thought to be minor on the coast (Wolff et al., 2008; Erbland et al., 2013; Shi et al., 2015). Taken 459 together, NO<sub>3</sub><sup>-</sup> in coastal surface snow might represent some post-depositional alteration. Even so, a 460 negative correlation between NO3<sup>-</sup> concentration and snow accumulation rate was found at the coast 461  $(R^2=0.42, p<0.01; Fig. 6a)$ , suggesting that overall the majority of the NO<sub>3</sub><sup>-</sup> appears to be preserved and 462 is driven-determined by snow accumulation.

463

#### 464 4.1.2 NO<sub>3</sub><sup>-</sup> in inland snowpack

465 In comparison with the coast, the correlation between  $C_{\text{archived}}$  and inverse snow accumulation is relatively weak in inland regions (Fig. 5d), suggesting more variable conditions in ambient 466 concentrations and dry deposition flux of NO<sub>3</sub><sup>-</sup>. In addition, the relationship of  $C_{\text{archived}}$  vs. inverse 467 468 accumulation-in inland is opposite to that of coast. Based on current understanding of the 469 post-depositional processing of  $NO_3$ , the negative correlation between  $C_{\text{archived}}$  and inverse snow 470 accumulation (Fig. 5d) suggests losses of NO3. The slope of the linear relationship indicates apparent NO<sub>3</sub><sup>-</sup> dry deposition flux of -44.5  $\pm$ 13.0 µeq m<sup>-2</sup> a<sup>-1</sup>, much larger than that of DML (-22.0  $\pm$ 2.8 µeq m<sup>-2</sup> 471 a<sup>-1</sup>), where the snow accumulation is generally lower than 100 kg m<sup>-2</sup> a<sup>-1</sup> (Pasteris et al., 2014). At 472 Kohnen Station (an inland site in East Antarctica), with snow accumulation of 71 kg m<sup>-2</sup>  $a^{-1}$ , the 473 emission flux of NO<sub>3</sub><sup>-</sup> is estimated to be -22.9  $\pm$ 13.7 µeq m<sup>-2</sup> a<sup>-1</sup> (Weller and Wagenbach, 2007), which 474 475 is also smaller in comparison with this observation. Weller et al. (2004) proposed that loss rate of  $NO_3^{-1}$ 476 does not depend on snow accumulation rate and the losses become insignificant at accumulation rates 477 above 100 kg m<sup>-2</sup> a<sup>-1</sup>. Among the inland sites, SP10 and Core2 (~800 km from the coast), featured by high snow accumulation rate (> 100 kg m<sup>-2</sup> a<sup>-1</sup>; Table 1 and Fig. 1), exhibit even higher values of 478 479  $C_{\text{archived}}$  and archived fluxes of NO<sub>3</sub><sup>-</sup> than those of the coastal sites. It is noted that the these two cases 480 influence the linear regression significantly (Fig. 5d). If the two sites are excluded, we can get a linear regression with the <u>a</u> slope of  $-27.7\pm9.2 \ \mu eq \ m^{-2} \ a^{-1}$ , which is comparable to previous reports in DML 481 482 (Pasteris et al., 2014).

483 The depths of inland snowpits cover past-several to tens of years' snow accumulation, thus allowing 484 for directly investigating  $NO_3^-$  emission rate. The difference between  $NO_3^-$  concentrations in the snow

485 layer accumulated during the most recent year (Fig. 4) and in the snow accumulated during the year 486 before the most recent year can represent the loss rate of  $NO_3^-$ . If it is assumed that snow accumulation 487 rate is relatively constant during past-recent decades at specific-sites, on average,  $36.7 \pm 21.3$  % of NO<sub>3</sub><sup>-</sup> 488 (in  $\mu$ eq L<sup>-1</sup>) was lost during one year, with (the two sites (SP10 and Core2) with snow accumulation >100 kg m<sup>-2</sup> a<sup>-1</sup> excluded from the calculation). The percentages are generally higher at 489 490 the sites with lower snow accumulation rate. Together with snow accumulation rate, the emission flux 491 of NO<sub>3</sub><sup>-</sup> is calculated to be -28.1  $\pm$ 23.0 µeq m<sup>-2</sup> a<sup>-1</sup>, close to the linear model prediction (-27.7  $\pm$ 9.2 µeq m<sup>-2</sup> a<sup>-1</sup>). The sSignificant losses can account for NO<sub>3</sub> profiles at inland sites, i.e., NO<sub>3</sub> concentration 492 493 decrease decreasing with increasing depths. Previous observations and modeling works suggested that 494 photolysis dominates the losses (Frey et al., 2009; Erbland et al., 2013; Shi et al., 2015). During 495 photolysis of  $NO_3^-$ , some of the photoproducts ( $NO_x$ ) are emitted into the gas phase (Davis et al., 2004; 496 France et al., 2011), and these products should undergo reoxidation by the local oxidants (e.g., 497 hydroxyl radical (OH), NO<sub>2</sub> + OH + M  $\rightarrow$  HNO<sub>3</sub> + M), forming gas phase HNO<sub>3</sub>. In inland Antarctica, 498 the dominant NO<sub>3</sub> species in the atmosphere is gaseous HNO<sub>3</sub> during summertime, while particulate 499 NO<sub>3</sub><sup>-</sup> is more important in winter (Legrand et al., 2017b; Traversi et al., 2017). The high levels of gas 500 phase HNO<sub>3</sub> in summer support the importance of the re-emission from snow through the photolysis of 501  $NO_3$  in affecting the atmospheric  $NO_3/NO_3$  budget (Erbland et al., 2013). On the one hand, the 502 gaseous HNO<sub>3</sub> can be efficiently co-condensed with water vapour onto the extensively developed 503 crystal ice layers on Antarctic plateaus (e.g., Fig. S1 in supporting information discussed above), 504 leading to an enrichment of  $NO_3^-$  in surface snow (Bock et al., 2016). On the other hand, a large 505 concentration of HNO<sub>3</sub> would enhance its reaction with sea-salt, leading to elevated particulate NO<sub>3</sub> 506 concentrations (Legrand et al., 2017b). The significant correlation between  $NO_3^-$  and H<sup>+</sup> in inland 507 Antarctic surface snow ( $R^2 = 0.65$ , p < 0.01) seems to support the importance of atmospheric gas phase 508 HNO<sub>3</sub> in affecting surface snow NO<sub>3</sub><sup>-</sup> concentrations, in particular NO<sub>3</sub><sup>-</sup> levels in the crystal ice 509 samples (Fig. 1).

510 Thus far, sSeveral modeling works have been performed to understand NO<sub>3</sub><sup>-</sup> recycling processes 511 across Antarctica (e.g., Erbland et al., 2015; Zatko et al., 2016; Bock et al., 2016), however, each 512 employs different assumptions and large uncertainty remains about in quantifying NO<sub>3</sub><sup>-</sup> recycling and 513 preservation. It is thought that emission and transport strength are the main factors controlling the 514 recycling of  $NO_3$ , while the former is associated with initial  $NO_3$  concentrations, UV and snow optical 515 properties, and the latter is linked with air mass movement (Wolff et al., 2008; Frey et al., 2009). As a 516 result, snow accumulation alone is likely insufficient to account for NO<sub>3</sub> variability in surface snow 517 (i.e., no significant correlation between  $NO_3^-$  concentration and snow accumulation; Fig. 6b).

518 The archived NO<sub>3</sub><sup>-</sup> fluxes vary considerably among inland sites, from ~3 to 333  $\mu$ eq m<sup>-2</sup> a<sup>-1</sup>, with 519 high values generally corresponding to high snow accumulation (Fig. 5e). However, the nearly 1:1 520 relationship between  $C_{\text{archived}}$  and NO<sub>3</sub><sup>-</sup> flux (Fig. 5f), suggests that accumulation rate is not the main 521 driver of the archived NO3<sup>-</sup> concentration. In inland Antarctica, the archived NO3<sup>-</sup> fraction is largely 522 influenced by the length of time that NO<sub>3</sub><sup>-</sup> was exposed to UV radiation (Berhanu et al., 2015), which 523 decreases exponentially in the snowpack. The *e*-folding depth,  $z_e$  value, is thought to be influenced by a 524 variety of factors, such as co-existent impurities (e.g., black carbon), bulk density and grain size (Zatko 525 et al., 2013). In addition, the snow albedo is also dependent on snow physical properties (Carmagnola 526 et al., 2013). Taken together, this suggests that the inland plateau is below a "threshold" of 527 accumulation rate such that the archived NO<sub>3</sub><sup>-</sup> flux cannot be explained by snow accumulation rate.

528

#### 529 4.2 Effects of coexisting ions on NO<sub>3</sub>

530 Atmospheric  $NO_3^{-1}$  in Antarctica is thought to be mainly associated with mid-latitude sources,

531 re-formed NO<sub>3</sub> driven by snow-sourced photolysis products, and/or stratospheric inputs (Savarino et 532 al., 2007; Lee et al., 2014; Traversi et al., 2017 and references therein). Although organic nitrates can 533 play an important role in the atmospheric NO<sub>v</sub> budget, multi-seasonal measurements of surface snow 534 NO3 correlate strongly with inorganic NOy species (especially HNO3) rather than organic (Jones et al., 2011). Here, Wwe investigate whether NO3<sup>-</sup> in snow is closely associated with coexisting ions (e.g., Cl<sup>-</sup>, 535  $SO_4^{2-}$ , Na<sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup> and Ca<sup>2+</sup>) since these ions have different main sources, e.g., Cl<sup>-</sup> and Na<sup>+</sup> are 536 predominantly influenced by sea salt, and SO42- is likely dominated by marine inputs (e.g., sea salt and 537 bio-activity source) (Bertler et al., 2005). In the snow, Cl<sup>-</sup>, Na<sup>+</sup> and SO<sub>4</sub><sup>2-</sup> are the most abundant ions in 538 addition to NO3<sup>-</sup>, and the potential association between NO3<sup>-</sup> and the three ions in the surface snow is 539 discussed here. 540

In surface snow, the non-sea salt fraction of  $SO_4^{2-}$  accounted for 75––99 % of its total 541 budget, with a mean of 95 %. The percentages are were relatively higher in inland regions than at 542 543 coastal sites. On the coast, a positive relationship was found between  $nssSO_4^{2-}$  and  $NO_3^{-}$  ( $R^2 = 0.32$ , p < 0.32) 0.01; Fig. 7a). Previous observations suggest that  $NO_3^-$  and  $nssSO_4^{-2-}$  peaks in the atmosphere and snow 544 are usually present in summer (Jourdain and Legrand, 2002; Wolff et al., 2008; Sigl et al., 2016; 545 546 Legrand et al., 2017a; Legrand et al., 2017b). However, the similar seasonal pattern of the two species is associated with distinct sources, i.e.,  $SO_4^{2^2}$  is mainly derived from marine biogenic emissions while 547 NO<sub>3</sub><sup>-</sup> is influenced by photolysis and tropospheric transport (Savarino et al., 2007; Lee et al., 2014; 548 Zatko et al., 2016). In the atmosphere, most of SO42- is typically found on the submicron particles, 549 550 while most of the  $NO_3^-$  is gaseous HNO<sub>3</sub> and the particulate  $NO_3^-$  is mainly on the intermediate size particles (Jourdain and Legrand, 2002; Rankin and Wolff, 2003; Legrand et al., 2017a; Legrand et al., 551 2017b). Thus, the correlation between  $NO_3^{-1}$  and  $SO_4^{-2-1}$  is unlikely explained by the sources or their 552 occurrence state in the atmosphere (i.e., gaseous and particulate phases). Laluraj et al. (2010) proposed 553 that the correlation between  $nssSO_4^{2-}$  vs.  $NO_3^{-}$  in ice ( $R^2 = 0.31$ , p < 0.01) could be associated with the 554 555 fine  $nssO_4^{2^2}$  aerosols, which <u>could</u> provide nucleation centers forming-the multi-ion complexes with 556 HNO<sub>3</sub> in the atmosphere. This assertion, however, should be examined further, considering that the complex chemistry of  $SO_4^{2^2}$  and  $NO_3^{-1}$  in the atmosphere is far from understood (e.g., Wolff, 1995; 557 Brown et al., 2006). Thus far, the mechanism of  $nssSO_4^{2-}$  influencing NO<sub>3</sub><sup>-</sup> in the snowpack, however, 558 is still debated, and it cannot be ruled out that  $nssSO_4^{2-}$  further affects mobilization of NO<sub>3</sub><sup>-</sup> during 559 and/or after crystallization (Legrand and Kirchner, 1990; Wolff, 1995; Röthlisberger et al., 2000). It is 560 noted that no relationship was found between nssSO4<sup>2-</sup> and NO3<sup>-</sup> in inland snow (Fig. 7d), possibly due 561 562 to the strong alteration of  $NO_3^-$  during post-depositional processes, as discussed in section 4.1.2.

In comparison with  $nssO_4^{2-}$  aerosols, the sea-salt aerosols (Na<sup>+</sup>) are coarser and can be removed 563 preferentially from the atmosphere due to a larger dry deposition velocity. High atmospheric sea salt 564 565 aerosol concentrations are expected to promote the conversion of gaseous HNO3 to the particulate 566 phase, considering that most of the  $NO_3^-$  in the atmosphere is in the gas phase (HNO<sub>3</sub>). In this case, particulate NO<sub>3</sub><sup>-</sup> can be efficiently lost via aerosol mechanisms. In addition, the saline ice also favors 567 568 the direct uptake of gaseous  $HNO_3$  to the ice surface. Changes in partitioning between gas phase (HNO<sub>3</sub>) and particulate phase will affect NO<sub>3</sub><sup>-</sup> levels due to the different wet and dry deposition rates of 569 570 the two species (Aw and Kleeman, 2003). Thus, sea salt aerosols play an important role in the 571 scavenging of gaseous HNO<sub>3</sub> from the atmosphere (Hara et al., 2005), and elevated NO<sub>3</sub> concentrations are usually accompanied by  $Na^+$  spikes in the snowpack (e.g., at Halley station, a 572 573 coastal site; Wolff et al., 2008). Here<u>Surprisingly</u>, no significant correlation was found between Na<sup>+</sup> 574 and NO<sub>3</sub><sup>-</sup> in coastal snow (Fig. 7b). The concentration profiles of NO<sub>3</sub><sup>-</sup> and Na<sup>+</sup> in coastal surface snow

575 are shown in Fig. 8, and NO<sub>3</sub><sup>-</sup> roughly corresponds to Na<sup>+</sup> in some areas, e.g., 50-150 km and 300-450 576 km distance inland, although in general they are not very coherent. It is noted that amongst the 4 snow 577 samples with Na<sup>+</sup> > 1.5  $\mu$ eq L<sup>-1</sup> (open circles in Fig. 8), only one sample co-exhibits a NO<sub>3</sub><sup>-</sup> spike. This is different from observations at Halley station, where  $Na^+$  peaks usually led to elevated  $NO_3^-$  levels in 578 579 surface snow in summer (Wolff et al., 2008). Of the 4 largest Na<sup>+</sup> spikes, one is a fresh snowfall sample 580 (dashed ellipse in Fig. 8), and this sample shows the highest Na<sup>+</sup> concentration (2.8  $\mu$ eq L<sup>-1</sup>) and low 581  $NO_3^-$  (0.75 µeq L<sup>-1</sup>). It is noted that  $NO_3^-$  concentration in this fresh snowfall is close to the model predictions (0.7  $\pm$ 0.07  $\mu$ eq L<sup>-1</sup>; section 4.1.1), validating that the simple linear deposition model (i.e., the 582 583 Eq. 6) can well depict the deposition and preservation of NO3<sup>-</sup> in coastal snowpack. At inland sites, no correlation was found between  $NO_3^-$  and  $Na^+$  (Fig. 7e), likely explained by the alteration of  $NO_3^-$ 584 585 concentration by post-depositional processing (discussed above).

586 In surface snow, nssCl<sup>-</sup> represents 0-64 % (mean = 40 %) of the total Cl<sup>-</sup>. On the coast, it is of 587 interest that nssCl<sup>-</sup> in the 4 samples with the highest Na<sup>+</sup> concentrations (open circles in Figs. 7b and 8) 588 are close to 0, and positive nssCl values were found for the other samples. The fractionation of Na<sup>+</sup> can occur due to mirabilite precipitation in sea-ice formation at <-8 °C (Marion et al., 1999), possibly 589 leading to the positive nssCl<sup> $\cdot$ </sup>. However, even if all of SO<sub>4</sub><sup>2-</sup> in sea water is removed via mirabilite 590 591 precipitation, only 12 % of sea salt Na<sup>+</sup> is lost (Rankin et al., 2002). Considering the smallest sea ice 592 extent in summertime in East Antarctica (Holland et al., 2014), the high Cl<sup>-</sup>/Na<sup>+</sup> ratio (mean = 2.1, well 593 above 1.17 of sea water, in µeq L<sup>-1</sup>) in surface snow is unlikely from sea salt fractionation associated 594 with mirabilite precipitation in sea-ice formation. In this case, nssCl<sup>-</sup> could be mainly related to the 595 deposition of volatile HCl, which is from the reaction of  $H_2SO_4$  and/or HNO<sub>3</sub> with NaCl (Röthlisberger 596 et al., 2003). In this case Thus, nssCl in snowpack can roughly represent the atmospherically deposited 597 HCl. In the summertime, most of the dechlorination (i.e., production of HCl) is likely associated with 598 HNO<sub>3</sub> due to its high atmospheric concentrations (Jourdain and Legrand, 2002; Legrand et al., 2017b). 599 ThusAccordingly, the observed relationship between NO<sub>3</sub> and nssCl<sup>-</sup> (Fig. 7c) appears to suggest that 600 HCl production can be enhanced by elevated HNO<sub>3</sub> levels in the atmosphere.

601 With regard to the crystal ice, no significant correlation was found between NO<sub>3</sub><sup>-</sup> and the coexisting 602 ions (e.g., Cl<sup>+</sup>, Na<sup>+</sup> and  $SO_4^{-2}$ ), suggesting that these ions are generally less influential on NO<sub>3</sub><sup>-</sup> in this 603 uppermost thin layer, compared to the strong air-snow transfer process of NO<sub>3</sub><sup>-</sup> (Erbland et al., 2013). It is noted that  $NO_3^-$  accounts for most of the calculated H<sup>+</sup> concentrations (81---97 %, mean = 89 %), and 604 a strong linear relationship was found between them ( $R^2 = 0.96$ , p < 0.01), suggesting that NO<sub>3</sub><sup>-</sup> is mainly 605 606 deposited as acid, HNO<sub>3</sub>, rather than in particulate form as salts (e.g., NaNO<sub>3</sub> and Ca(NO<sub>3</sub>)<sub>2</sub>). This 607 deduction is in line with the atmospheric observations at Dome C, where NO<sub>3</sub><sup>-</sup> was found to be mainly 608 in gaseous phase (HNO<sub>3</sub>) in summer (Legrand et al., 2017b). On average, the deposition of HNO<sub>3</sub> 609 contributes > 91 % of NO<sub>3</sub><sup>-</sup> in the crystal ice (the lower limit, 91 %, calculated by assuming all of the alkaline species (Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup> and Ca<sup>2+</sup>) are neutralized by HNO<sub>3</sub> in the atmosphere), 610 611 suggesting a dominant role of HNO<sub>3</sub> deposition in snow NO<sub>3</sub><sup>-</sup> concentrations. The elevated high 612 atmospheric NO<sub>3</sub><sup>-</sup> concentrations observed at Dome A (>100 ng m<sup>-3</sup>; 77.12° E and 80.42° S, Table S1 in supporting information) possibly indicate oxidation of gaseous NO<sub>x</sub> to HNO<sub>3</sub>, providing further 613 614 evidence that NO<sub>3</sub><sup>-</sup> recycling driven by photolysis plays an important role in its abundance in snowpack 615 on East Antarctic plateaus.

616

#### 617 5 Conclusions

618 Samples of surface snow, snowpits and the uppermost layer of crystal ice, collected on the traverse

619 from the coast to Dome A, East Antarctica, were used to investigate the deposition and preservation of

 $NO_3$  in snow. In general, a spatial trend of  $NO_3$  in surface snow was found on the traverse, with high

621 (low) concentrations on the plateau (coast). <u>Similarly, NO<sub>3</sub> concentrations in the atmosphere are higher</u>

622 <u>on the plateau than at coastal sites, with a range of 8 to 183 ng m<sup>-3</sup></u>. Extremely high  $NO_3^-$  levels (e.g., >

 $10 \ \mu eq \ L^{-1}$ ) were observed in the uppermost crystal ice layer, possibly associated with re-deposition of

624 the recycled  $NO_3$  – from snow-sourced  $NO_{x}$ . As for the snowpits,  $NO_3$  exhibits high levels in the top 625 layer and low concentrations at deeper depths in the inland region, while no clear trend was found on

626 the coast.

627 On the coast, the archived NO3<sup>-</sup> flux in snow is positively correlated with snow accumulation rate, 628 but negatively with NO<sub>3</sub><sup>-</sup> concentration. A linear model can well depict the relationship between 629 archived NO<sub>3</sub> and snow accumulation, supporting that atmospheric levels and dry deposition fluxes of 630  $NO_3$  are spatially homogeneous on the coast, and that dry deposition plays a minor role in snow  $NO_3$ 631 inputs. The dry deposition velocity and scavenging ratio for  $NO_3$  are estimated to be 0.5 cm s<sup>-1</sup> and 632 2200, respectively. In inland Antarctica, the archived NO3<sup>-</sup> fluxes, varying significantly among sites, are 633 largely dependent on NO<sub>3</sub><sup>-</sup> concentration. A weak correlation between snow accumulation and archived 634  $NO_3$  suggests variable ambient concentrations and dry deposition flux of  $NO_3$ , and the relationship is 635 opposite to that for the coast. This supports the idea that post-depositional processing dominates NO<sub>3</sub> 636 concentration and distribution in inland Antarctica (e.g., Erbland et al., 2013; Erbland et al., 2015; Shi 637 et al., 2015; Zatko et al., 2016).

638 The major ions, Cl<sup>-</sup>,  $SO_4^{2-}$  and Na<sup>+</sup>, originate from different sources from-than NO<sub>3</sub><sup>-</sup>, but could potentially affect the scavenging and preservation of NO<sub>3</sub>. In coastal surface snow, a positive 639 correlation between nssSO4<sup>2-</sup> and NO3<sup>-</sup> suggests the potential influence of fine aerosols on NO3<sup>-</sup> 640 641 formation and/or scavenging, while the coarse sea salt aerosol (e.g., Na<sup>+</sup>) is likely less influential. In 642 contrast to the coast, NO3<sup>-</sup> in inland surface snow is dominated by post-depositional processes, and the 643 effects of coexisting ions on NO3 appear to be rather minor. In inland surface snow, the strong 644 relationship between NO3<sup>-</sup> and H<sup>+</sup> suggests a dominant role of gaseous HNO3 deposition in determining 645 NO<sub>3</sub><sup>-</sup> concentrations.

646

#### 647 Associated content

- 648 Please see the file of Supporting Information.
- 649

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Snowpit	Latitude,	Longitude,	Elevation,	Distance	to	Annual snow	Depth,	Sampling	Sampling
No.	0	0	m	coast, km		accumulation,	cm	resolution,	year
						kg m <sup>-2</sup> a <sup>-1 1)</sup>		cm	
SP1	-70.52	76.83	1613	132		193.2	150	5.0	2010/2011
SP2	-71.13	77.31	2037	200		172.0	150	3.0	2012/2013
SP3	-71.81	77.89	2295	283		99.4	200	5.0	2012/2013
SP4	-72.73	77.45	2489	387		98.3	200	5.0	2012/2013
SP5	-73.40	77.00	2545	452		90.7	200	5.0	2012/2013
SP6	-73.86	76.98	2627	514		24.6	300	2.5	2012/2013
SP7	-74.50	77.03	2696	585		29.2	100	2.0	2012/2013
SP8	-74.65	77.01	2734	602		80.2	180	2.0	2010/2011
SP9	-76.29	77.03	2843	787		54.8	200	2.0	2012/2013
SP10	-76.54	77.02	2815	810		100.7	240	3.0	2010/2011
SP11	-77.13	76.98	2928	879		81.2	200	2.5	2012/2013
SP12	-77.26	76.96	2962	893		83.4	265	5.0	2009/2010
SP13	-77.91	77.13	3154	968		33.3	200	2.0	2012/2013
SP14	-78.34	77.00	3368	1015		87.6	216	3.0	2010/2011
SP15	-78.35	77.00	3366	1017		70.0	162	2.0	2009/2010
SP16	-79.02	76.98	3738	1092		25.4	200	2.5	2012/2013
SP17	-79.65	77.21	3969	1162		46.2	130	2.0	2010/2011
SP18	-80.40	77.15	4093	1250		24.2	300	2.0	2010/2011
SP19	-80.41	77.11	4092	1254		23.7	300	1.0	2009/2010
SP20	-80.42	77.12	4093	1256		23.5	300	2.5	2012/2013
Core 1 <sup>2)</sup>	-70.83	77.08	1850	168		127.0	-	-	1996/1997
Core 2 <sup>3)</sup>	-76.53	77.03	2814	813		101.0	-	-	1998/1999

**Table 1.** Snowpit information on the traverse from coastal Zhongshan Station to Dome A, EastAntarctica.

940 1) Annual snow accumulation rate is obtained from the field bamboo stick measurements (2009 - 2013),

941 updated from the report (Ding et al., 2011). Note that snow accumulation rate at the two ice core sites

are derived from ice core measurements.

943 2) Core 1, ice core data of previous report (Li et al., 1999; Xiao et al., 2004).

944 3) Core 2, ice core data of previous report (Li et al., 2009).



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948 Figure 1. Concentrations of NO<sub>3</sub><sup>-</sup> in snow (surface snow, crystal ice and snowpits; on the primary 949 y-axis) and atmosphere (on the secondary y-axis), with error bars representing one standard deviation 950 of NO<sub>3</sub><sup>-</sup> (1 $\sigma$ ) for individual snowpits. Also shown is the annual snow accumulation rate on the traverse 951 (red solid line; based on Ding et al. (2011)). Note that NO3<sup>-</sup> concentration in one crystal ice sample (red 952 dot)<del>, 16.7  $\mu$ eq L<sup>+</sup> in the parentheses,</del> is higher than the maximum value of the primary y-axis (NO<sub>3</sub><sup>-</sup> concentration = 16.7  $\mu$ eq L<sup>-1</sup> in the parentheses). 953





**Figure 2.** Concentrations of  $NO_3^-$  in surface snow across Antarctica. Note that the values of crystal ice around Dome A were not included. The data of DDU to Dome C is from Frey et al. (2009). The other surface snow  $NO_3^-$  concentrations are from the compiled data (Bertler et al., 2005 and references therein). Also illustrated are the locations of snowpits on the traverse route from Zhongshan to Dome A in this study (SP1 to SP20, solid short blue line; Table 1).



Figure 3. The full profiles of NO<sub>3</sub><sup>-</sup> concentrations for snowpits collected on the traverse from the coast
to Dome A, East Antarctica (SP1 is closest the coast; SP20 the furthest inland; see Figure 2). The
details on sampling of the snowpits refer to Table 1. The numbers in parentheses in each panel denote
the annual snow accumulation rates (kg m<sup>-2</sup> a<sup>-1</sup>). Note that the scales of x-axes for the snowpits SP1 –
SP9 and SP10 – SP-20 are different.





973 Figure 4. Mean concentrations of NO<sub>3</sub><sup>-</sup> for the entire snowpit depth (in square), the uppermost layer

974 covering one-year snow accumulation (in diamond) and the bottom layer covering a full annual cycle 975 of deposition (archived  $NO_3^-$  concentration,  $C_{archived}$ , in triangle).



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**Figure 5.** The relationships among<u>st</u> snow accumulation rate, the archived concentration ( $C_{archived}$ ), and flux of NO<sub>3</sub><sup>-</sup> in coastal (top row, (a), (b) and (c)) and inland (bottom row, (d), (e) and (f)) Antarctica. In panel (d), the linear fit in black line (y = -44.5x + 2.1) includes the full date data set, while the linear equation in red (y = -27.7x + 1.5) was obtained by excluding two cases (open circles) with snow accumulation rate larger than 100 kg m<sup>-2</sup> a<sup>-1</sup> (see the main text). The flux values are the product of  $C_{archived}$  of NO<sub>3</sub><sup>-</sup> and snow accumulation rate, namely the archived flux. Least squares regressions are noted with solid lines and are significant at p < 0.01. Error bars represent one standard deviation (1 $\sigma$ ).



990 Figure 6. The relationships between NO<sub>3</sub><sup>-</sup> concentration and inverse snow accumulation rate in surface

991 snow in coastal (panel (a)) and inland (panel (b)) Antarctica. Least squares regressions are noted with 992 solid line and are significant at p < 0.01.



Figure 7. Relationships between NO<sub>3</sub><sup>-</sup> and co-existing major ions in surface snow in coastal (top row, (a), (b) and (c)) and inland (bottom row, (d), (e) and (f)) Antarctica. Least squares regressions are noted with solid line and are significant at p < 0.01. The 4 samples with high Na<sup>+</sup> concentrations are denoted by blue open circles (b), the same as those in Figure 8 (the blue open circles). Note that the 4 samples were excluded in the plot of  $NO_3^-$  vs. nssCl<sup>-</sup> (c).



1003

**Figure 8.** Concentrations of NO<sub>3</sub><sup>-</sup> and Na<sup>+</sup> in surface snow samples on the coast. Four samples with high Na<sup>+</sup> concentrations are denoted by open circles, corresponding to those in Fig. 7b. Note that Na<sup>+</sup> concentrations in two samples, 2.5 and 2.8  $\mu$ eq L<sup>-1</sup> in parentheses, are above the maximum value of the secondary *y*-axis (Na<sup>+</sup> concentration). The sample in the dashed ellipse, with Na<sup>+</sup> concentration of 2.8  $\mu$ eq L<sup>-1</sup>, is the fresh snowfall.